Random Coordinate Underdamped Langevin Monte Carlo

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Abstract

The Underdamped Langevin Monte Carlo (ULMC) is a popular Markov chain Monte Carlo sampling method. It requires the computation of the full gradient of the log-density at each iteration, an expensive operation if the dimension of the problem is high. We propose a sampling method called Random Coordinate ULMC (RC-ULMC), which selects a single coordinate at each iteration to be updated and leaves the other coordinates untouched. We investigate the computational complexity of RC-ULMC and compare it with the classical ULMC for strongly log-concave probability distributions. We show that RC-ULMC is always cheaper than the classical ULMC, with a significant cost reduction when the problem is highly skewed and high dimensional. Our complexity bound for RC-ULMC is also tight in terms of dimension dependence.

1 Introduction

Langevin Monte Carlo (LMC) is a popular Monte Carlo sampling method, widely used in Bayesian statistics and machine learning (Andrieu et al., 2003). The goal is to construct a Markov chain that approximately generates i.i.d. samples from a target distribution given by (with some abuse of notation, we do not distinguish a distribution with its density)

\[ p_X(x) = \frac{1}{Z} e^{-f(x)} , \]

where \( Z \) is a normalizing constant that ensures \( \int p_X(x) \, dx = 1 \). Throughout the paper we assume \( f(x) \) is a convex function on \( \mathbb{R}^d \), and thus \( p_X(x) \) is a log-concave probability distribution.

Among the many Monte Carlo sampling methods, LMC (Rossky et al., 1978; Parisi, 1981; Roberts and Tweedie, 1996) stands out for its simplicity: For each iteration one updates the location of the particle by descending along the gradient and adding properly scaled Gaussian noise. For strongly log-concave distributions, it has been established in recent years that the empirical distribution of the iterate in LMC converges exponentially fast to the target distribution, with total computational cost \( \tilde{O}(d^2/\epsilon^2) \) to achieve \( \epsilon \) accuracy in Wasserstein distance (Dalalyan and Karagulyan, 2019; Durmus et al., 2019). Here and throughout the paper, we measure “cost” in terms of the total number of evaluations of a single element of the gradient, and assume that a full gradient evaluation requires about \( d \) times as much computation as a single component of the gradient.

To reduce the computational cost of sampling, the underdamped version of Langevin dynamics has recently been used to design the ULMC algorithm. By augmenting the state space with velocity variables, the underdamped Langevin dynamics converges to the equilibrium faster than the classical Langevin dynamics (Villani, 2006; Dolbeault et al., 2009; Baudoin, 2016; Monmarché, 2018; Cao et al., 2019). As a consequence, ULMC, the discrete version of the underdamped Langevin dynamics, also achieves faster convergence than LMC, the discrete version of the classi-
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cal Langevin dynamics [Cheng et al. 2018a; Dalalyan and Riou-Durand 2018] and [Eberle et al. 2018]. To be more specific, to obtain ϵ-accuracy in sampling from a log-concave pX, the computational complexity of ULMC is $O(d^{3/2}/\epsilon)$, whereas that of LMC is $O(d^2/\epsilon^2)$. This is an improvement in both $d$ and $\epsilon$.

This work aims at further improving the algorithm in terms of its dimension dependence, especially for the very high dimensional problems that arise often in practical applications (see [Ding et al. 2020b] for discussions of several examples). For these problems, ULMC requires a full evaluation of the gradient $\nabla f$ at each iteration, which often costs a factor of $O(d)$ greater than evaluating of a single component of the gradient. This factor arises when the expression of $\nabla f$ is not known explicitly, such as in partial differential equation (PDE) based inverse problems, where $f$ is given implicitly by solving the forward problem given as a PDE, and finite-difference approximation to the full gradient would be $d$ times more expensive than a single component. While automatic differentiation techniques have been developed, the cost of evaluation of the gradient often still leads to formidable computational and memory cost.

