MCMC-Based Bayesian Change Point Detection: a Comparison between the Gibbs and Metropolis-within-Gibbs Samplers

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1 Introduction

Change point detection solves the class of problems where one would like to identify times when abrupt changes in the underlying data generating process of a time series occurs. In practice, this setting is widely seen in areas such as stock analysis and climate studies [CG97; ENJ04]. In the literature, a wide variety of approaches, both frequentist and Bayesian, have been developed to efficiently solve the change point problems [KFE12; CGS92]. In the Bayesian approaches, the problem is often framed as estimating the posterior distribution of the change point locations. The common advantages of the Bayesian approach to statistical inference apply in the change point setting. Compared to the point estimates of the change point locations, the posterior distribution of the change point locations of a given sequence provides a more flexible and holistic understanding of the time series. For example, having access to the approximate posterior distribution of the change point locations allows us to estimate the probability that a change has occured at some given time.

One of the earliest Bayesian approaches to the change point problem is developed in [CGS92] through Markov Chain Monte Carlo (MCMC). Particularly, a Gibbs sampler is used to obtain samples for estimating the posterior distribution of the change point locations. It is worth noting that, although only the detection of a single change point is discussed in [CGS92], there is a natural extension to the multiple change point setting when the number of change points is known. Specifically, instead of sampling a single change point from the distribution of change points conditioned on all other parameters in the model, a set of change points can be sampled from the conditional distribution over all possible configurations of the change point locations. It is then obvious that the number of possible configurations of the change point locations grows linearly in the length of the time series and exponentially in the number of change points to be detected. As a result, the computational complexity of the Gibbs sampler becomes exponentially more expensive.

The scalability issue described above is present in virtually all change point detection methods. In fact, many of the more recent developments in MCMC-based approaches can be viewed as trying to bypass this computational bottleneck either through clever reformulation of the problem [Ste94; LL01] or the use of Metropolis-Hastings (MH) and potentially well-designed proposal distributions [Gre95; AL08]. The approach in [AL08] is one of the easiest and most intuitive attempts at circumventing the computational challenge posed in [CGS92]. Instead of computing the full conditional distribution of the change point locations, a uniform proposal over all possible configurations of change point locations is accepted/rejected based on the MH ratio. However, while the expensive full conditional distribution no longer needs to be evaluated, exploring this potentially enormous space of change point locations using a uniform proposal may not be the most efficient, thus posing a different challenge.

In this report, we compare the Bayesian change point detection problems proposed in [CGS92] and [AL08]. Using a piecewise constant model, we would like to explore, under different settings, whether there is a tradeoff between evaluating the expensive full conditional distribution and searching over a potentially large space
using a cheaper sampler that risks low acceptance rates. Through a number of experiments, we find that
although the approach in [CGS92] is more favourable in all models that are tested, the approach in [AL08]
offers very competitive performance. However, we note that only a uniform proposal distribution is used.

By designing more informative proposal distribution, as already been explored in [BF+18], proposal-based MCMC methods applied to the change point problems would be a very attractive choice when the number of possible change point configurations is too large to be handled by a Gibbs sampler.

2 Problem Setup

Below, we start by specifying the piecewise constant model and outlining the procedures of the two MCMC methods before presenting the discussing the experimental results.

2.1 Pairwise Constant Model

Consider a sequence of random variables (Y_1, \dots, Y_n) , where $\forall i \in \{1, \dots, n\}, Y_i \in \mathbb{R}$, such that

$$Y_i \sim \begin{cases} N(\mu_1, 1) & 1 \le i \le r_1 \\ N(\mu_2, 1) & r_1 < i \le r_2 \\ \vdots & & \\ N(\mu_{k+1}, 1) & r_k < i \le n \end{cases},$$

where k is known, $\mathbf{r} = (r_1, \dots, r_k)$ denotes the set of integer valued change point locations, and $\boldsymbol{\mu} = (\mu_1, \dots, \mu_{k+1})$ denotes the underlying means of each segment. Note that the jth segment is defined to be $\mathbf{Y}_{r_{j-1}+1:r_j} = (Y_{r_{j-1}+1}, \dots, Y_{r_j})$, with $r_0 = 0, r_{k+1} = n$ by convention. In addition, we assume that \mathbf{r} and $\boldsymbol{\mu}$ are independent and that the random variables are i.i.d. within each segment.

