Approximate Bayesian computation (ABC) gives exact results under the assumption of model error

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Summary

Approximate Bayesian computation (ABC) or likelihood-free inference algorithms are used to find approximations to posterior distributions without making explicit use of the likelihood function, depending instead on simulation of sample data sets from the model. In this paper we show that under the assumption of the existence of a uniform additive model error term, ABC algorithms give exact results when sufficient summaries are used. This interpretation allows the approximation made in many previous application papers to be understood, and should guide the choice of metric and tolerance in future work. ABC algorithms can be generalized by replacing the 0-1 cut-off with an acceptance probability that varies with the distance of the simulated data from the observed data. The acceptance density gives the distribution of the error term, enabling the uniform error usually used to be replaced by a general distribution. This generalization can also be applied to approximate Markov chain Monte Carlo algorithms. In light of this work, ABC algorithms can be seen as calibration techniques for implicit stochastic models, inferring parameter values in light of the computer model, data, prior beliefs about the parameter values, and any measurement or model errors.

Some key words: Approximate Bayesian computation; calibration; implicit inference; likelihood-free inference; Monte Carlo.

1. Introduction

Approximate Bayesian computation (ABC) algorithms are a group of methods for performing Bayesian inference without the need for explicit evaluation of the model likelihood function (Beaumont et al. (2002); Marjoram et al. (2003); Sisson et al. (2007)). The algorithms can be used with implicit computer models (Diggle & Gratton (1984)) that generate sample data sets rather than return likelihoods. ABC methods have become popular in the biological sciences with applications in genetics (see, for example, Siegmund et al. (2008); Foll et al. (2008)), epidemiology (Blum & Tran (2008); Tanaka et al. (2006)) and population biology (Ratmann et al. (2007); Hamilton et al. (2005); Cornuet et al. (2008)) most common. This popularity is primarily due to the fact that the likelihood function, which can be difficult or impossible to compute for some models, is not needed in order to do inference. However, despite their popularity little is known about the quality of the approximation they provide beyond results shown in simulation studies.

In this paper we give a framework in which the accuracy of ABC methods can be understood. The notation throughout this paper is as follows. Let \( \theta \) denote the vector of unknown model parameters we wish to infer, and let \( \eta(\cdot) \) denote the computer model. We assume \( \eta(\cdot) \) is stochastic, so that the model repeatedly run at \( \theta \) will give a range of possible model outputs, and write
\(X \sim \eta(\theta)\) to denote that \(X\) has the same distribution as the model run at \(\theta\). To distinguish the model output from the observed data, let \(D\) denote the observations. The aim is to calibrate the model to the data, in order to learn about the true value of the parameter. The Bayesian approach is to find the posterior distribution of \(\theta\) given \(D\), given by
\[
\pi(\theta \mid D) = \frac{\pi(D \mid \theta)\pi(\theta)}{\pi(D)}.
\]
Throughout this paper, \(\pi(\cdot)\) is used to denote different probability densities, and \(\pi(\cdot \mid \cdot)\) conditional densities, with the context clear from the arguments. Above, \(\pi(\theta)\) is the prior distribution, \(\pi(D \mid \theta)\) is the likelihood of the data under the model given parameter \(\theta\) (the probability distribution of \(\eta(\theta)\)), \(\pi(\theta \mid D)\) is the posterior distribution, and \(\pi(D)\) is the evidence for the model.

It is usual in Bayesian inference to find that the normalizing constant \(\pi(D)\) is intractable, and a wide range of Monte Carlo techniques have been developed to deal with this case (Liu (2001)). Doubly-intractable distributions are distributions which have a likelihood function \(\pi(D \mid \theta) = q(D \mid \theta)/c(\theta)\) which is known only up to a normalizing constant, \(c(\theta)\), which is intractable. Standard Monte Carlo techniques do not apply to these distributions, and Murray et al. (2006) have developed algorithms which can be used in this case. ABC methods are Monte Carlo techniques developed for use with completely-intractable distributions, where the likelihood function \(\pi(D \mid \theta)\) is not even known up to a normalizing constant. ABC algorithms, sometimes called likelihood-free algorithms, enable inference using only simulations generated from the model, and do not require any evaluation of the likelihood. The most basic form of the ABC algorithm is based on the rejection algorithm, and is as follows:

**Algorithm A: approximate rejection algorithm**

A1. Draw \(\theta \sim \pi(\theta)\)
A2. Simulate \(X\) from the model \(X \sim \eta(\theta)\)
A3. Accept \(\theta\) if \(\rho(X, D) \leq \delta\).