Other examples in which there is a factor-of-$d$ difference in evaluation cost between a full gradient and single component of the gradient come from problems with particular structures, such as graph-based problems. Given a graph with nodes $\mathcal{N} = \{1, 2, \ldots, d\}$ and directed edges $\mathcal{E} \subset \{(i, j) : i, j \in \mathcal{N}\}$, suppose there is a scalar variable $x_i$ associated with each node $i = 1, 2, \ldots, d$, and that the function $f$ has the form $f(x) = \sum_{(i,j) \in \mathcal{E}} f_{ij}(x_i, x_j)$. Then the partial derivative of $f$ with respect to $x_i$ is given by

$$\frac{\partial f}{\partial x_i} = \sum_{j : (i, j) \in \mathcal{E}} \frac{\partial f_{ij}}{\partial x_i}(x_i, x_j) + \sum_{l : (l, i) \in \mathcal{E}} \frac{\partial f_{li}}{\partial x_l}(x_l, x_i).$$

Note that the number of terms in the summations in this expression equals the number of edges in the graph that touch node $i$, the expected value of which is about $2/d$ times the total number of edges in the graph. Meanwhile, evaluation of the full gradient would require evaluation of both partial derivatives of each component function $f_{ij}$ for all edges in the graph, leading to a factor-of-$d$ difference in evaluation cost.

In this work, we target these problems in which single components of the gradient are much less expensive than full gradients by incorporating the random coordinate descent (RCD) method from optimization into underdamped Langevin sampling algorithm. RCD differs from gradient descent (GD) in that it updates just a single component, chosen at random, of the variable vector $x$ at each iteration. It takes a step in the negative gradient direction in just this component, leaving other components unchanged. (By contrast, gradient descent takes a step along the full negative gradient direction.) When there is a factor-of-$d$ difference in cost between evaluating the full gradient and a single component of the gradient, worst-case bounds for convex problems are better for RCD than for GD, the cost reduction being particularly significant when the dimension $d$ is high and $f$ is “skewed” in a sense to be defined later. Specifically, it was shown in [Nesterov 2012] that when the coordinate is chosen from a distribution weighted according to the directional Lipschitz constants, the complexity is reduced from $dk$ to $d\kappa_{\text{max}}$, where $\kappa$ and $\kappa_{\text{max}}$ are conditioning of $f$ and the maximum directional conditioning of $f$ respectively. Since $\kappa_{\text{max}} \leq \kappa$ for all functions [Wright 2015], RCD is always cheaper than GD. Further, when the dimension of the problem is high and $f$ is skewed in the sense that $\kappa \approx d\kappa_{\text{max}}$, the reduction in cost approaches a factor of $d$.

In this paper, we propose the random coordinate underdamped Langevin Monte Carlo (RC-ULMC) algorithm. We aim to improve the convergence of ULMC by utilizing cheaper steps, as in RCD, so we establish non-asymptotic convergence rates for RC-ULMC and compare with classical ULMC. Our main results are as follows:

1. The convergence rate of RC-ULMC depends on directional conditioning; see Theorem 6.2.
2. Comparing with ULMC, RC-ULMC is always cheaper than the classical ULMC, the change being $d^{3/2}\kappa^{3/2} \rightarrow (d^{3/2} + \kappa)\kappa^{1/2}$. This cost reduction is significant when $f$ is skewed and the dimension is high; see the discussion following Corollary 6.3.
3. The complexity bound of the RC-ULMC we obtain is tight in both $d$ and $\epsilon$; see Proposition 6.4.

The remainder of the paper is as follows. We review literature in Section 2 and summarize basic notations and assumptions in Section 3. In Section 4 we review ULMC and its convergence properties. In Section 5 we present our new method RC-ULMC. Our main results are presented in Section 6, where we discuss the non-asymptotic convergence rate, the numerical cost, the cost saving compared to the classical ULMC, and the tightness of the result. Computational results are presented in Section 7. Technical derivations and proofs appear in Supplementary Materials.

2 Related works

The non-asymptotic analysis of LMC and ULMC sampling methods has been an active area [Cheng et al. 2018a; Dalalyan and Riou-Durand 2018; Dalalyan and...
Karagulyan [2019] Durand et al. [2019]; and it has been established that ULMC gives a faster convergence under the same log-concavity and smoothness assumptions on the distribution (Cheng et al. [2018a] Dalalyan and Riou-Durand [2018]). When ULMC is modified with a better discretization scheme, e.g., the random midpoint method, the computational complexity can be even further reduced (Shen and Lee, 2019; He et al., 2020). For ULMC, it was established in Dalalyan and Riou-Durand [2018] that it achieves $\epsilon$ error in Wasserstein metric within $O\left(d^{3/2}K^{3/2}/\epsilon\right)$ iterations, where $O$ hides log factors. The total cost of ULMC is therefore $O\left(d^{3/2}K^{3/2}/\epsilon\right)$. In comparison, the RC-ULMC method proposed in this paper is always cheaper and the saving can be significant for highly skewed distributions in high dimension.