Let $f(\mu_i; y_j)$ be the likelihood of μ_i being the mean of a normal distribution with variance 1 given the observation y_j . Then given a sequence of realized observations $\mathbf{y} = (y_1, \dots, y_n)$ from the above data generating process, its likelihood is defined by

$$L(\mathbf{r}, \boldsymbol{\mu} \mid \mathbf{y}) = \prod_{i=1}^{k+1} \prod_{j=r_{i-1}+1}^{r_i} f(\mu_i; y_j).$$

Then once a set of prior distributions are defined on the parameters \mathbf{r} and $\boldsymbol{\mu}$, the goal of the Bayesian change point detection methods is to estimate the posterior distributions $p(\mathbf{r} \mid \mathbf{y})$ and $p(\boldsymbol{\mu} \mid \mathbf{y})$. We focus on the posterior distribution of the change point locations $p(\mathbf{r} \mid \mathbf{y})$.

For simplicity, we assume the priors on the segment means to be

$$\mu_i \sim N(m, 1), \forall i = 1, \dots, k+1, \text{ and } m \in \mathbb{R}.$$
 (1)

As a prior on the change point locations, we assume that r follows a discrete uniform distribution over all possible configurations of the change point locations. Specifically, for any given possible configuration r_c ,

$$p_{\boldsymbol{r}}(\boldsymbol{r}_c) = \frac{1}{\binom{n-1}{k}}.$$

Note that by our specification, it is not possible for the last observation in the sequence to be a change point.

2.2 MCMC Simulation Procedures

Under the above specification, to obtain a set of samples for r and μ , given (r^i, μ^i) , the Gibbs sampler described in [CGS92] generates the next sample (r^{i+1}, μ^{i+1}) through the following steps:

1. generate r^{i+1} from the probability mass function

$$p\left(\boldsymbol{r}\mid\boldsymbol{y},\boldsymbol{\mu^{i}}\right) = \frac{L\left(\boldsymbol{r},\boldsymbol{\mu^{i}}\mid\boldsymbol{y}\right)}{\sum_{j=1}^{\binom{n-1}{k}}L\left(\boldsymbol{r}^{j},\boldsymbol{\mu^{i}}\mid\boldsymbol{y}\right)},$$

where r^{j} in the denominator denotes the jth configuration from the permutation of all possible configurations;

2. generate μ^{i+1} by

$$\mu_j^{i+1} \sim N\left(\frac{m + \sum_{l=r_{j-1}+1}^{r_j^{i+1}} y_l}{r_j^{i+1} - r_{j-1}^{i+1}}, (1 + r_j - r_{j-1})^{-1}\right), \forall j \in \{1, \dots, k+1\}.$$
 (2)

Similarly, given $(\mathbf{r}^i, \boldsymbol{\mu}^i)$, the Metropolis-within-Gibbs (MWG) sampler described in [AL08] generates the next sample $(\mathbf{r}^{i+1}, \boldsymbol{\mu}^{i+1})$ through the following steps:

- 1. generate r' from the discrete uniform prior p_r ;
- 2. accept \mathbf{r}' as \mathbf{r}^{i+1} with probability min $(1, \beta(\boldsymbol{\mu}^i, \mathbf{r}^i, \mathbf{r}'))$, where

$$eta\left(oldsymbol{\mu}^{i},oldsymbol{r}^{i},oldsymbol{r}'
ight)=rac{L\left(oldsymbol{r}',oldsymbol{\mu}^{i}\midoldsymbol{y}
ight)}{L\left(oldsymbol{r}',oldsymbol{\mu}^{i}\midoldsymbol{y}
ight)};$$

- 3. otherwise set $\mathbf{r}^{i+1} = \mathbf{r}^i$;
- 4. generate μ^{i+1} following the last step of the Gibbs sampler.

Despite the use of conjugate priors when sampling the means of each segment, the trade-off between evaluating the expensive full conditional distribution of the change point locations and searching over a potentially large space using a cheaper sampler that risks low acceptance rates is still present. Specifically, the cost of computing $p\left(r\mid y,\mu^i\right)$ from the Gibbs sampler grows linearly in the length of the time series and exponentially in the number of change points to be detected. The search space size of all configurations of change point locations from the MWG sampler also grows in the same manner.

