# densEstBayes: density estimation via Bayesian inference engines

MATT P WAND

University of Technology Sydney

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## 1 Introduction

Probability density function estimation, or density estimation for short, based on a univariate sample is a problem of fundamental importance in statistics and data analysis. Approaches to the density estimation problem number in the hundreds and stretch over several decades, but shortcomings such as reliable choice of smoothing parameter are typical. Moreover, the vast majority of density estimation procedures are frequentist with no allowance made for the variability due to smoothing parameter choice. The R package densEstBayes provides high-quality Bayesian density estimates based the relatively recent Bayesian inference engine paradigm. It makes use of the Bayesian inference engines provided by the Star platform (Carpenter et al., 2017). Other options include slice sampling (e.g. Neal, 2003) and semiparametric mean field variational Bayes (e.g. Rohde & Wand, 2016). The infrastructure of densEstBayes is such that new and improved Bayesian inference engines can be incorporated when they are developed.

All density estimates provided by densEstBayes are accompanied with pointwise credible intervals that allow for variability in smoothing parameter selection. The methodology, based on binning, scales well to very large sample sizes. Full details of the Bayesian inference engine approach to density estimation are given in Wand & Yu (2020).

## 2 Illustrations for the 2011 Old Faithful Gevser Data

The OldFaithful dataset consists of the time intervals, in minutes, for all 3,507 adjacent pairs of eruptions of the Old Faithful Geyser in Yellowstone National Park, U.S.A., during 2011. The following R commands:

- > library(densEstBayes)
- > hist(OldFaithful2011,col = "gold",main = "",probability = TRUE,
- + xlab = "time interval between geyser eruptions (minutes)")

lead to the probability histogram shown in Figure 1:

A quick-to-compute Bayesian density estimate, based on the Bayesian inference engine paradigm described in Wand & Yu (2020), is obtained via the command:

```
> destSMFVB <- densEstBayes(OldFaithful2011.method = "SMFVB")
```

The method = "SMFVB" specification leads to semiparametric mean field variational Bayes (e.g. Rohde & Wand, 2016) being used for approximate Bayesian inference. A plot of the estimate is obtained using:

- > plot(destSMFVB,xlab = "time interval between geyser eruptions (minutes)",
- + main = "method = \"SMFVB\" (semiparametric mean field variational Bayes)")
- > rug(jitter(OldFaithful2011,amount = 0.2),col = "dodgerblue")

and is shown in Figure 2. The shaded region corresponds to approximate 95% pointwise credible intervals. A bimodal distribution is apparent, with the main mode at around 90 minutes and a secondary mode at around 65 minutes.

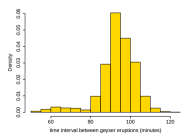


Figure 1: A probability histogram for the time intervals (minutes) between all 3,507 adjacent pairs of eruptions of the Old Faithful Geyser during the year 2011. The histogram bins correspond to the default call to the R function hist().

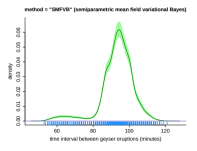


Figure 2: Bayesian density estimate for the 2011 Old Faithful geyser data introduced in Figure 1. The pale green shaded region corresponds to approximate 95% pointwise credible intervals. The data are shown as tick marks, with some jittering, at the base of the plot.

A density estimate based on the no-U-turn sampler (Hoffman & Gelman, 2014) as implemented in Stan (Carpenter et al., 2017). Bayesian inference engine is obtained and plotted via the commands:

> destNUTS <- densEstBayes(OldFaithful2011,method = "NUTS")
> plot(destNUTS,xlab = "time interval between geyser eruptions (minutes)",
+ main = "method = \"NUTS\" (no-U-turn sampler)")

## > rug(iitter(NldFaithful2011 amount = 0.2) col = "dodgerhlue")

with the result displayed in Figure 3. The estimate and variability band are very similar to those obtained via semiparametric mean field variational Bayes with method = "SMFVB". Even though method = "SMFVB" can provide a fast Bayesian density estimate, it involves solving a high-dimensional optimisation problem for which convergence is difficult to guarantee for central input data sets. Eurither details on this issue are given in Section 4.1 The Figure 3.