The combination of RCD and LMC (based on overdamped Langevin dynamics) has been recently explored in works (Shen et al., 2019; Ding et al., 2020b). This algorithm will be referred to as RC-OLMC (where “O” stands for overdamped). Compared with their result, the method in this paper converges faster both in terms of $d$ and $\epsilon$, similar to the saving obtained going from LMC to ULMC. This will be discussed further in Section 6.

Alternative sampling strategies have been developed without using the full gradient $\nabla f$ at each step. A standard approach is the Random Walk Metropolis algorithm, which combines a random walk proposal with Metropolis-Hastings acceptance-rejection step (Hastings, 1970), and thus only uses $f$ at each iteration. However, they are less efficient in high dimensions compared with gradient based methods (Mattingly et al., 2012; Pillai et al., 2012). There have been recent interests in ensemble based sampling methods, in particular in the context of data assimilation, inspired by the ensemble Kalman filter (Evensen, 2006), such as (Garbuno-Inigo et al., 2020; Iglesias et al., 2013). Unfortunately, none of these methods can be completely “gradient-free” and at the same time consistent for non-Gaussian distributions (Ding and Li, 2019a). To achieve consistency, one can try to incorporate weights to particles, as is done in importance sampling (Geweke, 1989) or sequential Monte Carlo (Doucet et al., 2001), however such methods often face the difficulty of high variance (Ding et al., 2020a).

When the log-density $f(x)$ has the form of $f(x) = \sum_{i=1}^{\mathcal{N}} f_i(x)$, one can randomly select a representative $\nabla f_i$ as a stochastic approximation to the full gradient, where $r$ is uniformly chosen from $\{1, \cdots, \mathcal{N}\}$. This leads to the stochastic gradient Langevin Monte Carlo method (Welling and Teh, 2011). Note that in general we can write the full gradient as $\nabla f = \sum_{i=1}^{d} \partial_i f e_i$ (where $e_i$ is the unit vector in $i$-th direction), and thus RCD and stochastic gradient, while used for different setups, share some similarity in reducing the cost of gradient evaluation.

Throughout the paper we do assume the log-concavity of $p_X$ (the convexity of $f$). For optimization problems, there are abundant works relaxing this assumption (Gelfand and Mitter [1991] Raginsky et al., 2017), but to our knowledge, no results on non-asymptotic convergence rate has been obtained for Bayesian sampling problems. One exception is (Cheng et al., 2018b) where the authors nevertheless require the convexity of $f$ outside a ball of finite-size.

3 Notations and assumptions

Throughout the paper we assume convexity and Lipschitz continuity of $f$.

**Assumption 3.1.** The function $f$ is second-order differentiable and $\mu$-strongly convex for some $\mu > 0$ and the gradient $\nabla f$ is $L$-Lipschitz. Specifically, we have: for all $x, x' \in \mathbb{R}^d$

$$f(x) - f(x') - \nabla f(x')^\top (x - x') \geq \frac{\mu}{2} |x - x'|^2$$

and

$$|\nabla f(x) - \nabla f(x')| \leq L|x - x'|.$$ (3)

Since the full gradient is Lipschitz continuous, so is its directional derivative. We denote directional Lipschitz constants by $L_i, i = 1, 2, \ldots, d$, meaning that

$$|\partial_i f(x + te_i) - \partial_i f(x)| \leq L_i |t|,$$ (4)

for any $i = 1, 2, \ldots, d$, any $x \in \mathbb{R}^d$, and any $t \in \mathbb{R}$.

Denote $\nabla^2 f$ the Hessian, then the assumption implies that

$$\mu I_d \preceq \nabla^2 f(x) \preceq L I_d, \quad |\partial_{ii} f(x)| \leq L_i,$$ 

where $\partial_{ii} f(x)$ is the $(i, i)$-element of $\nabla^2 f(x)$. We also define condition numbers:

$$\kappa = L/\mu \geq 1, \quad \kappa_i = L_i/\mu \geq 1, \quad \kappa_{\max} = \max_i \kappa_i.$$ (5)

As shown in Wright (2015), we have

$$\kappa_i \leq \kappa_{\max} \leq \kappa \leq d\kappa_{\max}.$$ (6)