Before presenting the experiment results, it is worth noting that the posterior distribution

$$p(\boldsymbol{r} \mid \boldsymbol{y}) = \int_{\mu_1} \cdots \int_{\mu_{k+1}} p(\boldsymbol{y} \mid \boldsymbol{\mu}, \boldsymbol{r}) p_{\boldsymbol{r}}(\boldsymbol{r}) p(\mu_1) \cdots p(\mu_{k+1}) d\mu_1 \cdots d\mu_{k+1}$$

$$= p_{\boldsymbol{r}}(\boldsymbol{r}) \left(\int_{\mu_1} p(\boldsymbol{y}_{1:r_1} \mid \mu_1) p(\mu_1) d\mu_1 \right) \cdots \left(\int_{\mu_{k+1}} p(\boldsymbol{y}_{r_k+1:n} \mid \mu_{k+1}) p(\mu_{k+1}) d\mu_{k+1} \right)$$

$$\coloneqq p_{\boldsymbol{r}}(\boldsymbol{r}) Q(1, r_1, \mu_1) \cdots Q(r_k + 1, n, \mu_{k+1}).$$

can be fully evaluated. Note that $p(\mu_i)$ denotes the prior distribution of the *i*th segment mean, and the likelihood of μ_i given all observations from the *i*th segment of the sequence is denoted

$$p(\mathbf{y}_{r_{i-1}+1:r_i} \mid \mu_i) = \prod_{j=r_{i-1}+1}^{r_i} f(\mu_i; y_j).$$

This posterior distribution can be evaluated using Baye's rule because the $Q(\cdot)$'s are the normalizing constants of the known posterior distributions of the segment means defined in Eq. (2). This allows us to more directly compare the quality of the samples generated from the two MCMC methods in the next section.

3 Experiment

As mentioned in the previous sections, both the length of the entire sequence and the number of change points to be detected have an impact on the number of possible configurations of change point locations and thus the computational cost of evaluating the conditional distribution of the change point locations. Therefore, to evaluate the performance of both MCMC-based Bayesian change point detection methods,

3.1 Discussion 3 EXPERIMENT

inference on five sequences of varying lengths and number of true change points are conducted using both the Gibbs and MWG approaches. The number of possible change point configurations range from 49 to 32509. The detailed specification and visualization of the five sequences are shown in Appendix A.

In this set of experiments, the *m* value in Eq. (1) is set to be the average of the true segment means of each sequence specified in Appendix A. Then inference on the change point locations of each sequence is run five times using both methods to ensure the results that we obtain are not by chance. For each of these runs, the same initial values are used for both the Gibbs sampler and the MWG sampler, although these initial values are not stored in the final output. Specifically, the initial change point locations are uniformly sampled and the initial segment means are the empirical means of the segments based on the initial change points. The Markov chains are run for one minute on the first and smallest time series, and for three minutes on all of the other four time series. Note that the relative time is measured using time.perf_counter provided in Python with the time spent on storing each output sample excluded. It is worth noting that the numba package is used for both methods to speed up computations by translating and pre-compiling some of the numpy functions. The experiments are run with an Intel i5 7200U processor and 8GB of memory. Code is available at https://github.com/NaitongChen/STAT520A_Project.

3.1 Discussion

Firstly, by checking the plots shown in Appendix B, we confirm that the number of samples produced by MWG is indeed orders of magnitude larger than that of Gibbs except for the first sequence. However, this is justified by that evaluating the conditional distribution of the change point locations is relatively cheap as there are only 49 possible locations of the change point.

To further study the actual values of the samples produced, we take a look at the trace plots of each change point location. Appendix C lists the trace plots of each change point, grouped by the particular sequences, across all five runs using both methods. After visually assessing these plots, the samples generated before the Makov chains have converges are removed, and the trace plots after removing the burin-in periods are shown in Appendix D. It is worth noting that this visual assessment is possible because we have knowledge of the ground truth for each sequence. For example, looking at Appendix C without knowing the true change point locations may lead to the conclusion that the corresponding Markov chain has converged since the beginning of the simulation.

From these trace plots, we see that the Markov chains in both methods converge almost immediately under the first two sequences. However, for the latter three sequences, MWG more often than not requires more time than Gibbs before reaching the high density regions. This is particularly evident for the sequence with 60 observations and 3 change points because the shorter segments make the segment means less stable. Even after the corresponding chains have converged, it is clear under the models with a larger search space of change point locations, MWG moves between states much less freequently than Gibbs does.

To better understand the behaviour of the two approaches, we take a closer look at the last round of simulation on the sequence with 60 observations and 3 change points (Figs. 7e and 12e). Using the same initialization, initially both MWG and Gibbs completely miss the last change point in the sequence. Indeed, putting a change point at location 30 corresponds to a local mode in the posterior distribution of the change point locations, regardless of whether it is the corresponding change point that is given this value. As a result, with a uniform proposal on the change point locations, MWG takes a very long time to generate a sample of higher mass from the more than 30000 possible configurations. Even after the Markov chain of MWG has converged to the region around the global mode, a similar problem is present. Since any set of change points around the global mode has a relatively high mass already, again with the uniform proposal, MWG fails at efficiently exploring this space as the probability of proposing another set of change point locations of comparable mass is low. This is clearly reflected in Fig. 12e.