Here, \(\rho(\cdot, \cdot)\) is a distance measure on the model output space, and \(\delta\) is a tolerance determining the accuracy of the algorithm. Accepted values of \(\theta\) are not from the true posterior distribution, but from an approximation to it, written \(\pi(\theta \mid \rho(D, X) \leq \delta)\). When \(\delta = 0\) this algorithm is exact and gives draws from the posterior distribution \(\pi(\theta \mid D)\), whereas as \(\delta \to \infty\) the algorithm gives draws from the prior. While smaller values of \(\delta\) lead to samples which better approximate the true posterior, they also lead to lower acceptance rates in step A3 than larger values, and so more computation must be done to get a given sample size. Consequently, the tolerance \(\delta\) can be considered as controlling a trade-off between computability and accuracy.

Several extensions have been made to the approximate rejection algorithm. If the data are high dimensional, then a standard change to the algorithm is to summarize the model output and data, using a summary statistic \(S(\cdot)\) to project \(X\) and \(D\) onto a lower dimensional space. Algorithm A is then changed so that step A3 reads

A3’. Accept \(\theta\) if \(\rho(S(X), S(D)) \leq \delta\).

Ideally, \(S(\cdot)\) should be chosen to be a sufficient statistic for \(\theta\). However, if the likelihood is unknown, then sufficient statistics cannot be identified. Summarizing the data and model output using a non-sufficient summary adds another layer of approximation on top of that added by the use of the distance measure and tolerance, but again, it is not known what effect any given choice for \(S(\cdot)\) has on the approximation.
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Beaumont et al. (2002) contains two innovations; they replace the discrete 0-1 cut-off in step A3 with a weighting scheme, using an Epanechnikov kernel to weight each value of $\theta$ according to the value of the metric $\rho(D, X)$, so that large weights are assigned to values of $\theta$ which produce model output close to the measurement and small weights to those values which produce output distant from $D$. They then use the weighted sample $(\hat{\theta}_i, w_i)$ to train a local-linear regression to model the posterior density. Blum & François (2008) have since extended this by using an adaptive heteroscedastic model to estimate the posterior from the weighted sample. While it has been shown that using a weighting scheme improves the accuracy of the approximate rejection algorithm in several situations, it is still unclear what the approximation represents or why any given weighting should be preferred over any other.

In this paper it is shown that the basic approximate rejection algorithm can be interpreted as performing exact inference in the presence of uniform model or measurement error. Similarly, the weighting scheme used in Beaumont et al. (2002) corresponds to an error with an Epanechnikov distribution. In other words, it is shown that ABC gives exact inference for the wrong model, and we give a distribution for the model error term for whatever choice of metric and tolerance are used. This interpretation allows us to show the effect a given choice of metric, tolerance and weighting have had in previous applications, and should provide guidance when choosing metrics and weightings in future work. It is also shown that Algorithm A can be generalized to give inference under the assumption of a completely flexible form for the model error. We discuss how to model the model error, and show how some models can be rewritten to give exact inference.

Finally, ABC has been extended by Marjoram et al. (2003) from the rejection algorithm to approximate Markov chain Monte Carlo algorithms, and by Sisson et al. (2007) and Beaumont et al. (2008) to approximate sequential Monte Carlo algorithms. We extend the approximate Markov chain Monte Carlo algorithm to give inference for a general form of error, and suggest methods for calculating Bayes factors and integrals for completely-intractable distributions.

2. INTERPRETING ABC

In this section a framework is described which enables the effect a given metric and weighting have in ABC algorithms to be understood. This will then allow us to improve the inference by carefully choosing a metric and weighting which more closely represents our true beliefs. The key idea is to assume that there is a discrepancy between the best possible model prediction and the data. This discrepancy represents either measurement error on the data, or model error describing our statistical beliefs about where the model is wrong. George Box famously wrote that ‘all models are wrong’, and in order to link models to reality it is necessary to account for this model error when performing inference. In the context of deterministic models, this practice is well established (Campbell (2006); Goldstein & Rougier (2008); Higdon et al. (2008)), and should also be undertaken when linking stochastic models to reality, despite the fact that the variability in the model can seemingly explain the data as they are.

The framework introduced here uses the best input approach, similar to that given in Kennedy & O’Hagan (2001). We assume that the measurement $D$ can be considered as a realization of the model run at its best input value, $\hat{\theta}$, plus an independent error term $\epsilon$

\[ D = \eta(\hat{\theta}) + \epsilon. \]  

The error $\epsilon$ might represent measurement error on $D$, or model error in $\eta(\cdot)$, or both, in which case we write $\epsilon = \epsilon_1 + \epsilon_2$. Discussion about the validity of Equation (1), and what $\epsilon$ represents
and how to model it are delayed until Section 3, and for the time being we simply consider $\epsilon$ to have density $\pi_\epsilon(\cdot)$. The aim is to describe our posterior beliefs about the best input $\hat{\theta}$ in light of the error $\epsilon$, the data $D$, and prior beliefs about $\hat{\theta}$. Consider the following algorithm:

**Algorithm B: probabilistic approximate rejection algorithm**

B1 Draw $\theta \sim \pi(\theta)$
B2 Simulate $X$ from the model $X \sim \eta(\theta)$
B3 Accept $\theta$ with probability $\frac{\pi_\epsilon(D - X)}{c}$.