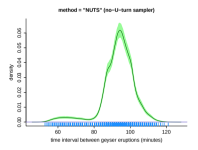


Figure 3: Bayesian density estimate for the data from Figure 1 using the no-U-turn sampler. The pale green shaded region corresponds to approximate 95% pointwise credible intervals. The data are shown as tick marks, with same rittering at the base of the plot.

density estimate is based on a version of Markov chain Monte Carlo sampling and at least a cursory check of the relevant chains is well-advised. The function checkChains() facilitates graphical checks of the chains corresponding to vertical slices of the density estimate at 5 equally-spaced abscissae. The vertical lines in Figure 4 indicate the locations of the default vertical slices. The command:

#### > checkChains(destNUTS)

produces the graphic given in Figure 5. In this case, all chains seem to be well-behaved and the method = "NUTS" Bayesian density estimate shown in Figures 3 and 4 is trustworthy.

The commands:

## > destDefault <- densEstBayes(OldFaithful2011) ; plot(destDefault)

produce and plot the default densEstBayes() estimate, and is based on slice sampling (method = "slice"). Usually the slice sampling-based estimate is computed more quickly than the Standependent estimates. This is because the underlying C++ code for method = "slice" focuses on the specific model used for Bayesian density estimation whereas the method = "HMC" and method = "NUTS" use a Bayesian inference engine that is designed to handle a very general class of Bayesian models.

# 3 Controlling the Estimation Procedure

Various settings for Bayesian density estimation via the densEstBayes() can be controlled using the control argument in calls to the function densEstBayes(). For example, the following

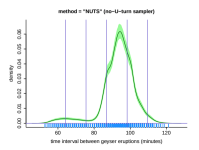


Figure 4: The same as for Figure 3 but with the addition of lines that indicate the positions of the vertical slices for which chains for the log-density estimate are monitored by the function check(ha) ins().

vertical slice		14	
of log-density	trace	lag 1	acf
function			
1/6 of range	Marchagalteinacha Aranemarkeinigeachd		
2/6 of range	irgidesirjetije Alabesirjetije		
3/6 of range	raciberaly landal, population		
4/6 of range	iti pirgilgi kiti kasalaat Kapisasi a laydisi yad		
5/6 of range			

Figure 5: The graphic produced by the command checkChains(destNUTS) where densNUTS is the densEstBayes() object corresponding to the estimate shown in Figure 5. The chains correspond to the estimates of the log-density function at the vertical slices shown in Figure 4. The second column is a trace (time series) plot of the chain. The third column is a scatterplot of the chain values against their previous (lag 1) values. The fourth column is the sample autocorrelation function, based on the R function acf().

command increases the warm-up size to 2,000, the lengths of the kept chains to 3,500 and then thins the kept chains by a factor of 5 when method = "HMC":

```
> destLongerChains <- densEstBayes(OldFaithful2011,method = "HMC",

+ control = densEstBayes.control(nWarm = 2000,

+ nKert = 3500.nThin = 51)
```

## > checkChains(destLongerChains)

vertical slice of log-density	trace	lag 1	acf
function			
1/6 of range	and and planting the state of t		
2/6 of range	Manage Manager		
3/6 of range			
4/6 of range			ellon market en
5/6 of range	kadaladikkalood Uppyrapilipini		***************************************

Figure 6: The chains corresponding to the density estimate object densLongerChains with a warm-up of size 2,000, kept samples of size 3,500 and a thinning factor of 5.

The default number of basis functions for the O'Sullivan penalized splines (e.g. Wand & Ormerod, 2008) is 50. For the 2011 Old Faithful geyser data it is likely that a much smaller basis is sufficient to resolve the structure in the data's density function and that 50 functions is an overkill for this particular dataset. To reduce the number of basis functions to 20 type:

```
> destSmallerBasis <- densEstBayes(OldFaithful2011,control =
+ densEstBayes.control(numBasis = 20))</pre>
```

The following code plots destSmallerBasis and compares it with the default density estimate:

```
> plot(destSmallerBasis,shade = FALSE,estCol = "blue",

+ xlab = "time interval between geyser eruptions (minutes)")
> rug(jitter(OldFaithful2Ol1,amount = 0.2),col = "dodgerblue")
> destDefault <- densEstBayes(OldFaithful2Ol1)
> plot(destDefault,shade = FALSE,estCol = "darkorange",add = TRUE)
> legend("topleft",legend = c("20 spline basis functions",
+ "50 spline basis functions"),lty = rep(1,2),
+ col = c("darkorange","blue"))
```

The full list of control parameters, and the purpose of each one, can be obtained by issuing the command:

> help(densEstBayes.control)

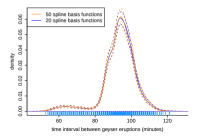


Figure 7: Bayesian density estimates for the 2011 Old Faithful geyser data with the densEst-Bayes() default of 50 basis functions and a simpler one with 20 basis functions specified using the control argument of densEstBayes().