In comparison, the Gibbs sampler gets out of the local mode much faster than MWG does. And even after the Gibbs sampler has reached the region of the global mode of the posterior, it seems to explore

this high mass region a lot more thoroughly. This is because the Gibbs sampler utilizes the full conditional distribution of the change point locations conditioning on all other parameters (segment means). In this particular case, despite missing the last change point completely, it very quickly identifies the first and third change point using the more information conditional distribution of the change point locations. Once two of the three change points have been identified, the segment means are immediately updated to be much closer to the true segment means, which then informs the Gibbs sampler to ultimately get out of the local mode.

From above, it seems like the more informative conditional distribution of the change point locations puts the Gibbs sampler at an advantage. However, we remind ourselves that the trace plots are compared iteration wise. In other words, even though MWG requires more samples to reach the region around the global mode, samples from MWG are generated much faster than those from Gibbs. We now check whether this faster sampling rate from MWG makes up for the difference in performance mentioned above.

As mentioned in Section 2, the true posterior distributions are available for each of the five sequences. We then use the plots in Appendix E to check how fast the estimated posteriors from both methods approach the true distribution. This distance is measured using the Wasserstein distance. We see that, except for the sequence with 20000 observations and 1 change point, the approximate posteriors computed using the Gibbs sampler is uniformly closer to those of MWG. This is because the Gibbs sampler explores the high mass region of the posterior more thoroughly with the help of the conditional distribution of the change point locations. In the case of Fig. 13b, the absence of the difference as shown in the other figures maybe be due to the fact that the sequence is much longer. Then the distances between the approximate and the true posteriors are averaged off by the vast majority of the change point locations with negligible mass.

To confirm our speculation above, we check the approximate and true marginal posterior distributions of each change point (Appendix F). A general trend in these plots is that the approximate marginal posterior distributions from the Gibbs sampler better capture the mode of each marginal distribution than those from MWG. Again this is because the Gibbs sampler explores the region of high mass better than MWG.

However, it is worth pointing out it is not the case that MWG does not have any advantages over the Gibbs sampler. To see this, we plot the effective sample sizes (ESS) and Monte Carlo standard errors (MCSE) of each sequence studied (shown in Appendix G). As suggested by [FJ+10], the ESS and MCSE values are computed using the overlapping batch mean method after splitting the number of observations $n = a_n b_n = \left(n^{\frac{1}{3}}\right) \left(n^{\frac{2}{3}}\right)$. We see that in the sequences with 20000 observations and 1 change point and with 100 observations and 3 change points, samples from MWG tends to have higher ESS's and lower MCSE's. Both of these sequences have a large search space of change point locations and do not exploit MWG's slow mixing rates. Under such circumstances, if the ground truth is unknown, samples from MWG may shed some more insights than those from Gibbs.

4 Conclusion

Through the experiments above, we find that the approach in [CGS92] is more favourable in all models that we have tested. Then when the computational cost of evaluating the conditional distribution of the change point locations is not prohibitive, using the Gibbs sampler approach will produce samples that better represent the posterior distributions of the change point locations. However, we note that the approach in [AL08] offers very competitive performance, especially considering only a uniform proposal distribution is used. By designing more informative proposal distribution, as already been explored in [BF+18], proposal-based MCMC methods applied to the change point problems would be a very attractive choice when the computational load is too heavy to be handled by the fully Gibbs approach.

A Description and Visualization of Datasets

See below for the visualization of the five sequences whose change points are to be detected using MCMC-based Bayesian change point detection methods. The blue dots are the observed values and the orange lines show the underlying means of each segment. The number of observation (n), number of change points (k), true segment means (μ) , true change point locations (r), and the number of possible configurations of change points (T) are shown in the plots below. Note that the x-axis indicate the index or order in time, and the y-axis are the values of the observations.

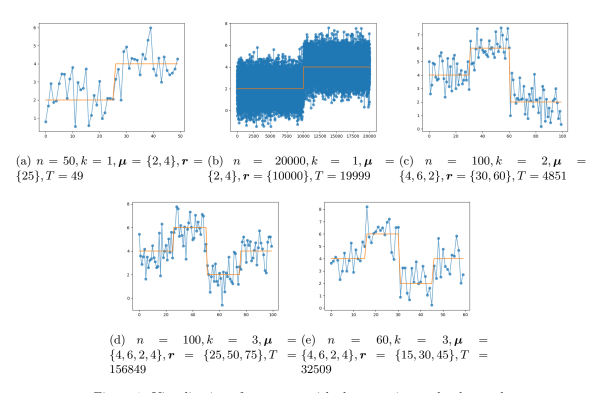


Figure 1: Visualization of sequences with change points to be detected.