Here, $c$ is a constant chosen to guarantee that $\pi_\epsilon(D - X)/c$ defines a probability. For most cases we will expect $\epsilon$ to have a modal value of 0, and so taking $c = \pi_\epsilon(0)$ will make the algorithm valid and also ensure efficiency by maximizing the acceptance rate. If $D$ and $X$ are both real valued arrays of matching dimension, then $D - X$ is simply the arithmetic pairwise difference. However, if $D$ and $X$ are not real-valued, for example, $D$ and $X$ could both be gene sequences, then $D - X$ represents the difference between the two data sets (we could write $D - X = \rho(D, X)$) and $\pi_\epsilon(\cdot)$ is a distribution on this space of differences.

The main innovation in this paper is to show that Algorithm B gives exact inference for the statistical model described above by Equation (1). This is essentially saying that ABC gives exact inference, but for the wrong model.

**Theorem 1.** Algorithm B gives draws from the posterior distribution $\pi(\hat{\theta} \mid D)$ under the assumption that $D = \eta(\hat{\theta}) + \epsilon$ and $\epsilon \sim \pi_\epsilon(\cdot)$ independently of $\eta(\hat{\theta})$.

**Proof.** Let

$$I = \begin{cases} 
1 & \text{if } \theta \text{ is accepted} \\
0 & \text{otherwise.} 
\end{cases}$$

We then find that

$$\text{pr}(I = 1 \mid \theta) = \int \text{pr}(I = 1 \mid \eta(\theta) = x, \theta)\pi(x \mid \theta)dx$$

$$= \int \frac{\pi_\epsilon(D - x)}{c}\pi(x \mid \theta)dx.$$ 

This gives that the distribution of accepted values of $\theta$ is

$$\pi(\theta \mid I = 1) = \frac{\pi(\theta) \int \pi_\epsilon(D - x)\pi(x \mid \theta)dx}{\int \pi(\theta') \int \pi_\epsilon(D - x)\pi(x \mid \theta')dx d\theta'}.$$ 

To complete the proof we must find the posterior distribution of the best model input $\hat{\theta}$ given the data $D$ under the assumption of model error. Note that $\pi(D \mid \eta(\hat{\theta}) = x) = \pi_\epsilon(D - x)$ which implies that the likelihood of $\theta$ is

$$\pi(D \mid \hat{\theta}) = \int \pi(D \mid \eta(\hat{\theta}) = x, \hat{\theta})\pi(x \mid \hat{\theta})dx$$

$$= \int \pi_\epsilon(D - x)\pi(x \mid \hat{\theta})dx.$$
Approximate Bayesian computation (ABC) gives exact results under the assumption of model error. Consequently, the posterior distribution of $\theta$ is

$$\pi(\theta \mid D) = \frac{\pi(\theta) \int \pi(x|\theta)(D-x) \pi(x|\hat{\theta})dx}{\int \pi(\theta) \int \pi(x|\theta)(D-x) \pi(x|\hat{\theta})dx d\theta}$$

which matches the distribution of accepted values from Algorithm B.

To illustrate the algorithm, we consider the toy example used in Sisson et al. (2007) and again in Beaumont et al. (2008) where analytic expressions can be calculated for the approximations.

**Example 1.** Assume the model is a mixture of two normal distributions with a uniform prior for the mean:

$$\eta(\theta) \sim \frac{1}{2}N(\theta, 1) + \frac{1}{2}N(\theta, \frac{1}{100}), \, \theta \sim \mathcal{U}[-10, 10].$$

Further assume that we observe $D = 0$, but that there is measurement error $\varepsilon$ on this data. If $\varepsilon \sim \mathcal{U}[-\delta, \delta]$, which is the assumption made when using Algorithm A with $\rho(x, 0) = |x|$, then it is possible to show that the approximation is

$$\pi(\theta \mid \varepsilon \sim \mathcal{U}[-\delta, \delta], D = 0) \propto \Phi(\delta - \theta) - \Phi(-\delta - \theta) + \Phi(10(\varepsilon - \theta)) - \Phi(-10(\varepsilon + \theta))$$

for $\theta \in [-10, 10]$, where $\Phi(\cdot)$ is the standard Gaussian distribution function. Note that this is slightly different to the distribution given in Beaumont et al. (2008). An alternative to assuming uniform error, is to suppose that the error has a normal distribution $\varepsilon \sim N(0, \delta^2/3)$. It can then be shown that the posterior distribution of $\theta$ is

$$\pi(\theta \mid \varepsilon \sim N(0, \frac{\delta^2}{3}), D = 0) \propto \frac{1}{2} \phi(\theta; 0, 1 + \frac{\delta^2}{3}) + \frac{1}{2} \phi(\theta; 0, \frac{1}{100} + \frac{\delta^2}{3})$$

truncated onto $[-10, 10]$. This is the approximation found when using Algorithm B with a Gaussian acceptance kernel, where $\phi(\cdot; \mu, \sigma^2)$ is the probability density function of a Gaussian distribution with mean $\mu$ and variance $\sigma^2$. The value of the variance, $\delta^2/3$, is chosen to be equal to the variance of a $\mathcal{U}[-\delta, \delta]$ random variable. For large values of the tolerance $\delta$, the difference between the two approximations can be significant (see Figure 1), but in the limit as $\delta$ tends to zero, the two approximations will be the same, corresponding to zero error.