## 4 Convergence Issues

Each of the methods used by densEstBayes() for achieving Bayesian density estimation rely on either an iterative algorithm (method = "SMFVB") or Markov chain Monte Carlo sampling (method = "HMC", method = "NUTS" and method = "Sicc") and, hence, are prone to convergence issues. We now provide some illustrations of such issues.

## 4.1 Fixed point iteration for semiparametric mean field variational Bayes

When method = "SMFVE" is specified in the call to deneEstBayes() then a fixed point iterative scheme is used to obtain the semiparametric mean field variational Bayes approximate posterior distributions of spline basis coefficients. For the default number of spline basis functions, the approximation involves a 52-dimensional Multivariate Normal distribution with a full covariance matrices are determined using the fixed point iterative scheme listed at the bottom of page 29 of Rohde and Wand (2016). Often convergence is quite rapid, and a fast and high quality Bayesian density estimate results. However, noting that the number of free parameters is  $52 + \frac{1}{2} \times 52 \times 53 = 1,430$ , guaranteeing convergence for general input data is a tall order. The following code illustrates this problem, starting with simulation of 100 observations from the 5th density function in Table 1 of Marron and Wand (1992), which is such that outliers tend to present:

```
> set.seed(2) ; x <- rMarronWand(100,5)
> destSMFVB <- densEstBayes(x,method = "SMFVB")</pre>
```

The following error message is returned:

#### Error.

Call to densEstBayes() with method="SMFVB" has failed due to a singular matrix and/or non-convergence problem. Use method="slice", method="NUTS" or method="HMC" instead. If, instead, the default of method = "slice" is used then the following code:

```
> destDefault <- densEstBayes(x); plot(destDefault)
> rug(x,col = "dodgerblue")
> xg <- seq(destDefault$range.x[1],destDefault$range.x[2],length = 1001)
> trueDensg <- dMarronWand(xg,5)
> lines(xg,trueDensg,col = "indianred3")
> legend("topleft",legend = ("estimate", "truth"),lty = rep(1,2),
+ col = c("daxkgreen", "indianred3")
```

produces the estimate shown in Figure 8. The density function from which the data were simulated is shown for comparison

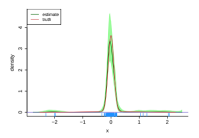


Figure 8: The default densEstBayes() estimate for data simulated from the 5th density function in Table 1 of Marron and Wand (1992) via the function 'xMarronwand'). The true density function is also shown. The shaded region corresponds to pointwise 95% credible intervals.

## 4.2 Markov chain Monte Carlo convergence issues

Consider density estimation for a dataset consisting of the widths of 93 makes of cars. The data are part of the R package MASS (Ripley, 2020) and the following code obtains and prints the sorted data:

```
> library(MASS) ; print(sort(Cars93$Width))
which results in:
```

- A Bayesian density estimate based on a default call to densEstBayes() and plot involves the code:

and leads to the estimate shown in Figure 9 (note that the estimate depends on random draws within a Markov chain Monte Carlo scheme and is subject to change for different calls to densEstBayes(). The density estimate is not very visually appealing and is close to a linear

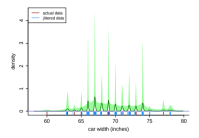


Figure 9: Bayesian density estimate for data on widths (inches) of 93 cars from the data frame Cars 93 in the R package MASS. This estimate is produced from a call to the default version of the function densestBayes. On the rug plots at the base of the plot show both the actual data, which has several clusters of identical observations, and a sittering of the data.

combination of point masses at each of the unique data values. To check the chains on which this estimate is based we issue:

## > checkChains(destCarWidths)

and obtain the graphic shown in Figure 10, which indicates that the Bayesian density estimation procedure is suffering from identifiability problems.

The culprit seems to be the degree to which these data have been discretised. The command:

## > table(Cars93\$Width)

leads to the output:

```
60 63 64 65 66 67 68 69 70 71 72 73 74 75 77 78
1 5 1 3 10 15 7 11 7 5 7 4 11 1 2 3
```

and shows that there are only 16 unique values in a dataset of size 93.