B Sample Size Comparison

See below for a comparison of the size of the samples generated by each MCMC sampler for each of the five sequences. For each model, the medians (solid lines), the 25th and the 75th percentiles (dashed lines) of the number of samples generated by each MCMC sampler are plotted across the amount of time elapsed since the start of the simulation. Note that in this particular plot, for each sequence, the samples of each MCMC sampler is trimmed to the minimum sample sizes across all five runs.

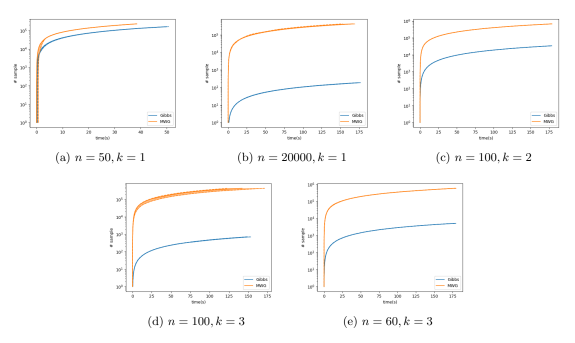


Figure 2: Number of samples generated by each MCMC sampler for each of the five sequences.

C Trace plots of entire sequences

Each of the five figures below show the trace plots of the change points across the five runs for each sequence. Note that the trace plots include all samples generated by both MCMC samplers. Each color represents a distinct change point. For each plot, the x-axis indicate the number of the sample, and the y-axis represent the values of each sample (locations of change points).

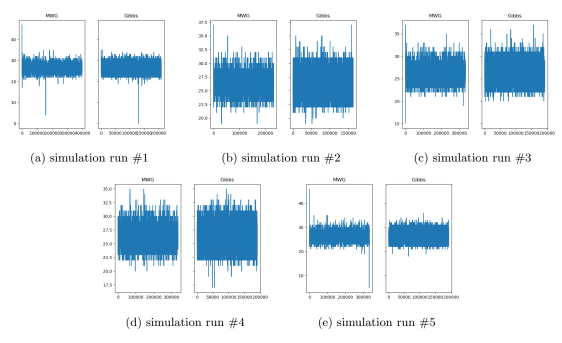


Figure 3: Trace plots of entire simulations for the sequence with $n = 50, k = 1, \mu = \{2, 4\}, r = \{25\}, T = 49$.

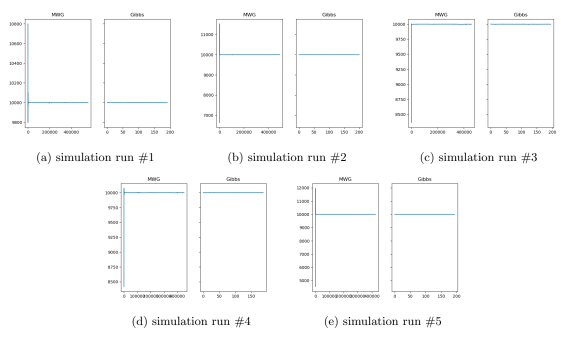


Figure 4: Trace plots of entire simulations for the sequence with $n=20000, k=1, \mu=\{2,4\}, r=\{10000\}, T=19999$.

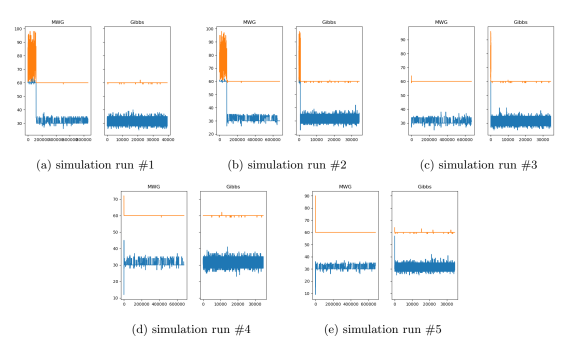


Figure 5: Trace plots of entire simulations for the sequence with $n=100, k=2, \boldsymbol{\mu}=\{4,6,2\}, \boldsymbol{r}=\{30,60\}, T=4851.$

