University of Toronto Department of Economics



Working Paper 618

Hamiltonian Sequential Monte Carlo with Application to Consumer Choice Behavior

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September 12, 2018

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September 7, 2018

Abstract

Practical use of nonparametric Bayesian methods requires the availability of efficient algorithms for implementation for posterior inference. The inherently serial nature of Markov Chain Monte Carlo (MCMC) imposes limitations on its efficiency and scalability. In recent years there has been a surge of research activity devoted to developing alternative implementation methods that target parallel computing environments. Sequential Monte Carlo (SMC), also known as a particle filter, has been gaining popularity due to its desirable properties. SMC uses a genetic mutation-selection sampling approach with a set of particles representing the posterior distribution of a stochastic process. We propose to enhance the performance of SMC by utilizing Hamiltonian transition dynamics in the particle transition phase, in place of random walk used in the previous literature. We call the resulting procedure Hamiltonian Sequential Monte Carlo (HSMC). Hamiltonian transition dynamics in the context of MCMC procedures. The rationale behind HSMC is to translate such gains to the SMC environment. We apply both SMC and HSMC to a panel discrete choice model with a nonparametric distribution of unobserved individual heterogeneity. We contrast both methods in terms of convergence properties and show the favorable performance of HSMC.

JEL: C11, C14, C15, C23, C25

Keywords: Particle filtering, Bayesian nonparametrics, mixed panel logit, discrete choice

^{*}We would like to thank the participants of the 11th International Conference on Computational and Financial Econometrics (CFE), University of London, UK, 2017, for insightful comments and suggesions. Computations were performed on the Niagara supercomputer at the SciNet HPC Consortium. SciNet is funded by: the Canada Foundation for Innovation under the auspices of Compute Canada; the Government of Ontario; Ontario Research Fund - Research Excellence; and the University of Toronto.

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1 Introduction

In Bayesian statistics, parameters are treated as random variables and all forms of uncertainty are expressed in terms of probability. A nonparametric Bayesian model is a model whose parameter space has infinite dimensionality. For any given finite data set, only a finite subset of the available parameters is invoked whereby its dimensionality is allowed to grow with the sample size (Orbanz and Teh, 2010). Nonparametric (and semiparametric) models allow us to avoid the arbitrary and possibly unverifiable assumptions inherent in parametric models. A recent detailed exposition of Bayesian nonparametric methods is provided in Ghosal and van der Vaart (2017).

A critical issue for the practical use of nonparametric Bayesian models is the availability of efficient algorithms to implement posterior inference. The last several decades have witnessed an explosive growth of numerical implementation methods in Bayesian analysis. The cornerstone of such methods has been Markov Chain Monte Carlo (MCMC) and its variants. Nonetheless, the inherently *serial* nature of MCMC, whereby a new draw of the desired parameter chain can only be taken conditional on completing the preceding draw, imposes limitations on the implementational efficiency and scalability of such methods. Yet, the speed of microprocessor cores measured by their GHz frequency has been virtually stable since the mid-2000s, following decades of rapid growth (Rupp, 2018). During the last ten years or so, improvements in computing performance have not originated from processor speed but rather from *parallelization*¹.

In recent years there has been a surge of research activity devoted to developing alternative implementation methods that target (massively) parallel computing environments. In this paper we focus on one particular stream of research in this area: Sequential Monte Carlo (SMC), also known as a particle filter (Doucet et al., 2001). SMC uses a genetic mutation-selection sampling approach with a set of particles representing the posterior distribution of a stochastic process. SMC is highly parallelizable as the core computational load involving

¹Recent trends include shared memory multi-core CPUs, GPUs, and distributed memory highperformance clusters.

the model likelihood is performed by individual particles independently of one another. Due to their computational complexity, Bayesian nonparametric methods stand to benefit substantially from such approaches.

SMC algorithms were initially developed to solve filtering problems that arise in nonlinear state space models (Doucet et al., 2001). In economics, the SMC approach has become a popular method of inference for dynamic systems that benefit from real-time updating of the posterior approximation via recursive importance sampling updates (Kim et al., 1998; Fernández-Villaverde and Rubio-Ramírez, 2007; Creal, 2012; Lopes and Carvalho, 2013; Herbst and Schorfheide, 2014; Blevins, 2016). Chopin (2002) adapted SMC to conduct posterior inference for a static Euclidean parameter vector. This approach was further extended by Fearnhead (2004), Ulker et al. (2010), Carvalho et al. (2010), Bouchard-Côté et al. (2017), and Griffin (2017) to static nonparametric mixture models, which is also our modeling context. The extent to which SMC is parallelizable in a related parametric environment and the corresponding computational gains are elaborated in Durham and Geweke (2014).

SMC is typically implemented in three phases: (i) particle reweighting (correction phase), (ii) particle resampling (selection phase), and (iii) particle transition (mutation phase). We detail each phase further below. In this paper, we propose to enhance the performance of SMC by utilizing Hamiltonian transition dynamics in the particle mutation phase, in place of random walk transitions used in SMC in the previous literature. We call the resulting procedure Hamiltonian Sequential Monte Carlo (HSMC). Hamiltonian transition dynamics have been shown to yield superior mixing and convergence properties relative to random walk transition dynamics in the context of serial MCMC procedures (Neal, 2011). In particular, Hamiltonian dynamics use information about the first derivative of the likelihood function and construct a proposal draw using a sequence of steps, unlike random walk (RW) one-step proposals that do not use derivative information. The rationale behind HSMC is to extend such gains to the SMC environment. We apply both SMC and HSMC to a panel discrete choice model with a nonparametric distribution of unobserved individual heterogeneity. We contrast both methods in terms of convergence properties and show the favorable performance of HSMC.

The remainder of this paper is organized as follows. In section 2 we provide the background for MCMC methods and in section 3 a review of SMC. Hamiltonian transition dynamics are detailed in section 4. We discuss Bayesian nonparametrics in section 5. Within the context of a Bayesian nonparametric mixture model, we introduce HSMC combining SMC with Hamiltonian dynamics in section 6. We then apply both SMC and HSMC to a nonparametric discrete choice model in section 7, comparing the performance of both approaches. Section 8 concludes.