The next set of commands allow for the slice sampling warmup and kept sample sizes to be much longer. A thinning factor of 10 is applied to the kept chains.

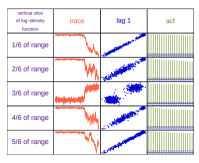


Figure 10: The chains of the log-density estimates at five equally-spaced vertical slices for the slice sampling-based Bayesian density displayed in Figure 9.

The resulting density estimate is shown in Figure 11 and indicates the slice sampling is converging to the linear combination of point masses density estimate. The command:

#### > checkChains(destLongerChains)

produces the Figure 12 graphic. The trace plots are flatter, but still not very satisfactory. The issue seems to be a type of degeneracy that arises in this Bayesian density estimation procedure when data on a continuous variable are discretised.

If we break the ties in the data by adding a small amount of random noise to each of the observations via code such as:

```
> carWidthsJittered <- jitter(Cars93$Width,amount = 0.2)
> destJitteredData <- densEstBayes(carWidthsJittered)
> plot(destJitteredData,xlab = "car width (inches)")
> rug(Cars93$Width,col = "darkred")
> rug(cars93$Width,col = "darkred")
> legend("topleft",legend = c("actual data", "jittered data"),
+ tty=rep(1,2),col=c("darkred", "dodgerblue")
```

then a density estimate such as shown in Figure 13 is shown. This estimate is in keeping with the continuous nature of the car widths variable. A check of the Figure 13 chains via the command:

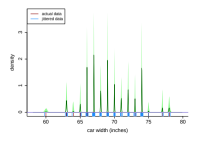


Figure 11: The same as for Figure 11 except that the chains for the slice sampling are specified to have a warmup of length 10,000, kept chains of length 10,000 and a thinning factor of 10.

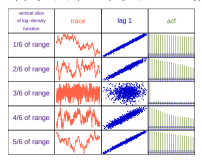


Figure 12: The chains of the log-density estimates at five equally-spaced vertical slices for the slice sampling-based Bayesian density displayed in Figure 12.

## > checkChains(destJitteredData)

is shown in Figure 14 and shows that good Markov chain Monte Carlo behaviour is achieved.

To summarise, density estimation via Monte Carlo-based Bayesian inference engines can be affected by issues such as data discretisation and convergence checks via the checkChains() function are worthwhile.

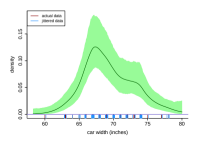


Figure 13: The same as for Figure 11 but with the default call to densEstBayes() applied to the fittered data rather than the actual data.

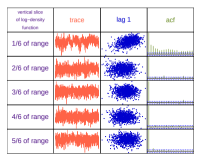


Figure 14: The chains of the log-density estimates at five equally-spaced vertical slices for the slice sampling-based Bayesian density displayed in Figure 13.

# 5 Display Options

After obtaining a Bayesian density estimation object using the daneEstBayes() function there are various options for displaying the estimate. There are tweak factors such as the colour of the estimate curve and whether to use shading or curves for the variability band. Another display aspect is restriction to a sub-region. Some of these display options are explained and illustrated here.

The following commands obtain a density estimate of the 2011 Old Faithful gevser data

and plot the estimate with user-specified colours. The level of the pointwise credible intervals is set to 99%

The result is shown in Figure 15.

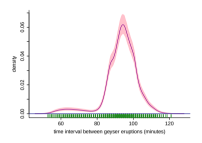


Figure 15: Bayesian density estimate for the 2011 Old Faithful geyser data. The pink shaded region corresponds to approximate 99% pointwise credible intervals. The data are shown as tick marks, with some jittering, at the base of the plot.

Now suppose that we want to zoom in on the region where the time interval between geyser eruptions is between 70 minutes and 100 minutes. Due to the normalization requirement, it is necessary to compute the estimate over the range of the data and then use the predict() function to obtain the plotting vectors:

```
> myKgrid <- seq(TO,100,length = 501)
> predDbj <- predict(dest,newdata = myKgrid,cred.fit = TRUE,credLev = 0.99)
> myEstGrid <- predDbj$fit
> myLowGrid <- predDbj$fredLow.fit
> myUnoGrid <- predDbj$credLow.fit
```

The abscissae plotting vector myXgrid and the ordinate plotting vectors myEstGrid, myLowGrid and myUppGrid can now be used to assemble the required graphic — with dashed curves used for the variability band:

The result is shown in Figure 16.