We note that both inequalities, $\kappa_{\max} \leq \kappa$ and $\kappa \leq d\kappa_{\max}$ are sharp. If $\nabla^2 f$ is a diagonal matrix, then $L_{\max} = L$, both being the largest eigenvalue of $\nabla^2 f$, so that $\kappa_{\max} = \kappa$. This is the case when all coordinates are independent of each other, for example $f = \sum_i \lambda_i x_i^2$. On the other hand, if $f$ is highly
skewed, such as $f = (\sum_i x_i)^2$, so $\nabla^2 f = e \cdot e^\top$ (where $e = [1, 1, \ldots, 1]^\top$), then $L = dL_{\text{max}}$ and $\kappa = d\kappa_{\text{max}}$.

Furthermore, since $\kappa_i \geq 1$, we have for $p > 0$ that

$$(d - 1) + \kappa_{\text{max}}^p \leq \sum_{i=1}^d \kappa_i^p \leq \kappa_{\text{max}}^p \leq \kappa_{\text{max}}^p.$$  

Both bounds are tight. In the case when $f = \kappa_1 |x_1|^2 + \sum_{i=2}^d |x_2|^2$ with $\kappa_1 > 1$, $\kappa_{\text{max}} = (d - 1) + \kappa_{\text{max}}^p$. On the other hand, when $f = \sum_i x_i^2$, we have $\kappa_i = \kappa_{\text{max}} = 1$, then $\sum_{i=1}^d \kappa_i^p = d\kappa_{\text{max}}^p = d\kappa_{\text{max}}^p$.

And we say $f$ is highly skewed if

$$\sum_{i=1}^d \kappa_i^p \approx (d - 1) + \kappa_{\text{max}}^p .$$  

(7)

The extreme example in this setting is to set $f(x) = \frac{1}{2} (dx_i^2 + \sum_{i=2}^d x_i^2)$ with $d \gg 1$. In this example, $\kappa_{\text{max}} = d$ and $\sum_{i=1}^d \kappa_i^p = (d - 1) + \kappa_{\text{max}}^p$.

To measure the distance between two probability distributions, we use the Wasserstein distance.

**Definition 3.1.** The Wasserstein distance $W_p$ (for any $p \geq 1$) between probability measures $\mu$ and $\nu$ is defined as

$$W_p(\mu, \nu) = \left( \inf_{(X, Y) \in \Gamma(\mu, \nu)} \mathbb{E}[|X - Y|^p] \right)^{1/p} ,$$  

where $\Gamma(\mu, \nu)$ is the set of distribution of $(X, Y) \in \mathbb{R}^{2d}$ whose marginal distributions, for $X$ and $Y$ respectively, are $\mu$ and $\nu$.

In this paper, we will use the 2-Wasserstein metric $W_2$.

### 4 Classical ULMC

Underdamped Langevin dynamics is characterized by the following SDE:

$$\begin{cases}
  dX_t = V_t \, dt , \\
  dV_t = -2V_t \, dt - \gamma \nabla f(X_t) \, dt + \sqrt{4\gamma} \, dB_t,
\end{cases}$$  

(8)

where $\gamma > 0$ is a parameter to be tuned, and $B_t$ is the Brownian motion. Here we use the parametrization form of [Cheng et al., 2018] (alternative parametrizations are used in Dalalyan and Riou-Durand [2018]; Shen and Lee [2019]). Denoting by $q(x, v, t)$ the probability density function of $(X_t, V_t)$, we have that $q$ satisfies the Fokker-Planck equation

$$\partial_t q = \nabla \cdot \left( \frac{-v}{2v + \gamma \nabla f} q + \left[ \begin{array}{cc} 0 & 0 \\ 0 & 2\gamma \end{array} \right] \nabla q \right).$$  

It is well known that under mild conditions, $q$ converges to $p(x, v) \propto \exp(-(f(x) + |v|^2/2\gamma))$ (see e.g., Villani [2006]; Dolbeault et al. [2009]; Baudoin [2016]; Cao et al. [2019]), and thus the marginal density function for $x$ becomes the target distribution $p_X(x)$.