2 Markov Chain Monte Carlo

Consider a general class of models that is parametrized by a Euclidean vector $\theta \in \Theta$ with posterior density $\pi(\theta)$ assumed known up to θ and an integrating constant². Formally, this class of models can be characterized by a family \mathcal{P}_{θ} of probability measures on a measurable space (Θ, \mathcal{B}) where \mathcal{B} is the Borel σ -algebra. The purpose of Markov Chain Monte Carlo (MCMC) methods is to formulate a Markov chain on the parameter space Θ for which, under certain conditions, $\pi(\theta) \in \mathcal{P}_{\theta}$ is the invariant (also called "equilibrium") distribution. The Markov chain of draws of θ can be used to construct simulation-based estimates of the required integrals and functionals $h(\theta)$ of θ that are expressed as integrals. These functionals include objects of interest for inference on θ such as quantiles of $\pi(\theta)$.

The Markov chain sampling mechanism specifies a method for generating a sequence of random variables $\{\theta_r\}_{r=1}^R$, starting from an initial point θ_0 , in the form of conditional distributions for the draws $\theta_{r+1}|\theta_r \sim Q(\theta_r)$. Under relatively weak regularity conditions (Robert and Casella, 2004), the average of the Markov chain converges to the expectation under the stationary distribution:

$$\lim_{R \to \infty} \frac{1}{R} \sum_{r=1}^{R} h(\theta_r) = E_{\pi}[h(\theta)].$$

A Markov chain with this property is called ergodic. As a means of approximation the analyst relies on large but finite number of draws $R \in \mathbb{N}$ which can be selected in applications based

 $^{^{2}}$ For the sake of simplicity in the notation of this section we suppress the dependence of the posterior on data and other parameters not directly sampled.

on various criteria.

The conditional distribution $Q(\theta_r)$ can be obtained from a given (economic) model and its corresponding posterior. In many cases of interest, the model likelihood in $\pi(\theta)$ has a complicated form which precludes direct sampling from $\pi(\theta)$. In such case, the Metropolis-Hastings (M-H) principle is often utilized for drawing $\theta_{r+1}|\theta_r$ from $Q(\theta_r)$; see Chib and Greenberg (1995) for a detailed overview. Suppose we have a proposal-generating density $q(\theta_{r+1}^*|\theta_r)$ where θ_{r+1}^* is a proposed state given the current state θ_r of the Markov chain. The M-H principle stipulates that θ_{r+1}^* be accepted as the next state θ_{r+1} with the acceptance probability

$$\alpha(\theta_r, \theta_{r+1}^*) = \min\left[\frac{\pi(\theta_{r+1}^*)q(\theta_r|\theta_{r+1}^*)}{\pi(\theta_r)q(\theta_{r+1}^*|\theta_r)}, 1\right],\tag{1}$$

otherwise $\theta_{r+1} = \theta_r$. Then the Markov chain satisfies the so-called detailed balance condition

$$\pi(\theta_r)q(\theta_{r+1}^*|\theta_r)\alpha(\theta_r,\theta_{r+1}^*) = \pi(\theta_{r+1}^*)q(\theta_r|\theta_{r+1}^*)\alpha(\theta_{r+1}^*,\theta_r)$$

which is sufficient for ergodicity. $\alpha(\theta_{r+1}^*, \theta_r)$ is the probability of the move $\theta_r | \theta_{r+1}^*$ if the dynamics of the proposal generating mechanism were to be reversed. The proposal-generating density $q(\theta_{r+1}^*|\theta_r)$ is often chosen to be sampled easily. The popular Gibbs sampler arises as a special case when the M-H sampler is factored into conditional densities. The proposal draws $\theta_{r+1}^*|\theta_r$ from $q(\theta_{r+1}^*|\theta_r)$ in (1) are generated in one step.

3 Sequential Monte Carlo

A key challenge for MCMC methods, in particular in high-dimensional parameter spaces, is to find a good proposal density for the acceptance probability (1). Sequential Monte Carlo, also known as particle filter, encompasses a set of simulation-based methods that address this problem by constructing proposal densities sequentially with a number of desirable properties. SMC provides a convenient and computationally attractive numerical characterization of the posterior distribution.

The essence of SMC, with so-called data-tempering used here, can be summarized as follows (Herbst and Schorfheide, 2016). Let $p(Y|\theta)$ denote the likelihood and $p(\theta)$ the prior density.

The notation $Y_{1:N} = (Y_1, \ldots, Y_N)$ will be used as shorthand for vectors. Let ϕ_m , with $m = 1, \ldots, R_{\phi}$, be a sequence that slowly increases from zero to one. A sequence of posteriors can be constructed by sequentially adding observations to the likelihood function,

$$\pi_m^{(D)}(\theta) = \frac{p(Y_{1:\lfloor\phi_m N\rfloor}|\theta)p(\theta)}{\int p(Y_{1:\lfloor\phi_m N\rfloor}|\theta)p(\theta)d\theta}, \ \phi_m \uparrow 1, \ m = 1, \dots, R_{\phi},$$
(2)

where $\lfloor x \rfloor$ is the largest integer that is less than or equal to x. If ϕ_1 is close to zero then the $p(\theta)$ can provide an efficient proposal density for π_1 . SMC seeks to efficiently exploit $\pi_m(\theta)$ as a suitable proposal density for $\pi_{m+1}(\theta)$. As a result, SMC algorithms generate weighted draws from the sequence of posteriors $\{\pi_m(\theta)\}_{m=1}^{R_{\phi}}$. The weighted draws are called *particles*. Denote the overall number of particles by R_j . At any given stage m, the posterior $\pi_m(\theta)$ is represented by a swarm of particles $\{\theta_m^j, w_m^j\}_{j=1}^{R_j}$ in the sense that for the Monte Carlo average,

$$\overline{h}_{m,N_j} = \frac{1}{R_j} \sum_{j=1}^{R_j} w_m^j h(\theta^j) \stackrel{a.s.}{\to} E_\pi \left[h(\theta_m) \right].$$

Given the set of particles at stage m - 1, SMC proceeds in three steps: (i) correction: reweighting of the stage m - 1 particles to reflect the posterior at stage m; (ii) selection: resampling the particles with elimination of low-weight particles and multiplication of highweight particles; and (iii) mutation: propagating the particles forward using a Markov transition kernel. The details on each step are given further below in section 6.