### 3. Model discrepancy

The interpretation of ABC given by Theorem 1 allows us to revisit previous analyses done using the ABC algorithm, and to understand the approximation in the posterior in terms of the distribution implicitly assumed for the error term. If the approximate rejection algorithm (Algorithm A) was used to do the analysis, we can see that this is equivalent to using the acceptance probability

$$\frac{\pi_c(r)}{c} = \begin{cases} 1 & \text{if } \rho(r) \leq \delta \\ 0 & \text{otherwise} \end{cases}$$

where $r$ is the distance between the simulated and observed data. This says that Algorithm A gives exact inference for the model which assumes a uniform measurement error on the region defined by the 0-1 cut-off, i.e.,

$$\varepsilon \sim \mathcal{U}\{x : \rho(x, D) \leq \delta\}.$$
If $\rho(\cdot, \cdot)$ is a Euclidean metric, $\rho(D, x) = (x - D)^T(x - D)$, this is equivalent to assuming uniform measurement error on a ball of radius $\delta$ about $D$.

The weighting scheme used in Beaumont et al. (2002), and since used in numerous application papers, used an Epanechnikov kernel $K_\delta(\|x - D\|)$ to weight each value of $\theta$. The form of this density is

$$K_\delta(r) = \frac{3}{4\delta} \left(1 - \frac{r^2}{\delta^2}\right) I_{r \leq \delta}. \quad (2)$$

The kernel gives the probability density function of the error term, and thus analysis done using an Epanechnikov kernel gives results which assume error with this distribution. In most situations, it is likely to be a poor choice for a model of the measurement error, because the tails of the distribution are short, with zero mass outside of the interval $[-\delta, \delta]$.

There are two ways we can choose to view the error term: either as measurement error or model error. Interpreting $\epsilon$ to represent measurement error is relatively straightforward, as scientists usually hold beliefs about the distribution and magnitude of measurement error on their data. For most problems, assumptions of uniform measurement error will be inappropriate, and so using Algorithm A with a 0-1 cut-off will be inappropriate. But we have shown how to replace this uniform assumption with a distribution which more closely represents the beliefs of the scientist. Although the distribution of the measurement error will often be completely specified by the scientist, for example zero-mean Gaussian error with known variance, it is possible to include unknown parameters for the distribution of $\epsilon$ in $\theta$ and infer them along with the model parameters. Care needs to be taken to choose the constant $c$ so that the acceptance rate in step B3 is less than one for all values of the parameter, but other than this it is in theory simple to infer error parameters along with the model parameters. So for example, if $\epsilon \sim N(0, \sigma^2)$, where $\sigma^2$ is unknown, we could include $\sigma^2$ in $\theta$. 

![Fig. 1. The posterior distributions found when using Algorithm A (solid line) and Algorithm B (dashed line) with a Gaussian acceptance kernel. The left plot is for $\delta = 1$ and the right plot for $\delta = 0.1$.](image)
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Some models have sampling or measurement error built into the computer code so that the
model output includes a realization of this noise. Rather than coding the noise process into the
model, it will sometimes be possible to rewrite the model so that it outputs the latent underlying
signal. If the likelihood of the data given the latent signal is computable (as it often is), then it
may be possible to analytically account for the noise with the acceptance probability \( \pi_i(\cdot) \). ABC
methods have proven most popular in fields such as genetics, epidemiology, and population bi-
ology, where a common occurrence is to have data generated by sampling a hidden underlying
tree structure. In many cases, it is the partially observed tree structure which causes the likeli-
hood to be intractable, and given the underlying tree the sampling process will have a known
distribution. If this is the case (and if computational constraints allow), we can use the proba-
bilistic ABC algorithm to do the sampling to give exact inference without any assumption of
model error. Note that if the sampling process gives continuous data, then exact inference using
the rejection algorithm would not be possible, and so this approach has the potential to give a
significant improvement over current methods.