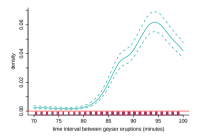


Figure 16: Bayesian density estimate for the 2011 Old Faithful geyser data over the restricted interval of 70 to 100 minutes. The dashed curves correspond to approximate 99% pointwise credible intervals. The data are shown as tick marks, with some jittering, at the base of the plot.

Finally, we provide an example involving two Bayesian density estimates plotted on the same axes. The data are part of the data frame <u>carAuction</u> within the R package HBW (Harczlak, Ruppert and Wand, 2019), which contains observations for 51 variables on 72,983 cars purchased at automobile auctions by dealerships. Here we study the variable carAuction5odomRead, which is the odometer redaing, in miles, of the car at the time of purchase and divide the data according to whether or not the car is a considered a bad buy or good buy by the dealership. The following code creates the required data vectors.

```
> library(HRW)
```

- > odomReadGoodBuv <- carAuction\$odomRead[carAuction\$IsBadBuv == 0]
- > odomReadBadBuv <- carAuction\$odomRead[carAuction\$IsBadBuv == 1]

Bayesian density estimates for each data are obtain via:

```
> destGoodBuy <- densEstBayes(odomReadGoodBuy,method = "SMFVB")
> destBadBuy <- densEstBayes(odomReadBadBuy,method = "SMFVB")</pre>
```

Plotting vectors to display the two density estimates on the same set of axes are computed using:

```
> myOdomReadGrid <- seq(20000,120000,length = 1001)
> predObjGoodBuy <- predict(destGoodBuy,newdata = myOdomReadGrid, + cred.fit = TRUE)
> estGridGoodBuy <- predObjGoodBuy$fit
```

```
> estGridBadBuv <- predObiBadBuv$fit
> lowGridRadRuy <- predObiRadRuy$credLow fit
> unnGridRadRuv <- nredOhiRadRuv$credUnn fit
We are now ready to make the desired plot:
> vlimVal <- range(c(lowGridGoodRuv unnGridGoodRuv
                     lowGridRadRuy unnGridRadRuy))
> par(mfrow = c(1 1))
> mvGoodColour <- "blue" : mvBadColour <- "darkorange"
> plot(0.type = "n".btv = "l".xlim = range(mv0domReadGrid).vlim = vlimVal.
       xlab = "odometer reading (miles)".vlab = "density")
> abline(h = 0.col = "slateblue")
> lines(mvOdomReadGrid.estGridGoodBuv.col = mvGoodColour.lwd = 3)
> lines(mvOdomReadGrid lowGridGoodRuy col = mvGoodColour lyd = 3 lty = 2)
> lines(mvOdomReadGrid.uppGridGoodBuv.col = mvGoodColour.lwd = 3.1tv = 2)
> lines(mvOdomReadGrid.estGridBadBuv.col = mvBadColour.lwd = 3)
> lines(mvOdomReadGrid.lowGridBadBuv.col = mvBadColour.lwd = 3.1tv = 2)
> lines(mvOdomReadGrid.uppGridBadBuv.col = mvBadColour.lwd = 3.1tv = 2)
> legend("topleft".legend = c("good buy at auction"."bad buy at auction").
         ltv = rep(1.2) \cdot lwd = rep(3.2) \cdot col = c(mvGoodColour.mvBadColour))
```

cred fit = TRUE)

> lowGridGoodBuy <- predUbjGoodBuy\$credLow.fit > uppGridGoodBuy <- predUbjGoodBuy\$credUpp.fit

> predObiBadBuy <- predict(destBadBuy.newdata = myOdomReadGrid.

The result is shown in Figure 17 and indicates that the density function of odometer reading values for bad buy cars is shifted to the right compared to that for good buy cars.

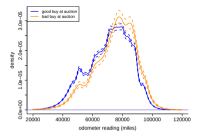


Figure 17: Bayesian estimates of the density functions of odometer readings of 72,983 cars sold at auctions by automobile dealerships in U.S.A. The data have been divided according to whether or not the car is classified as a good buy (64,007 cars) or a bad buy (8,976 cars). The dashed curves correspond to pointwise 95% credible intervals.

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