4 Hamiltonian Dynamics

Hamiltonian (or Hybrid) Monte Carlo (HMC) is a class of MCMC methods featuring multistep distant proposals whose path follows the evolution of Hamiltonian dynamics. HMC has its roots in the physics literature where it was introduced for simulating molecular dynamics (Duane et al., 1987). It has since become popular in a number of application areas including statistical physics (Gupta et al., 1988; Akhmatskaya et al., 2009), computational chemistry (Tuckerman et al., 1993), and as a generic tool for Bayesian statistical inference (Neal, 1993, 2011; Ishwaran, 1999; Liu, 2004; Beskos et al., 2010). HMC is most applicable in situations when a suitable importance sampler is not available or practical to implement and one would thus typically need to rely on random walk sampling. HMC has been shown to yield samples far more efficient than obtained by the random walk Metropolis-Hastings mechanism (Rasmussen, 2003; Neal, 2011).

In contrast to the one-step proposals drawn in MCMC, Hamiltonian Monte Carlo (HMC) uses a *sequence* of steps in constructing the proposal whereby the last step in the sequence becomes the proposal draw. The proposal sequence is generated using difference equations of the law of motion yielding high acceptance probability even for proposals that are relatively distant from the current draw in the parameter space. This facilitates efficient exploration of the parameter space with the resulting Markov chain.

Consider a vector of parameters of interest $\theta \in \mathbb{R}^d$ distributed according to the posterior density $\pi(\theta)$. Let $\gamma \in \mathbb{R}^d$ denote a vector of auxiliary parameters with $\gamma \sim N(0, M)$, distributed Gaussian with mean vector 0 and covariance matrix M, independent of θ . Denote the joint density of (θ, γ) by $\pi(\theta, \gamma)$. Then the negative of the logarithm of the joint density of (θ, γ) is given by the Hamiltonian equation³

$$H(\theta, \gamma) = -\ln \pi(\theta) + \frac{1}{2} \ln \left((2\pi)^d |M| \right) + \frac{1}{2} \gamma' M^{-1} \gamma.$$
(3)

Hamiltonian Monte Carlo (HMC) is formulated in the following three steps that we will describe in detail further below:

- 1. Draw an initial auxiliary parameter vector $\gamma_r^0 \sim N(0, M)$;
- 2. Transition from (θ_r, γ_r) to $(\theta_r^L, \gamma_r^L) = (\theta_{r+1}^*, \gamma_{r+1}^*)$ according to the Hamiltonian dynamics;
- 3. Accept $(\theta_{r+1}^*, \gamma_{r+1}^*)$ with probability $\alpha(\theta_r, \gamma_r; \theta_{r+1}^*, \gamma_{r+1}^*)$, otherwise keep (θ_r, γ_r) as the next MC draw.

³In the physics literature, θ denotes the position (or state) variable and $-\ln \pi(\theta)$ describes its potential energy, while γ is the momentum variable with kinetic energy $\gamma' M^{-1} \gamma/2$, yielding the total energy $H(\theta, \gamma)$ of the system, up to a constant of proportionality. M is a constant, symmetric, positive-definite "mass" matrix which is often set as a scalar multiple of the identity matrix.

Step 1 provides a stochastic initialization of the system akin to a random walk (RW) draw. This step is necessary in order to make the resulting Markov chain $\{(\theta_r, \gamma_r)\}_{r=1}^R$ irreducible and aperiodic (Ishwaran, 1999). In contrast to RW, this so-called refreshment move is performed on the auxiliary variable γ as opposed to the original parameter of interest θ , setting $\theta_r^0 = \theta_r$. In terms of the HMC sampling algorithm, the initial refreshment draw of γ_r^0 forms a Gibbs step on the parameter space of (θ, γ) accepted with probability 1. Since it only applies to γ , it will leave the target joint distribution of (θ, γ) invariant and subsequent steps can be performed conditional on γ_r^0 (Neal, 2011).

Step 2 constructs a sequence $\{\theta_r^k, \gamma_r^k\}_{k=1}^L$ according to the Hamiltonian dynamics starting from the current state (θ_r^0, γ_r^0) and setting the last member of the sequence as the HMC new state proposal $(\theta_{r+1}^*, \gamma_{r+1}^*) = (\theta_r^L, \gamma_r^L)$. The role of the Hamiltonian dynamics is to ensure that the M-H acceptance probability (1) for $(\theta_{r+1}^*, \gamma_{r+1}^*)$ is kept close to 1. As will become clear shortly, this corresponds to maintaining the difference $-H(\theta_{r+1}^*, \gamma_{r+1}^*) + H(\theta_r^0, \gamma_r^0)$ close to zero throughout the sequence $\{\theta_r^k, \gamma_r^k\}_{k=1}^L$. This property of the transition from (θ_r, γ_r) to $(\theta_{r+1}^*, \gamma_{r+1}^*)$ can be achieved by conceptualizing θ and γ as functions of continuous time tand specifying their evolution using the Hamiltonian dynamics equations⁴

$$\frac{d\theta_i}{dt} = \frac{\partial H(\theta, \gamma)}{\partial \gamma_i} = \left[M^{-1} \gamma \right]_i , \qquad (4)$$

$$\frac{d\gamma_i}{dt} = -\frac{\partial H(\theta, \gamma)}{\partial \theta_i} = \nabla_{\theta_i} \ln \pi(\theta), \tag{5}$$

for i = 1, ..., d, where ∇_{θ_i} denotes the derivative of $\ln \pi(\theta)$ with respect to θ_i . For any discrete time interval of duration s, (4)–(5) define a mapping T_s from the state of the system at time t to the state at time t + s. For practical applications of interest these differential equations (4)–(5) in general cannot be solved analytically and instead numerical methods are required. The Stormer-Verlet (or leapfrog) numerical integrator (Leimkuhler and Reich, 2004) is one such popular method, discretizing the Hamiltonian dynamics as

$$\gamma(t + \varepsilon/2) = \gamma(t) + (\varepsilon/2)\nabla_{\theta} \ln \pi(\theta(t)), \tag{6}$$

$$\theta(t+\varepsilon) = \theta(t) + \varepsilon M^{-1} \gamma(t+\varepsilon/2), \tag{7}$$

$$\gamma(t+\varepsilon) = \gamma(t+\varepsilon/2) + (\varepsilon/2)\nabla_{\theta} \ln \pi(\theta(t+\varepsilon)), \tag{8}$$