Example 2. To illustrate the idea of rewriting the model in order to do analytic sampling,
we describe a version of the problem considered in Plagnol & Tavaré (2004). Their aim was to
use the primate fossil record to date the divergence time of the primates. They used an inho-
ogeneous branching process to model speciation, with trees rooted at time \( t = \tau \), and simul-
lated forwards in time to time \( t = 0 \), so that the depth of the tree, \( \tau \), represents the divergence
time of interest. The branching process is parametrized by \( \lambda \), which can either be estimated and
fixed, or treated as unknown and given a prior distribution. Time is split into geologic epochs
\( \tau < t_k < \cdots < t_i < 0 \), and the data consist of counts of the number of primate species that have
been found in each epoch of the fossil record, \( D = (D_1, \ldots, D_k) \). Fossil finds are modelled by
a discrete marking process on the tree, with each species having equal probability \( \alpha \) of being
preserved as a fossil in the record. If we let \( N_i \) be the cumulative number of branches that ex-
ist during any point of epoch \( i \), then the model used for the fossil finds process can be written
as \( D_i \sim \text{Binomial}(N_i, \alpha) \). The distribution of \( N = (N_1, \ldots, N_{14}) \) cannot be calculated explicit-
ly and so we cannot use a likelihood based approach to find the posterior distribution of the
unknown parameter \( \theta = (\lambda, \tau, \alpha) \). The ABC approach used in Plagnol & Tavaré (2004) was to
draw a value of \( \theta \) from its prior, simulate a sample tree and fossil finds, and then count the num-
ber of simulated fossils in each epoch to find a simulated value of the data \( X \). They then accepted
\( \theta \) if \( \rho(D, X) \leq \delta \) for some metric \( \rho(\cdot, \cdot) \) and tolerance \( \delta \). This gives an approximation to the pos-
terior distribution of the parameter given the data and the model, where the approximation can
be viewed as model or measurement error.

However, instead of approximating the posterior, it is possible in theory to rewrite the model
and perform the sampling analytically to find the exact posterior distribution:

1. Draw \( \theta = (\lambda, p, \alpha) \sim \pi(\cdot) \)
2. Simulate a tree \( T \) using parameter \( \lambda \) and count \( N \)
3. Accept \( \theta \) with probability \( \prod_{i=1}^k \left( \frac{N_i}{D_i} \right) \alpha_i^D (1 - \alpha)^{N_i - D_i} \).

This algorithm gives exact draws from the posterior distribution of \( \theta \) given \( D \), and in theory
there is no need for any assumption of measurement error. Note that \( \theta \) can include parameter \( \alpha \)
for the sampling rate, to be inferred along with the other model parameters. However, this makes
finding a normalizing constant in step 3 difficult. Without a normalizing constant to increase the
acceptance rate, applying this algorithm directly will be slow for many values of \( D \) and \( k \) (the
choice of prior distribution and number of parameters we choose to include in \( \theta \) can also have
a significant effect on the efficiency). A practical solution would be to add an error term and
assume the presence of measurement error on the data (which is likely to exist in this case), in
order to increase the acceptance probability in step 3. Approaching the problem in this way, it is
possible to carefully model the error on \( D \) and improve the estimate of the divergence time.

Using \( \epsilon \) to represent measurement error is straightforward, whereas using \( \epsilon \) to model the model
discrepancy (to account for the fact the model is wrong) is harder to conceptualize and not as
commonly used. For deterministic models, the idea of using a model error term when doing
calibration or data assimilation is well established and described for a Bayesian framework in
Kennedy & O’Hagan (2001). They assume that the model run at its ‘best’ input, \( \eta(\hat{\theta}) \), is sufficient
for the model when estimating \( \hat{\theta} \). In other words, knowledge of the model run at its best input
provides all the available information about the system for the purpose of prediction. If this is the
case, then we can define \( \epsilon \) to be the difference between \( \eta(\hat{\theta}) \) and \( D \), and assume \( \epsilon \) is independent
of \( \eta(\hat{\theta}) \). Note that the error is the difference between the data and the model run at its best input,
and does not depend on \( \theta \). If we do not include an error term \( \epsilon \), then the best input is the value of
\( \theta \) that best explains the data, given the model. When we include an error term which is carefully
modelled and represents our beliefs about the discrepancy between \( \eta(\cdot) \) and reality, then it can be
argued that \( \theta \) represents the ‘true’ value of \( \theta \), and that \( \pi(\hat{\theta} \mid D, \epsilon \sim \pi(\cdot)) \) should be our posterior
distribution for \( \hat{\theta} \) in light of the data and the model.

For deterministic models, Goldstein & Rougier (2008) provide a framework to help think
about the model discrepancy. To specify the distribution of \( \epsilon \), it can help to break the discrep-
ancy down into various parts: physical processes not modelled, errors in the specification of the
model, imperfect implementation etc. So for example, if \( \eta(\cdot) \) represents a global climate model
predicting average temperatures, then common model errors could be not including processes
such as clouds, \( CO_2 \) emissions from vegetation etc., error in the specification might be using an
unduly simple model of economic activity, and imperfect implementation would include using
grid cells too large to accurately solve the underlying differential equations. In some cases it
may be necessary to consider model and measurement error, \( \epsilon + \epsilon \) say, and model each process
separately. For stochastic models, as far as we are aware, no guidance exists about how to model
the error, and indeed it is not clear what \( \epsilon \) should represent.