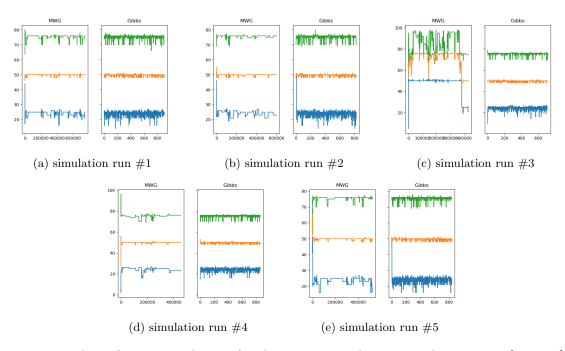


Figure 6: Trace plots of entire simulations for the sequence with $n=100, k=3, \boldsymbol{\mu}=\{4,6,2,4\}, \boldsymbol{r}=\{25,50,75\}, T=156849.$

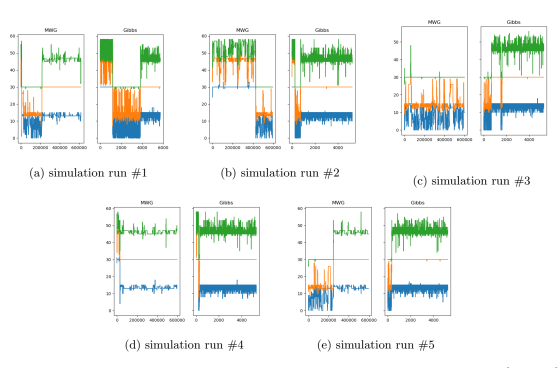


Figure 7: Trace plots of entire simulations for the sequence with $n=60, k=3, \boldsymbol{\mu}=\{4,6,2,4\}, \boldsymbol{r}=\{15,30,45\}, T=32509.$

D Trace plots after burn in

Each of the five figures below show the trace plots of the change points across the five runs for each sequence. Note that the trace plots exlude samples generated before the corresponding Markov chains have converged. Each color represents a distinct change point. For each plot, the x-axis indicate the number of the sample, and the y-axis represent the values of each sample (locations of change points). Empty plots indicate that the simulation has terminated before the Markov chain converged.

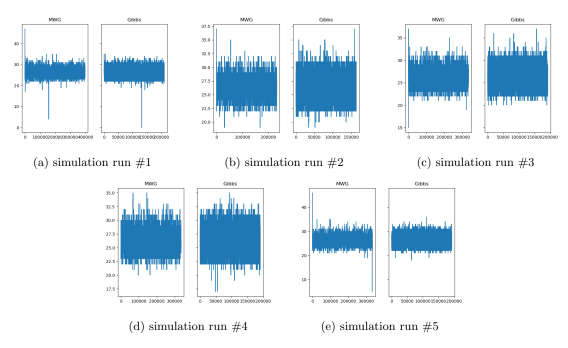


Figure 8: Trace plots post burn-in for the sequence with $n = 50, k = 1, \mu = \{2, 4\}, r = \{25\}, T = 49$.

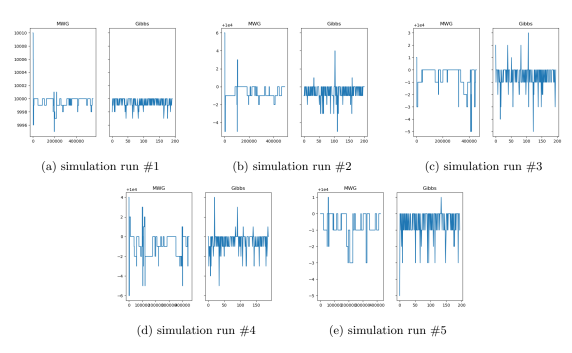


Figure 9: Trace plots post burn-in for the sequence with $n=20000, k=1, \boldsymbol{\mu}=\{2,4\}, \boldsymbol{r}=\{10000\}, T=19999.$

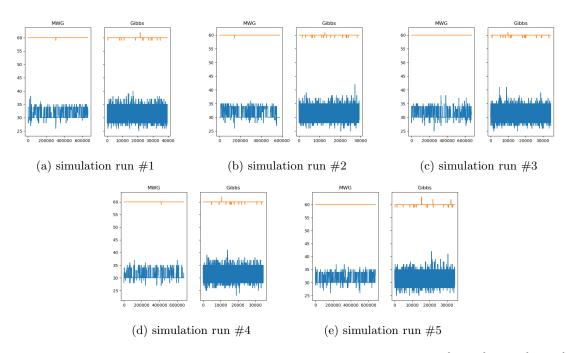


Figure 10: Trace plots post burn-in for the sequence with $n=100, k=2, \boldsymbol{\mu}=\{4,6,2\}, \boldsymbol{r}=\{30,60\}, T=4851.$