⁴In the physics literature, the Hamiltonian dynamics describe the evolution of (θ, γ) that keeps the total energy $H(\theta, \gamma)$ constant.

for some small $\varepsilon \in \mathbb{R}$. From this perspective, γ plays the role of an auxiliary variable that parametrizes (a functional of) $\pi(\theta, \cdot)$ providing it with an additional degree of flexibility to maintain the acceptance probability close to one for every k. Even though $\ln \pi(\theta_r^k)$ can deviate substantially from $\ln \pi(\theta_r^0)$, resulting in favorable mixing for θ , the additional terms in γ in (3) compensate for this deviation maintaining the overall level of $H(\theta_r^k, \gamma_r^k)$ close to constant over $k = 1, \ldots, L$ when used in accordance with (6)–(8), since $\frac{\partial H(\theta, \gamma)}{\partial \gamma_i}$ and $\frac{\partial H(\theta, \gamma)}{\partial \theta_i}$ enter with the opposite signs in (4)–(5). In contrast, without the additional parametrization with γ , if only $\ln \pi(\theta_r^k)$ were to be used in the proposal mechanism as is the case in RW style samplers, the M-H acceptance probability would often drop to zero relatively quickly.

Step 3 applies a Metropolis correction to the proposal $(\theta_{r+1}^*, \gamma_{r+1}^*)$. In continuous time, or for $\varepsilon \to 0$, (4)–(5) would keep $-H(\theta_{r+1}^*, \gamma_{r+1}^*) + H(\theta_r, \gamma_r) = 0$ exactly resulting in $\alpha(\theta_r, \theta_{r+1}^*) = 1$ but for discrete $\varepsilon > 0$, in general, $-H(\theta^*, \gamma^*) + H(\theta, \gamma) \neq 0$ necessitating the Metropolis step. A key feature of HMC is that the generic M-H acceptance probability (1) can be expressed in a simple tractable form using only the posterior density $\pi(\theta)$ and the auxiliary parameter Gaussian density $\phi(\gamma; 0, M)$. The transition from (θ_r^0, γ_r^0) to (θ_r^L, γ_r^L) via the proposal sequence $\{\theta_r^k, \gamma_r^k\}_{k=1}^L$ taken according to the discretized Hamiltonian dynamics (6)–(8) is a deterministic proposal, placing a Dirac delta probability mass $\delta(\theta_r^k, \gamma_r^k) = 1$ on each (θ_r^k, γ_r^k) conditional on (θ_r^0, γ_r^0) . The system (6)–(8) is time reversible and symmetric in (θ, γ) , which implies that the forward and reverse transition probabilities $q(\theta_r^L, \gamma_r^L|\theta_r^0, \gamma_r^0)$ and $q(\theta_r^0, \gamma_r^0|\theta_r^L, \gamma_r^L)$ are equal: this simplifies the Metropolis-Hastings acceptance ratio in (1) to the Metropolis form $\pi(\theta_{r+1}^*, \gamma_{r+1}^*)/\pi(\theta_r^0, \gamma_r^0)$. From the definition of the Hamiltonian $H(\theta, \gamma)$ in (3) as the negative of the log-joint densities, the joint density of (θ, π) is given by

$$\pi(\theta,\gamma) = \exp\left[-H(\theta,\gamma)\right] = \pi(\theta)\left(\left(2\pi\right)^d |M|\right)^{-1/2} \exp\left(-\frac{1}{2}\gamma' M^{-1}\gamma\right).$$
(9)

Hence, the Metropolis acceptance probability takes the form

$$\begin{aligned} \alpha(\theta_r, \gamma_r; \theta_{r+1}^*, \gamma_{r+1}^*) &= \min\left[\frac{\pi(\theta_{r+1}^*, \gamma_{r+1}^*)}{\pi(\theta_r^0, \gamma_r^0)}, 1\right] \\ &= \min\left[\exp\left(-H(\theta_{r+1}^*, \gamma_{r+1}^*) + H(\theta_r^0, \gamma_r^0)\right), 1\right]. \end{aligned}$$

The expression for $\alpha(\theta_r, \gamma_r; \theta_{r+1}^*, \gamma_{r+1}^*)$ shows, as noted above, that the HMC acceptance probability is given in terms of the difference of the Hamiltonian equations $H(\theta_r^0, \gamma_r^0) - H(\theta_r^0, \gamma_r^0)$

 $H(\theta_{r+1}^*, \gamma_{r+1}^*)$. The closer can this difference be kept to zero, the closer the acceptance probability approaches one. A key feature of the Hamiltonian dynamics (4)–(5) in Step 2 is that they maintain $H(\theta, \gamma)$ constant over the parameter space in continuous time conditional on $H(\theta_r^0, \gamma_r^0)$ obtained in Step 1, while their discretization (6)–(8) closely approximates this property for discrete time steps $\varepsilon > 0$ with a global error of order ε^2 corrected by the Metropolis update in Step 3 (Neal, 2011). The goal of the Hamiltonian proposal transition dynamics is thus to maintain the proposal acceptance probability at or close to one even for a relatively long proposal sequence.

The acceptance ratio can only be maintained at exactly one if the proposal trajectory evolution were continuous. However, due its discretization into individual steps, the acceptance probability in general deviates from one due to discretization errors. The length of the proposal sequence can then be tuned using $\varepsilon > 0$ and L to achieve a desired acceptance rate, analogously to the RW environment. The Hamiltonian dynamics approximately keeps the joint density $\pi(\theta, \gamma)$ of θ and γ constant, permitting changes in the marginal density $\pi(\theta)$. Due to this feature, the proposal sequence does not move along a "straight" trajectory in the parameter space Θ of θ , but rather along a "curve". This ensures that the proposal sequence does not travel "too far" into the tails and stays in regions with non-zero probability.