To complicate matters further, for many models the dimension of \( D \) and \( X \) will be large,
making it likely that the acceptance rate \( \pi(\epsilon \mid X - D) \) will be small. As noted above, in this case it
is necessary to summarize the model output and the data using a multidimensional summary \( S(\cdot) \).
Using a summary means that rather than approximating \( \pi(\theta \mid D) \), the algorithms approximate
\( \pi(\theta \mid S(D)) \). The interpretation of \( \epsilon \) as model or measurement error still holds, but now the error
is on the measurement \( S(D) \) or the model prediction \( S(X) \). If each element of \( S(\cdot) \) has an
interpretation in terms of a physical process, this may make it easier to break the error down into
independent components. For example, suppose that we use \( S(x) = (x, s_{xx}), \) the sample mean
and variance of \( X \), and that we then use the following acceptance density

\[
\pi(\epsilon(S(X) - S(D)) = \pi_1(\bar{X} - \bar{D})\pi_2(s_{XX} - s_{DD}).
\]

This is equivalent to assuming that there are two sources of model error. Firstly, the mean predic-
tion is assumed to be wrong, with the error distributed with density \( \pi_1(\cdot) \). Secondly, it assumes
that the model prediction of the variance is wrong, with the error having distribution \( \pi_2(\cdot) \). It
also assumes that the error in the mean prediction is independent of the error in the variance
prediction. This independence is not necessary, but helps with visualization and elicitation. For
this reason it can be helpful to choose the different components of \( S(\cdot) \) so that they are close
to independent (independence may also help increase the acceptance rate). Another possibility
for choosing \( S(\cdot) \) is to use principal component analysis (if \( \dim(X) \) is large) to find a smaller
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number of uncorrelated summaries of the data which may have meaningful interpretations. In
general however, it is not known how to choose good summaries. Joyce & Marjoram (2008)
have suggested a method for selecting between different summaries and for deciding how many
summaries it is optimal to include. However, more work is required to find summaries which are
informative, interpretable and for which we can describe the model error.

Finally, once we have specified a distribution for \( \epsilon \), we may find the acceptance rate is too small
to be practicable and that it is necessary to compromise (as in Example 2 above). A pragmatic
way to increase the acceptance rate is to use a more disperse distribution for \( \epsilon \). This moves us
from the realm of using \( \epsilon \) to model an error we believe exists, to using it to approximate the true
posterior. This is currently how most ABC methods are used. However, even when making a
pragmatic compromise, the interpretation of the approximation in terms of an error will allow us
to think more carefully about how to choose between different compromise solutions.

Example 3. One of the first uses of an ABC algorithm was by Pritchard et al (1999), who used
a simple stochastic model to study the demographic history of the \( Y \) chromosome, and used an
approximate rejection algorithm to infer mutation and demographic parameters for their model.
Their data consisted of 445 \( Y \) chromosomes sampled at eight different loci from a mixture of
populations from around the world, which they summarized by just three statistics: the mean
(across loci) of the variance of repeat numbers \( V \), the mean effective heterozygosity \( H \), and
the number of distinct haplotypes \( N \). The observed value of the summaries for their sample
was \( D \equiv (V, H, N)^T = (1.149, 0.6358, 316)^T \). They elicited prior distributions for the mutation
rates from the literature, and used diffuse priors for population parameters such as the growth rate
and the effective number of ancestral \( Y \) chromosomes. Population growth was modelled using
a standard coalescent model growing at an exponential rate from a constant ancestral level, and
various different mutation models were used to simulate sample values for the three summaries
measured in the data. They then applied Algorithm A using the metric

\[
\rho(D, X) = \prod_{i=1}^{3} \frac{D_i - X_i}{D_i}
\]

(3)

where \( X \) is a triplet of simulated values for the three summaries statistics. They used a tolerance
value of \( \delta = 0.1 \), which for their choice of metric corresponds to an error of 10% on each
measurement. This gives results equivalent to assuming that there is independent uniform
measurement error on the three data summaries, so that the true values of the three summaries have
the following distributions

\[ V \sim U[1.0341, 1.2624], \quad H \sim U[0.58122, 0.71038], \quad N \sim U[284, 348]. \]

Beaumont et al. (2002) used the same model and data set to compare the relative performance of
Algorithm A with an algorithm similar to Algorithm B, using an Epanechnikov density applied
to the metric value (3) for the acceptance probability \( \pi_\epsilon(\cdot) \). They set a value of \( \delta \) (the cut-off in
Algorithm A and the range of the support for \( \epsilon \) in Algorithm B) by using a quantile \( P_\delta \) of the
empirical distribution function of simulated values of \( \rho(D, X) \), i.e., \( P_{0.01} \) means they accepted
the 1% of model runs with values closest to \( D \). They concluded that Algorithm B gives more
accurate results than Algorithm A, meaning that the distribution found using Algorithm B is
closer to the posterior found when assuming no measurement error (\( \delta = 0 \)).