Denoting by $h > 0$ the time stepsize, we have that for $t \in [mh, (m + 1)h]$, (8) is equivalent to

$$\begin{align*}
  X(t) &= X(mh) + \frac{1 - e^{-2th}}{2} V(mh) - \frac{\gamma}{2} \int_{mh}^t \left(1 - e^{-2(t-s)}\right) \nabla f(X(s)) \, ds \\
  V(t) &= V(mh)e^{-2t} - \gamma \int_{mh}^t e^{-2(t-s)} \nabla f(X(s)) \, ds + \sqrt{4\gamma e^{-2t}} \int_{mh}^t e^{2s} \, dB_s ,
\end{align*}$$  

(9)

The sampling method, ULMC, can be viewed as a numerical solver for (9)-(10) based on the Euler approximation. Denoting by $(x^m, v^m)$ the numerical approximation to $(X(mh), V(mh))$, and replacing $X(s)$ in (9)-(10) by $x^m$, Euler approximation yields that $(x^{m+1}, v^{m+1}) \in \mathbb{R}^{2d}$ are two Gaussian random vectors with the following expectation and covariance:

$$\begin{align*}
  E x^{m+1} &= x^m + \frac{1}{2} (1 - e^{-2h}) v^m - \frac{\gamma}{2} \left( h - \frac{1}{2} (1 - e^{-2h}) \right) \nabla f(x^m) , \\
  E v^{m+1} &= v^m e^{-2h} - \frac{\gamma}{2} \left( 1 - e^{-2h} \right) \nabla f(x^m) , \\
  \text{Cov} (x^{m+1}) &= \gamma \left[ h - \frac{3}{4} - \frac{1}{4} e^{-4h} + e^{-2h} \right] I_d , \\
  \text{Cov} (v^{m+1}) &= \gamma \left[ 1 - e^{-4h} \right] I_d , \\
  \text{Cov} (x^{m+1}, v^{m+1}) &= \frac{\gamma}{2} \left[ 1 + e^{-4h} - 2e^{-2h} \right] I_d .
\end{align*}$$  

(11)

Here $E$ denotes the expectation, and $\text{Cov}(a, b)$ denotes the covariance of $a$ and $b$ (abbreviated to $\text{Cov}(a)$ when $b = a$), and $I_d$ is the identity matrix in $\mathbb{R}^d$. We thus draw $x^{m+1}, v^{m+1}$ from this Gaussian distribution numerically to update the iteration. We summarize ULMC in Algorithm

The algorithm converges exponentially when $f$ is strongly convex with Lipschitz continuous gradient; see Dalalyan and Riou-Durand [2018]. The original statement uses a different parametrization. We translate the result to the current one in Supp. and restate the result here.

**Theorem 4.1.** [Dalalyan and Riou-Durand, 2018, Theorem 2] Assume $f$ satisfies Assumption 3.1 and
We denote the discrete distribution from which \( r \) is chosen by \( \Phi \), with \( \phi_i \) being the probability of component \( i \) being chosen, that is,

\[
\Phi := \{ \phi_1, \phi_2, \ldots, \phi_d \},
\]

where \( \phi_i > 0 \) for all \( i \) and \( \sum_{i=1}^{d} \phi_i = 1 \). Denoting by \( h_i \) the stepsize when \( i \)-th direction is chosen, we choose \( h_i \) to be inversely proportional to \( \phi_i \), as follows:

\[
h_i = \frac{h}{\phi_i}, \quad i = 1, 2, \ldots, d,
\]

where \( h > 0 \) is a parameter that can be viewed as the expected stepsize. We also define the total elapsed time after \( m \) steps as

\[
T_m = \sum_{n=0}^{m-1} h_r^n.
\]

The initial iterate \((x^0, v^0)\) is drawn from a distribution \( q_0 \), which can be any distribution that is easy to draw from (e.g., a normal distribution).

Because only component \( r^m \) is updated at iteration \( m \), we have for \( t \in [T_m, T_{m+1}] \) that

\[
X_r(t) = X_r(T_m) + \frac{1 - e^{-2t}}{2} V_r(T_m)
\]

\[
- \gamma^2 \int_{T_m}^{t} (1 - e^{-2(t-s)}) \partial_{r^m} f(X(s)) \, ds
\]

\[
+ \sqrt{\gamma^2} \int_{T_m}^{t} (1 - e^{-2(t-s)}) \, dB_s,
\]

(15)

\[
V_r(t) = V_r(T_m) e^{-2(t-T_m)}
\]

\[
- \gamma^2 \int_{T_m}^{t} e^{-2(t-s)} \partial_{r^m} f(X(s)) \, ds
\]

\[
+ \sqrt{\gamma^2} \int_{T_m}^{t} e^{-2(t-s)} \, dB_s,
\]