Each proposal sequence in HMC and its extensions starts with a "refreshment" of the kinetic auxiliary variable γ newly drawn from the Gaussian distribution N(0, M) where M is the mass matrix. This draw determines the "direction" in which the proposal sequence will propagate through the parameter space. The stochastic nature of γ prevents the chain from getting stuck at the original point or too close to it.

4.1 Constraints

Parameter space constraints can be incorporated into the HMC proposal mechanism via "hard walls" representing a barrier against which the proposal sequence, simulating a particle movement, bounces off elastically. Constraints thus do not provide grounds for proposal rejection, eliminating any associated redundancies. Heuristically, the constraint is checked at each step of the proposal sequence and if it is violated then the trajectory of the sequence is reflected off the hard wall posed by the constraint. This facilitates efficient exploration of the parameter space even in parameter spaces that are constrained in a complex way.

5 Bayesian Nonparametric Mixture Modeling

Consider an exchangeable sequence $Y \equiv Y_{1:N}$ of random variables defined over a measurable space (Φ, \mathcal{D}) where \mathcal{D} is a σ -field of subsets of Φ . Denote the joint density of Y implied by an economic model by $f(Y;\theta)$ where $\theta \in \Theta$ is a Euclidean parameter. Further denote by G_0 the prior distribution of θ over a measurable space (Θ, \mathcal{B}) with \mathcal{B} being a σ -field of subsets of Θ , where G_0 admits a density g_0 .

In a parametric Bayesian model, the joint density of Y and θ is defined as

$$p(Y;\theta,G_0) = f(Y;\theta)g_0,$$
(10)

Conditioning on observed realizations y of Y turns $f(Y;\theta)$ into the likelihood function $p(\theta|y)$ and $p(Y;\theta,G_0)$ into the posterior density $\pi(\theta|G_0,y)$.

In the class of nonparametric Bayesian mixture models⁵ considered here, the joint density of Y and θ is defined as a mixture

$$p(Y; \theta, G) = \int f(Y; \theta) dG(\theta),$$

where G is the mixing distribution over θ . The distribution G is now random which leads to a flexibility of the resulting mixture model. The model parameters θ are no longer restricted to follow any given pre-specified distribution as was stipulated by the fixed G_0 in the parameteric case. The parameter space now also includes the random infinite-dimensional G with the additional need for a prior distribution for G. The Dirichlet Process prior is a popular alternative due to its numerous desirable properties; we proceed with its description in the next section.

⁵A commonly used technical definition of nonparametric Bayesian models are probability models with infinitely many parameters (Bernardo and Smith, 1994).

5.1 Dirichlet Process Prior

In a seminal paper, Fergusson (1973) introduced the Dirichlet process (DP) prior for random measures whose support is large enough to span the space of probability distribution functions and that leads to analytically manageable posterior distributions. Antoniak (1974) further elaborated on using the DP as the prior for the mixing proportions of a simple distribution.

A DP prior for G is determined by two parameters: a distribution G_0 that defines the "location" of the DP prior, and a positive scalar precision parameter α . The distribution G_0 may be viewed as a baseline prior that would be used in a typical parametric analysis. The flexibility of the DP prior model environment stems from allowing G – the actual prior on the model parameters – to stochastically deviate from G_0 . The precision parameter α determines the concentration of the prior for G around the DP prior location G_0 and thus measures the strength of belief in G_0 . For large values of α , a sampled G is very likely to be close to G_0 , and vice versa.

More specifically, let $\mathcal{M}(\Psi)$ be a collection of all probability measures on Ψ endowed with the topology of weak convergence. The space $\mathcal{M}(\mathcal{M}(\Psi))$ is then the collection of all probability measures (i.e. priors) on $\mathcal{M}(\Psi)$ together with the topology of weak convergence derived from $\mathcal{M}(\Psi)$. Let $G_0 \in \mathcal{M}(\Psi)$ and let α be a positive real number. Following Fergusson (1973), a *Dirichlet Process* on (Ψ, \mathcal{B}) with a base measure G_0 and a concentration parameter α , denoted by $DP(G_0, \alpha) \in \mathcal{M}(\mathcal{M}(\Psi))$, is a distribution of a random probability measure $G \in \mathcal{M}(\Psi)$ over (Ψ, \mathcal{B}) such that, for any finite measurable partition $\{\Psi_i\}_{i=1}^J$ of the sample space Φ , the random vector $(G(\Psi_1), ..., G(\Psi_J))$ is distributed as $(G(\Psi_1), ..., G(\Psi_J)) \sim Dir(\alpha G_0(\Psi_1), ..., \alpha G_0(\Psi_J))$ where $Dir(\cdot)$ denotes the Dirichlet distribution. We write $G \sim DP(G_0, \alpha)$ if G is distributed according to the Dirichlet process $DP(G_0, \alpha)$.

5.2 Dirichlet Process Mixture Model

Bayesian nonparametric mixture models have been widely applied to solving problems such as clustering, density estimation and topic modeling. These models make relatively very weak assumptions about the underlying process that generated the observed data. When more data are collected, the complexity of these models can change accordingly. In the Bayesian mixture modeling framework it is possible to infer the number of components to model the data and therefore it is unnecessary to explicitly restrict the number of components a-priori (Görür and Rasmussen, 2010).

For a nonparametric continuous density estimation the discrete Dirichlet process is typically convolved with a continuous kernel. There are many various ways of doing so. We follow the approach laid out by Ghosal and van der Vaart (2017, section 5.1) based on previous literature on Bayesian nonparametrics cited therein. For each $\theta \in \Theta \subset \mathbb{R}^d$, let $f(Y|\theta)$ be a probability density function of Y, where Y is an observable random variable. The density (10) where G is endowed with the Dirichlet process prior, is known as a Dirichlet process mixture (DPM). Realizations of the DP are discrete with probability one and hence a DPM can be viewed as a countably infinite mixture (Ghosal and van der Vaart, 2017).