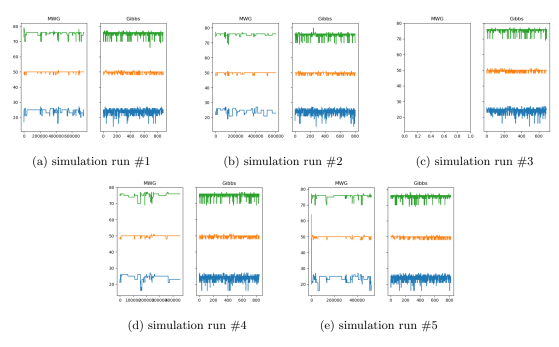


Figure 11: Trace plots post burn-in for the sequence with $n=100, k=3, \mu=\{4,6,2,4\}, r=\{25,50,75\}, T=156849.$

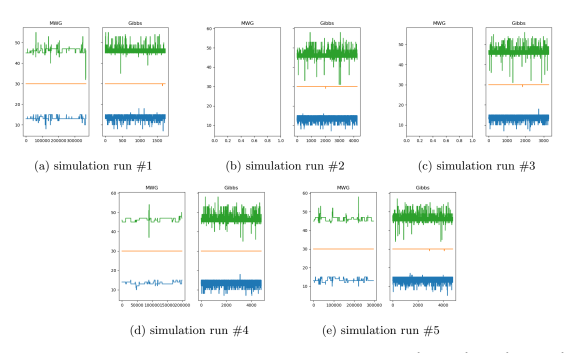


Figure 12: Trace plots post burn-in for the sequence with $n = 60, k = 3, \mu = \{4, 6, 2, 4\}, r = \{15, 30, 45\}, T = 32509.$

E Wasserstein Distance to the True Posterior

See below for the comparison between the two MCMC samplers on the Wasserstein distance between the approximated posterior and the true posterior. Approximate posteriors are the empirical distributions of the change point locations using all samples available post burn-in after a certain amount of time has passed since the start of the simulation (shown on the x-axis of the plots). Note these plots do not include simulation runs where the corresponding Markov chains have not converged by the end of the simulation period.

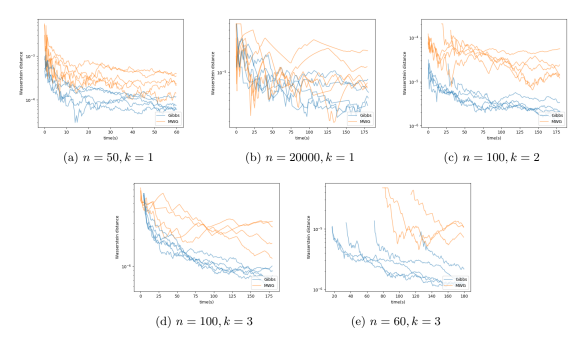


Figure 13: Wasserstein distance between the estimated posterior distribution and the true distribution.

F Marginal Posterior Approximation

See below for a comparison between each of the MCMC sampler's marginal posterior distribution and the true posterior distribution. Note that the green lines denoted BF indicate the true marginal posterior distributions computed using brute force. The approximate marginal posteriors are the empirical distributions using all samples obtained after brun-in and before the end of the simulations. If a simulation fails to converge, the marginal posteriors are not computed. Note that the x-axis indicate the index of the sequence and the y-axis show the associated mass. A plot is made for each change point. Note also that since the majority of the mass is negligible, for the sequence with 20000 observations, only the area around the global mode is shown.

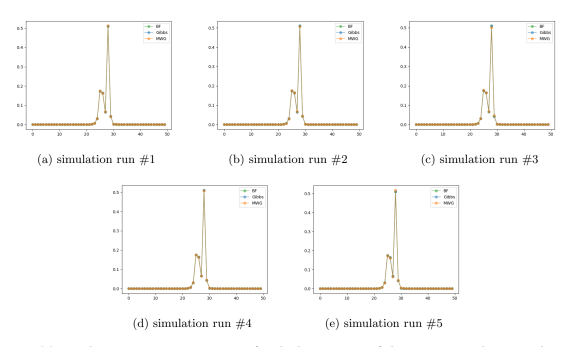


Figure 14: Marginal posterior approximations of each change point of the sequence with $n=50, k=1, \mu=\{2,4\}, r=\{25\}, T=49$.