(16)

\[
X_i(t) = X_i(T_m), \quad V_i(t) = V_i(T_m), \quad i \neq r^m.
\]

(17)

We call \([15,17]\) Random Coordinate Underdamped Langevin dynamics (RC-ULD). To obtain a practical algorithm, we apply the Euler approximation to these dynamics. Denoting by \((x_{r^m}^m, v_{r^m}^m)\) the numerical approximation to \((X(T_m), V(T_m))\), we replace \( \partial_{r^m} f(X(s)) \) in \([15,17]\) by \( \partial_{r^m} f(x^m) \), so that \( x_{r^m}^{m+1} = x_{r^m}^m, \quad v_{r^m}^{m+1} = v_{r^m}^m \), for \( i \neq r^m \), and \((x_{r^m}^{m+1}, v_{r^m}^{m+1})\) are two Gaussian random variables with the following expectation and covariance:

\[
E x_{r^m}^{m+1} = x_{r^m}^m + \frac{1}{2} \left( 1 - e^{-2h_{r^m}} \right) v_{r^m}^m
\]

\[
- \frac{\gamma}{2} \left( h_{r^m} - \frac{1}{2} (1 - e^{-2h_{r^m}}) \right) \partial_{r^m} f(x^m),
\]

(18)

Then, \((x_{r^m}^{m+1}, v_{r^m}^{m+1})\) is drawn according to this Gaussian distribution for the update. We summarize the RC-ULMC approach in Algorithm 2.

6 Main results

We have three main results regarding the underlying dynamics (RC-ULD), and the RC-ULMC algorithm.
Algorithm 2 Random Coordinate Underdamped Langevin Monte Carlo (RC-ULMC)

Input: $h$ (time stepsize); $\gamma$ (parameter); $d$ (dimension); $\nabla f(x)$; probability set $\Phi := \{\phi_1, \phi_2, \ldots, \phi_d\}$; $M$ (stopping index).

Initial: $(x^0, v^0)$ i.i.d. sampled from the initial distribution induced by $q_0(x, v)$. For $m = 0, 1, \ldots, M$ do
1. Draw $r$ randomly from 1, 2, $\ldots, d$ according to $\Phi$.
2. Update $(x^{m+1}, v^{m+1})$ as follows:
   - $x_i^{m+1} = x_i^m, v_i^{m+1} = v_i^m$ for $i \neq r$;
   - sample $(x_r^{m+1}, v_r^{m+1})$ as Gaussian variables according to (18).
end for
Output: $\{x^m\}$.

In Section 6.1, we discuss convergence of the RC-ULD (15)-(17). This SDE can be viewed as the continuum version of the RC-ULMC algorithm. Only with the convergence of this SDE can we hope for the convergence of RC-ULMC. In Section 6.2, we describe the non-asymptotic convergence properties of RC-ULMC. From this result, we can determine an optimal strategy for selecting the coordinate $r^m$ at each iteration. We will also compare our results with those for classical ULMC, showing that the bounds for RC-ULMC are always better. Moreover, when $f$ is highly skewed — for example when $\kappa_1 = \kappa_{\text{max}} \gg 1$ and $\kappa_i \approx 1$ for $i \geq 2$ — the total cost is $\tilde{O}(d^{3/2} + \kappa_{\text{max}}) \kappa^{1/2}/\epsilon$, as compared to $\tilde{O}(d^{3/2} \kappa^{3/2}/\epsilon)$ for ULMC.

We provide an example in Section 6.3 to show that our bounds are tight.

6.1 Convergence of RC-ULD

Our first result is on the convergence of (15)-(17), the RC-ULD that incorporates random coordinate selection.

Denote $X^m = X(T^m)$, $V^m = V(T^m)$ and denote the probability filtration by $\mathcal{F}^m = \{x^0, v^0, s \leq m, B_s \leq T^m\}$. Then we have the following result about its geometric ergodicity.