For a sample size N, let Y_i with i = 1, ..., N be distributed according to the density kernel

$$p_{i,G} = \int f_i(Y_i|\theta) dG(\theta),$$

where $G \sim DP(G_0, \alpha)$. The resulting model can be equivalently written in terms of N latent variables θ_i as

$$\begin{split} Y_i | \theta_i, G &\sim f_i(Y_i | \theta_i), \\ \theta_i | G &\sim G, \\ G &\sim DP(G_0, \alpha). \end{split}$$

The model can also be represented in terms of allocation variables $s_1, ..., s_N$ that link the observations to the components of the mixture model:

$$Y_i|s_i^*, G = k \sim f_i(Y_i|\theta_k^*), \ i = 1, \dots, N,$$

where s_i^* and θ_k^* are the distinct values of s_i and θ_k , respectively. MCMC posterior inference for DPM models has been detailed in and number of studies, including Neal (2011) and Ghosal and van der Vaart (2017).

6 Hamiltonian Sequential Monte Carlo

Here we first provide the details of an SMC algorithm suited for our context and then propose its extension to form HSMC. SMC generally consists of three phases, as described above: (i) *correction*, (ii) *selection*, and (iii) *mutation*. The state-of-the-art procedure for the *correction* phase for a Bayesian static nonparametric model with a Dirichlet Process (DP) prior and a non-conjugate likelihood is Algorithm 2 of Griffin (2017), which we use for particle *correction* in both SMC and HSMC:

Correction phase:

Let $m_{i,k}$ denote the number of $s_{1:i}$ associated with the mixture component k, let m_0 denote the prior value for $m_{N,k}$ for all k, and let K_i denote the number of mixture components for $s_{1:i}$. For i = 1, ..., N:

Step 1: For all particles $j = 1, ..., R_j$, perform steps 1a and 1b. Step 1a: Sample $\theta_{new} \sim G_0$, and $s_i^{*(j)}$ conditional on $y_{1:i}$ and $s_{1:(i-1)}^{*(j)}$ from

$$q(k) \propto \begin{cases} m_{k,i-1}^{(j)} f_i(y_i | \theta_k^{*(j)}) \text{ if } 1 \le k \le K_{i-1}^{(j)} \\ m_0 f_i(y_i | \theta_{new}) \text{ if } k = K_{i-1}^{(j)} + 1. \end{cases}$$

Step 1b: Calculate the unnormalized weight

$$\xi_i^{(j)} = m_0 f_i(y_i | \theta_{new}) + \sum_{k=1}^{K_{i-1}^{(j)}} m_{k,i-1}^{(j)} f_i(y_i | \theta_k^{*(j)})$$

Step 2: Reweight the particles according to the weights

$$w_i^{(j)} = \frac{\xi_i^{(j)}}{\sum_{j=1}^{N_j} \xi_i^{(j)}}.$$

Although a number of alternative *selection* schemes have been proposed in the literature, we utilize the popular Residual Resampling as described in Chopin (2004).

Selection phase:

Reproduce each particle $int\{R_j w_i^{(j)}\}$ times, where $int\{x\}$ stands for the integer part of x. Complete the particle vector by R_j^r independent draws from $R_j^r = R_j - int\{R_j w_i^{(j)}\}$ draws from the multinomial distribution which reproduces the jth particle with probability $(R_j w_i^{(j)} - int\{R_j w_i^{(j)}\})/R_j^r$.

Mutation phase:

Propagate each parameter $\theta_k^{*(j)}$ for $j = 1, ..., R_j$ and $k = K_i^{(j)}$ according to Hamiltonian transition dynamics as described in section 4.

In contrast, SMC uses the random walk transition kernel in the mutation phase. To the best of our knowledge, HSMC has not been proposed in the previous literature.

7 The Nonparametric Mixed Logit Model

The mixed logit can approximate any random utility model (McFadden and Train, 2000) and remains popular among practitioners for its analytical tractability. Mixed logit models can be obtained under different behavioral specifications, and each derivation provides a particular interpretation of the model fundamentals. Any behavioral specification whose choice probabilities take its particular form is called a mixed logit model (Train, 2009).

7.1 Model Environment

There are N individuals, i = 1, ..., N, choosing in each of T time periods, t = 1, ..., T, one out of J alternatives, j = 1, ..., J. Let y_{it} denote the choice of individual i at time t. The latent utility of individual i at time t of choice j is given by

$$u_{itj} = \boldsymbol{\beta}_i' \mathbf{x}_{itj} + \varepsilon_{itj},$$

with latent iid residual $\varepsilon_{itj} \sim F_{\varepsilon}$ where F_{ε} is the Extreme Value Type 1 distribution. The first element of β_i is normalized to zero for identification purposes. Then, conditional on the vector of covariates $x_{it} = (x_{it1}, \ldots, x_{itJ})'$ and the vector of parameters β_i , the probability of choosing y_{it} at time t is given by

$$L_{it}(y_{it}|\boldsymbol{\beta}_i, \mathbf{x}_{it}) = rac{\exp(\boldsymbol{\beta}'_i \mathbf{x}_{ity_{it}})}{\sum_{j=1}^J \exp(\boldsymbol{\beta}'_i \mathbf{x}_{itj})}$$

and the probability of choosing the vector $y_i = (y_{i1}, \ldots, y_{iT})'$ is given by

$$K(y_i|\boldsymbol{\beta}_i, \mathbf{x}_i) = \prod_{t=1}^T L_{it}(y_{it}|\boldsymbol{\beta}_i, \mathbf{x}_{it}).$$

The mixed logit model specification is obtained by expressing the choice probabilities in the form

$$P_i(y_i|\mathbf{x}_i) = \int K(y_i|\boldsymbol{\beta}_i, \mathbf{x}_i) f(\boldsymbol{\beta}_i) d\boldsymbol{\beta}_i \;.$$

The mixed logit probability is a weighted average of the logit formula evaluated at different values of β_i , where the weights are given by the density $f(\beta_i)$.

Under the Bayesian nonparametric mixture model approach, we specify the model for the distribution of β_i as follows:

$$y_i | \boldsymbol{\beta}_i \sim K(y_i | \boldsymbol{\beta}_i, \mathbf{x}_{it}),$$
$$\boldsymbol{\beta}_i | \boldsymbol{G} \sim \boldsymbol{G},$$
$$\boldsymbol{G} \sim DP(\boldsymbol{G}_0, \alpha).$$

with G_0 a standard Normal distribution and α obtained implicitly by setting $m_0 = 1$.