The conclusion that Algorithm B is preferable to Algorithm A for this model is perhaps not
surprising in light of what we now know, as it was not taken into account that both algorithms
used the same value of \( \delta \). For Algorithm A this corresponds to assuming a measurement error
with variance \( \delta^2 / 3 \), whereas using acceptance probability (2) is equivalent to assuming a
4. APPROXIMATE MARKOV CHAIN MONTE CARLO

For problems which have a tightly constrained posterior distribution (relative to the prior), repeatedly drawing independent values of $\theta$ from its prior distribution in the rejection algorithm can be inefficient. For problems with a high dimensional $\theta$ this inefficiency is likely to make the application of a rejection type algorithm impracticable. The idea behind Markov chain Monte Carlo (MCMC) is to build a Markov chain on $\theta$ and correlate successive observations so that more time is spent in regions of high posterior probability. Most MCMC algorithms, such as the Metropolis-Hastings algorithm, depend on knowledge of the likelihood function which we have assumed is not known. Marjoram et al. (2003) give an approximate version of the Metropolis-Hastings algorithm, which approximates the acceptance probability by using simulated model output with a metric and a 0–1 cut-off to approximate the likelihood ratio. This, as before, is equivalent to assuming uniform error on a set defined by the metric and the tolerance. As above, this algorithm can be generalized from assuming uniform measurement error to an arbitrary error term. Below, are two different algorithms to perform MCMC for the model described by Equation (1). The difference between the two algorithms lies in the assumption about on which variable to belong to the space of parameter values $\Theta$.

Algorithm C: probabilistic approximate MCMC 1

C1 At time $t$, propose a move from $\theta_t$ to $\theta'$ according to transition kernel $q(\theta_t, \theta')$.
C2 Simulate $X' \sim \eta(\theta')$.
C3 Set $\theta_{t+1} = \theta'$ with probability

$$r(\theta_t, \theta' | X') = \frac{\pi(\theta')}{\pi(\theta_t)} \min \left( 1, \frac{q(\theta', \theta_t)\pi(\theta_t)}{q(\theta_t, \theta')\pi(\theta')} \right),$$

(4)

otherwise set $\theta_{t+1} = \theta_t$.

An alternative approach is to introduce the value of the simulated output as an auxiliary variable and construct the Markov chain on the space $\Theta \times X$, where $X$ is the space of model outputs.

Algorithm D: probabilistic approximate MCMC 2

D1 At time $t$, propose a move from $\psi_t = (\theta_t, X_t)$ to $\psi' = (\theta', X')$ with $\theta'$ drawn from transition kernel $q(\theta_t, \theta')$, and $X'$ simulated from the model using $\theta'$:

$$X' \sim \eta(\theta')$$

D2 Set $\psi_{t+1} = (\theta', X')$ with probability

$$r((\theta_t, X_t), (\theta', X')) = \min \left( 1, \frac{\pi(\theta')q(\theta', \theta_t)\pi(\theta_t)}{\pi(\theta')q(\theta_t, \theta')\pi(\theta_t)} \right),$$

(5)

otherwise set $\psi_{t+1} = \psi_t$. 

433 measurement error with variance $\delta^2/5$. Therefore, using Algorithm B uses measurement error only 60% as variable as that assumed in Algorithm A, and so it is perhaps not surprising that Algorithm B gives more accurate results in this case.
Approximate Bayesian computation (ABC) gives exact results under the assumption of model error.

Proof of convergence. To show that these Markov chains converge to the required posterior
distribution, it is sufficient to show that the chains satisfy the detailed balance equations
\[ p(s, t)\pi(t) = p(t, s)\pi(s) \quad \text{for all } s, t, \]
where \( p(\cdot, \cdot) \) is the transition kernel of the chain and \( \pi(\cdot) \) the required stationary distribution.

For Algorithm C the transition kernel is the product of \( q(\theta, \theta') \) and the acceptance rate. To
calculate the acceptance rate, note that in Equation (4) the acceptance probability is conditioned
upon knowledge of \( X' \) and so we must integrate out \( X' \) to find \( r(\theta, \theta') \). This gives the transition
kernel for the chain:
\[ p(\theta, \theta') = q(\theta, \theta') \int_{\mathcal{X}} \frac{\pi_c(D - X')}{c} \min\left(1, \frac{q(\theta', \theta)\pi(\theta)}{q(\theta, \theta')\pi(\theta')}\right) \pi(X' \mid \theta) dX'. \]
The target stationary distribution is
\[ \pi(D \mid \theta) = \int_{\mathcal{X}} \pi_c(D - X)\pi(X \mid \theta) dX. \]
It is then simple to show that the Markov chain described by Algorithm C satisfies the detailed
balance equations.