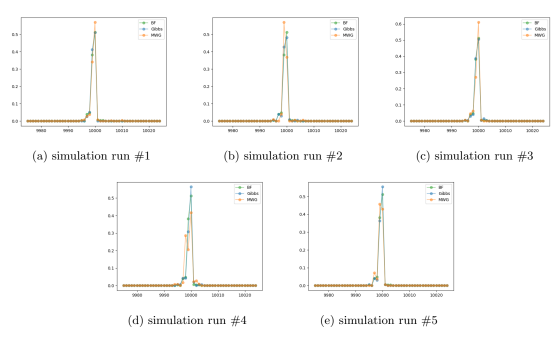


Figure 15: Marginal posterior approximations of each change point of the sequence with $n=20000, k=1, \mu=\{2,4\}, r=\{10000\}, T=19999.$

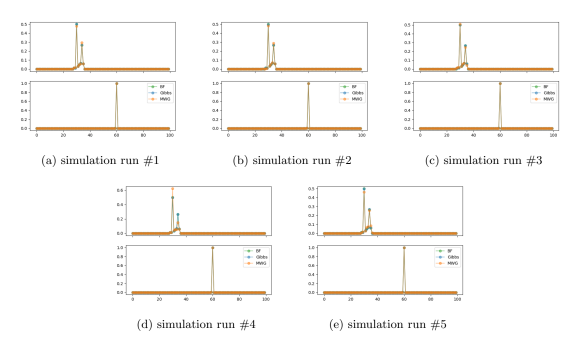


Figure 16: Marginal posterior approximations of each change point of the sequence with $n=100, k=2, \mu=\{4,6,2\}, r=\{30,60\}, T=4851.$

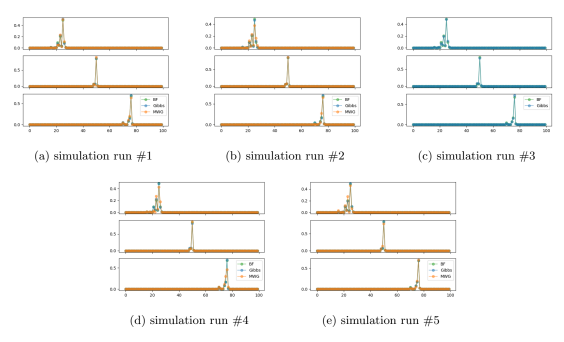


Figure 17: Marginal posterior approximations of each change point of the sequence with $n=100, k=3, \mu=\{4,6,2,4\}, r=\{25,50,75\}, T=156849.$

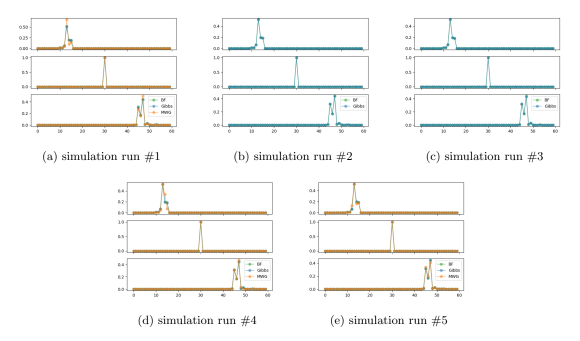


Figure 18: Marginal posterior approximations of each change point of the sequence with $n = 60, k = 3, \mu = \{4, 6, 2, 4\}, r = \{15, 30, 45\}, T = 32509.$

G Effective Sample Sizes and Monte Carlo Standard Errors

See below for the effective sample sizes (ESS) and Monte Carlo standard errors (MCSE) across all simulations for each of the sequenced using both samplers. The x-axes can be ignored and the y-axes show the values of interest as indicated in the plot of each subfigure. These values are computed using all samples obtained after brun-in and before the end of the simulations. If a simulation fails to converge, the ESS and MCSE values are not computed.

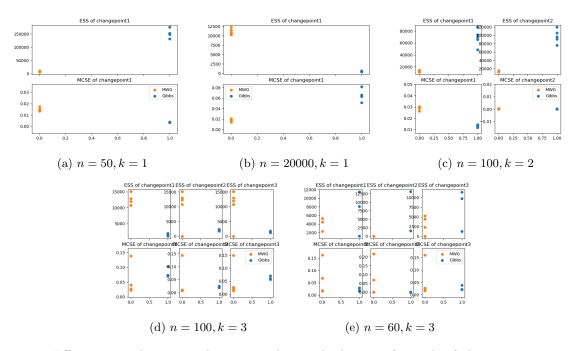


Figure 19: Effective sample sizes and monte carlo standard errors for each of the sequences across all simulations.

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