Fox et al. (2012) showed that the mixed logit model is nonparametrically identified. Fox and Gandhi (2016) analyze a nonparametric estimator for the case when the observable random variables have a discrete support. Fox et al. (2016) propose a computationally attractive projection-based estimator of the joint distribution of random coefficients over a fixed support grid in structural models including the mixed logit. The Bayesian framework allows for continuous support of observable random variables and does not impose the fixed support grid restriction on the parameter space.

In Bayesian multinomial choice modeling, MCMC has so far been the dominant approach to inference (Kim et al., 2004; Burda et al., 2008; Keane and Wasi, 2013; Li and Ansari, 2014). Although SMC has been utilized in analysis of generic Bayesian DPM models, we are not aware of its application to Bayesian multinomial discrete choice model. In the sequel, we will estimate the distribution of β_i by both HSMC and SMC in a real-world application. We will then evaluate and compare the convergence properties of both methods.

7.2 Data

Our empirical analysis is based on the IRI Academic Dataset (Bronnenberg et al., 2008), containing panel data of grocery stores purchases in two U.S. cities. We chose to focus on the purchases of mayonnaise since this product category is composed of relatively few well defined homogenous items. In our data set, two dominating brands cover 87% of the market: Hellman's (46%) and Kraft (41%). The remainder of the market is served by "private label" (8%), Cains (3%), and "other" (2%).

We use the time period from June 2010 through December 2012, totalling 138 weeks. During this time period, the data contains a stable choice set without introducing new or discontinuing old products in the set of the choice alternatives. Each of the 2,684 households in our sample was recorded as making mayonnaise purchases on average for 7.86 weeks. The distribution of weeks observed in the sample for all households is shown in Figure 1.



Figure 1: Weeks Observed Making Mayonnaise Purchases

Similarly to Thomadsen (2016), we assume that consumers choose among a set of "top" alternatives, or else choose an outside option if they choose a product in the category that does not belong to a top alternative. The "top" alternatives are selected by ranking the alternatives by the number of purchases made by the panelists, include alternatives starting from the most popular ones and going in decreasing order of popularity until the set of included top alternatives covers all major ones. Thus, we consider the following six alternatives plus an outside option, as given in Table 1.

Label	Name	Frequency	%	Cum. %
1	Hellmann's Real Mayonnaise	7,041	24.42	24.42
2	Kraft Miracle Whip	$6,\!189$	19.88	44.30
3	Kraft Miracle Whip Low Fat	3,031	11.35	55.65
4	Hellmann's Light Mayonnaise	1,752	5.26	60.91
5	Kraft Soybean Mayonnaise	1,015	4.43	65.34
6	Hellmann's Soybean Mayonnaise	786	2.05	67.38
0	outside option	$12,\!533$	32.62	100.00

Table 1: Choice Set

The panel contains information about product attributes and consumer characteristics. Given the high degree of product homogeneity within any given alternative category, in addition to brand we only included price among the product attributes. The price dispersion for each product in the choice set is shown in Figure 2. The Label of the alternatives corresponds to the product name code listed in Table 1. Although on a given choice occasion price is only observed for the selected alternative, we infer the prices of the remaining alternatives of the choice set from observations of other customers who selected such alternatives in any given store.

In order to keep dimensionality of the parameter vector low, we have selected income as a key consumer characteristic. Table 2 specifies the ordinal coding for the income ranges in our dataset and Figure 3 shows a histogram of the income data codes in our sample. We have dropped all households whose income data was missing.

The utility of the outside option has been normalized to zero for identification purposes. The model contains an individual-specific intercept for each of the choice alternatives, other than the outside option. With three random parameters (intercept, price, income) per each of the six choice alternatives, our model contains 18 parameters whose joint distribution we seek to estimate nonparametrically.



Figure 2: Choice Set Price Dispersion

Code	Household Income per Year
1	\$00,000 to \$ 9,999
2	\$10,000 to \$11,999
3	12,000 to $14,999$
4	\$15,000 to \$19,999
5	\$20,000 to \$24,999
6	\$25,000 to \$34,999
7	\$35,000 to \$44,999
8	\$45,000 to \$54,999
9	\$55,000 to \$64,999
10	\$65,000 to \$74,999
11	\$75,000 to \$99,999

 Table 2: Income Codes

7.3 Implementation

In the implementation, we have run both HSMC and SMC for one hour of wallclock time. The implementation was run with a Coarray Fortran 2008 code using Intel 2016 compiler on 4 nodes of a 40-core 2.4 GHz Linux cluster (Loken et al., 2010). We used 20 steps in



Figure 3: Household Income Distribution

constructing the Hamiltonian proposal in HSMC and tuned the step size to achieve transition acceptance rate of about 80%. Theoretical analysis of optimal step sizes and acceptance rates for HMC is provided in Beskos et al. (2010). We introduced the data in ten batches of equal size, one per 100 iterations. We tuned the SMC step size to achieve transition acceptance rates of about 30% (Roberts et al., 1997). Due to the Hamiltonian transition dynamics, HSMC takes somewhat longer than SMC to complete one full iteration but features superior mixing properties. During the run, HSMC completed about 8,800 iterations while SMC completed 10,650 iterations.

Our model is nonparametric and in this context each particle represents a mixture of parametric kernels, where the number of kernels is stochastic and fluctuates during the implementation run. HSMC sampled on average 21 kernels per each particle mixture while SMC sampled on average 24 kernels. We used 3,200 particle mixtures and thus each procedure utilized on average over 67,000 parameter vectors, each of which had 18 dimensions.

7.4 Performance Comparison

For the assessment of the performance a posterior sampler, it is standard practice to rely on convergence diagnostics obtained by examining the sampling output (Cowles and Carlin, 1996). A typical MCMC diagnostic starts several Markov chains at overdispersed initial values, and monitors convergence by comparing between-chain and within-chain variances for selected scalar quantities (Plummer et al., 2006). However, the bulk of such diagnostics is not applicable to particle-based samplers, including SMC and HSMC, as a significant proportion of sample chain paths are discontinued during the implementation during the resampling phase. In the absence of chains of parameter draws of full equal length the chains cannot be compared in terms of their sampling behavior. Furthermore, very few diagnostics have been designed to asses convergence outside of Euclidean spaces. Yet, in our case we are interested in convergence of a nonparametric object, the distribution of β_i .