For Algorithm D, the transition kernel is
\[ p((\theta, X), (\theta', X')) = q(\theta, \theta')\pi(X' \mid \theta') \min\left(1, \frac{\pi_c(D - X')q(\theta', \theta)\pi(\theta')}{\pi_c(D - X)q(\theta, \theta')\pi(\theta)}\right). \quad (6) \]
The Markov chain in this case takes values on in \( \Theta \times \mathcal{X} \) and the required stationary distribution is
\[ \pi(\theta, X \mid D) = \frac{\pi_c(D - X)\pi(X \mid \theta)\pi(\theta)}{\pi(D)} \quad (7) \]
It can then be shown that Equations (6) and (7) satisfy the detailed balance equations.  □

While Algorithm C is more recognisable as a generalization of the approximate MCMC algo-
rithm given in [Marioram et al. (2003)], Algorithm D is likely to be more efficient in most cases.
This is because the ratio of model error densities that occurs in acceptance rate (5) is likely to
result in larger probabilities than those given by Equation (4) which simply has a \( \pi_c(D - x)/c \)
term instead. Algorithm D also has the advantage of not requiring a normalizing constant.

5. Extensions

5.1. Monte Carlo integration

Suppose our aim is to calculate expectations of the form
\[ E(f(\theta) \mid D) = \int f(\theta)\pi(\theta \mid D)d\theta \]
where the expectation is taken with respect to the posterior distribution of \( \theta \). The simplest way
to approach this is to draw a sample of \( \theta \) values \( \{\theta_i\}_{i=1,\ldots,n} \) from \( \pi(\theta \mid D) \) using Algorithm B,
C or D and then approximate using the sum \( n^{-1} \sum f(\theta_i) \). However, a more stable estimate can
be obtained by using draws from the prior weighted by \( \pi_c(D - X_i) \) as in Algorithm B. For each
\( \theta \) drawn from the prior in step B1, assign it weight \( w_i = \pi(D - X_i) \). Then an estimator of the
required expectation is
\[ \frac{\sum f(\theta_i)w_i}{\sum w_i} \].

Note that all values of \( \theta \) drawn from the prior are used in the sum; there is no rejection step. This is a direct extension of the estimate given in [Beaumont et al. (2002)] which used Epanechnikov kernels to weight each value of \( \theta \).

### 5.2. Approximate model selection

The theoretical acceptance rate from the rejection algorithm (Algorithm A with \( \delta = 0 \)) is equal to the model evidence \( \pi(D) \). The evidence from different models can then be used to calculate Bayes factors which can be used to perform model selection ([Kass & Raftery (1995)]). It is possible to approximate the value of \( \pi(D) \) by using the acceptance rate from Algorithm A. By doing this for two or more competing models, we can perform approximate model selection, although in practice this approach can be unstable ([Wilkinson (2007)]). The estimate of \( \pi(D) \) can be improved and made interpretable by using the weighted estimate

\[ \frac{1}{n} \sum_{i=1}^{n} \frac{1}{m} \sum_{j=1}^{m} \pi(\epsilon(D - X_{ij}^{\theta})) \]

where \( X_{1i}, \ldots, X_{mi} \sim \eta(\theta_i) \) and \( \theta_1, \ldots, \theta_n \sim \pi(\cdot) \). This gives a more stable estimate than simply taking the acceptance rate, and also tends to the exact value (as \( n, m \to \infty \)) for the model given by Equation (1).

### 6. Discussion

It has been shown in this paper that approximate Bayesian computation algorithm can be considered to give exact inference under the assumption of model error. However, this is only part of the way towards a complete understanding of ABC algorithms. In the majority of the application papers using ABC methods, summaries of the data and model output have been used to reduce the dimension of the output. It cannot be known whether these summaries are sufficient for the data, and so in most cases the use of summaries means that there is another layer of approximation. While this work allows us to understand the error assumed on the measurement of the summary, it says nothing about what effect using the summary rather than the complete data has on the inference. More work is required to discover this, and to help guide the choice of which summaries to use.

The use of a model error term when making inferences is important if one wants to move from making statements about the model to statements about reality. There has currently been only minimal work done on modelling the discrepancy term for stochastic models. One way to approach this is to view the model as deterministic, outputting a density \( \pi_{\theta}(x) \) for each value of the input \( \theta \) (many realizations of \( \eta(\theta) \) would be needed to learn \( \pi_{\theta}(x) \)). The discrepancy term \( \epsilon \) can then be considered as representing the difference between \( \pi_{\theta}(x) \) and the true variability inherent in the physical system (at least the variability given the level we choose to model it at). Finally, it should be possible to generalize approximate sequential Monte Carlo methods in a similar way to that done for the approximate rejection and approximate Markov chain Monte Carlo algorithms.
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