7.4.1 PACE Diagnostic

For the purpose of comparing the convergence properties of HSMC and SMC we will use the *Partition-based approximation for convergence evaluation* (PACE) diagnostic procedure developed by VanDerwerken and Schmidler (2017) to address the limitations of other diagnostics discussed above. The PACE statistic involves initializing J sets of particles at overdispersed locations in the state space. At a given iteration, the trajectories of all particle draws are pooled together. The distance between the sample distributions of each particle set is quantified by comparing the within-set and across-set probabilities over a partition of the parameter space. When the particle sets are stationary, the proportion of within-set draws belonging to a given partition element will be approximately equal across the particle sets. Heuristically, when each particle set results in approximately the same posterior probability for a given partition element of the parameter space then the sets can be regarded as having converged. In contrast, when different particle sets imply different posterior probabilities for a given parameter space partition element then none of the sets can be guaranteed to have converged. The PACE statistic is based on a comparison of approximate posterior probabilities over a parameter space partition. The posterior probabilities can be quantified using any number of particles without requiring that the chains of individual particle draws be of full equal length. The particles are thus free to be resampled in the SMC selection phase. Furthermore, the parameter space can be either Euclidean or a function space which renders PACE suitable for nonparametric estimation problems.

VanDerwerken and Schmidler (2017) proposed the PACE statistic using an adaptive parameter space partition whereby pooled sampler draws are suitably clustered in order to construct the partition. We used a simpler version in which the partition is constructed over a fixed equidistant grid over the parameter space in a non-adaptive manner. This saves on computation time substantially and avoids introducing an ad-hoc clustering procedure which may act differently in HSMC and SMC obscuring the differences stemming from these two procedures alone. Correspondingly, we used a mean-absolute deviation (MAD) measure as the PACE distance function.

In our application we estimate the density of β_i , which is 18 dimensional. Obtaining PACE in such relatively high-dimensional space turned out computationally prohibitive. For the sake of feasibility, we have implemented PACE for the bivariate distributions of all pairwise combinations of elements in β_i which reflect at least to some extent the information contained in the joint distribution of β_i beyond the univariate margins. We then calculated the average PACE statistic as a function of Monte Carlo iterations. The results are presented in Figure 4. Both methods seem to have converged well within the first half of the run. Throughout the run PACE of HSMC has dominated SMC by a substantial margin, attesting to the superior mixing properties of HSMC.

7.4.2 Estimated Distribution of Coefficients

Here we also report the output on estimated distribution of the mixed multinomial logit coefficients. Table 3 provides summary statistics, mean and standard deviation of a benchmark parametric model Mlogit, SMC, and HSMC. The parametric model Mlogit was implemented



Figure 4: PACE

by the command $mlogit^6$ in R (Croissant and Réunion, 2012).

However, the summary statistics obscure important information about the shape of the estimated densities of the coefficients, which is undetectable in the parametric model. The estimated coefficient densities are presented in Figure 5. The probability mass of the HSMC densities are generally somewhat farther away from zero (prior mean) than SMC densities, suggesting that the former has explored the parameter space and updated the posterior more effectively than the latter. The overall pattern of the densities reveals that the income coefficients, $\beta_{i,13} - \beta_{i,18}$, are much closer to zero than the intercept or price coefficients. Nonetheless, several income coefficient densities feature a prominent left tail, suggesting a negative income effect. The intercept coefficients, $\beta_{i,1} - \beta_{i,6}$, tend to have more probability mass distributed on the positive side of the real line, while the price coefficients, $\beta_{i,7} - \beta_{i,12}$, on the negative side. This pattern is in general agreement with the parametric benchmark

⁶mlogit does not take into account the time dimension. With our dataset, mlogit failed to converge for alternative-specific price coefficients and hence we were only able to obtain output for a common price parameter across all choice alternatives.

		Mlogit		SMC		HSMC	
Coefficient	Alternative	Mean	St.dev.	Mean	St.dev.	Mean	St.dev.
	1	2.428	0.121	0.000	0.024	0.0275	0.046
	2	2.817	0.109	0.005	0.026	0.0310	0.052
Intercept	3	1.201	0.142	-0.001	0.024	0.0107	0.036
	4	0.985	0.162	-0.006	0.025	-0.0118	0.035
	5	1.248	0.163	-0.005	0.025	0.0013	0.030
	6	3.516	0.216	-0.004	0.024	-0.0053	0.030
	1	-1.407	0.035	-0.011	0.028	-0.0288	0.048
	2	-1.407	0.035	-0.014	0.035	-0.0394	0.066
Price	3	-1.407	0.035	-0.014	0.032	-0.0355	0.053
	4	-1.407	0.035	-0.013	0.032	-0.0265	0.044
	5	-1.407	0.035	-0.015	0.033	-0.0321	0.050
	6	-1.407	0.035	-0.013	0.029	-0.0221	0.032
	1	-0.029	0.011	-0.002	0.016	-0.0004	0.015
	2	-0.088	0.011	-0.002	0.018	-0.0030	0.016
Income	3	0.015	0.025	-0.008	0.023	-0.0118	0.024
	4	0.042	0.026	-0.011	0.024	-0.0127	0.024
	5	-0.009	0.028	-0.010	0.027	-0.0128	0.023
	6	-0.018	0.020	-0.010	0.025	-0.0064	0.020

Table 3: Summary of Estimated Coefficients

Mlogit model estimates obtained in R.

8 Conclusions

In this paper, we have proposed Hamiltonian Sequential Monte Carlo (HSMC), which uses Hamiltonian transition dynamics in particle mutation phase, in place of random walk transitions used in Sequential Monte Carlo (SMC), in the context of a Bayesian nonparametric mixture model. HSMC combines the advantages of SMC in terms of convenience of approximation of complex posterior shapes and parallelizability with the benefits of superior convergence properties stemming from Hamiltonian transition dynamics utilizing information about the first derivative of the likelihood function. We have applied SMC and HSMC to a panel discrete choice model with a nonparametric distribution of unobserved individual heterogeneity, using the IRI panel data set. We have contrasted both methods in terms of convergence properties and showed the favorable performance of HSMC.



Figure 5: Estimated coefficient densities: HSMC (solid line) and SMC (dashed line)

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