**Theorem 6.1.** Suppose that $f$ satisfies Assumption 3.1 and that
$$\gamma \leq \frac{1}{L}, \quad h \leq \frac{\gamma \mu \min\{\phi_i\}}{240}. \quad (20)$$

Denote $q_m(x, v)$ the probability density function of iteration $m$ of RC-ULMC and define $W_m = W_2(q_m, p)$. Then we have
$$W_m \leq 4 \exp \left(-\frac{\mu \gamma m h}{8}\right) W_0 + 40 \gamma^{1/2} h \sqrt{\sum_{i=1}^{d} \frac{\kappa_i^2}{\phi_i^2}}. \quad (21)$$

The proof can be found in Supp. 3. This result indicates that the Wasserstein distance between $q_m$ and the target distribution decays exponentially except for an error term of size $O(h)$. The convergence rate is given by $\mu \gamma$, and with the choice $\gamma = 1/L$, this quantity is the inverse condition number $1/\kappa$ of the objective function (see (5)). The second term in (21) reflects...
the discretization error, with its size being determined by the directional condition number $\kappa_i$ (see (5)) and the random selection probability distribution $\Phi$.

This theorem not only allows us to estimate the number of iterations required to achieve a preset accuracy, but also suggests that we choose $\{\phi_i\}$ in a way that minimizes the bound.

**Corollary 6.3.** Suppose that the conditions of Theorem 6.2 hold and $\gamma = 1/L$. We have the following estimates.

- For any $\epsilon > 0$, the number of needed iterations $M$ to attain $W_M \leq \epsilon$ is
  \[
  M = \Theta \left( \frac{\epsilon}{\mu} \frac{1}{\kappa} \sqrt{\frac{d}{\sum_{i=1}^{d} \kappa_i^2 / \phi_i^2}} \log \left( \frac{W_0}{\epsilon} \right) \right). \tag{22}
  \]

- The optimal choice of $\phi_i$ is
  \[
  \phi_i = \frac{L_j^{2/3}}{\sum_{j=1}^{d} L_j^{2/3}}, \quad i = 1, 2, \ldots, d. \tag{23}
  \]

In this case, the number of iterations required is
  \[
  M = \Theta \left( \frac{\epsilon}{\mu} \frac{1}{\kappa} \sqrt{\frac{d}{\sum_{i=1}^{d} \kappa_i^2 / \phi_i}} \log \left( \frac{W_0}{\epsilon} \right) \right). \tag{24}
  \]

The proof can be found in Supp. 5. Suppose the objective function $f$ is well-conditioned in every direction, so that $\kappa_i = O(1)$ for all $i$. Then according to both (22) and (24), we see that the cost is roughly $O(d^{3/2}/\epsilon)$. This order is the same as for the classical ULMC shown in Theorem 4.1.

When $f$ is not as well-conditioned, meaning $\kappa_i$ are not uniformly small in every direction, then RC-ULMC can have a significant advantage over the classical ULMC. In practice, if we have some a priori estimate of $\kappa_i$, we can choose the optimal $\Phi$ and the cost estimate will be given by (24). Of course, such a priori information might not be available, in such case, we can choose uniformly the coordinate at each iteration: $\phi_i = 1/d$. We compare below the cost of RC-ULMC in these two scenarios with $O(d^{1/2}/\sqrt{\kappa}/\epsilon^{2/3})$, the cost of the classical ULMC, as shown in Theorem 4.1.

Case 1: Uniform sampling, where we choose $\phi_i = 1/d$. From (22), we found that the cost is
  \[
  \tilde{O} \left( \frac{d^{1/2}}{\mu^{1/2}} \sqrt{\sum_{i=1}^{d} \kappa_i^2} \right), \tag{25}
  \]
  where the $\tilde{O}$ ignores log terms. Since $\sum_{i=1}^{d} \kappa_i^2 \leq d \kappa_{\text{max}}^2 \leq d \kappa^2$, we observe that RC-ULMC is always cheaper than ULMC. Furthermore, when $f$ is highly skewed (7) with $p = 2$, then (25) is reduced to $\tilde{O} \left( \frac{d(\sqrt{3} + \kappa_{\text{max}}) \kappa^{1/2}}{\mu^{1/2}} \right)$. This bound indicates that RC-ULMC is significantly cheaper than ULMC when both $d$ and $\kappa$ are large.

Case 2: With the optimal choice of $\Phi$ (using (23)), the cost of RC-ULMC is equivalent to $M$ in (24). We still have that RC-ULMC is always cheaper than ULMC. Furthermore, when $f$ is highly skewed (7) with $p = 3/2$, then we have upper bound $\left( \sum_{j=1}^{d} \kappa_j^{2/3} \right)^{3/2} \approx (d - 1) + \kappa_{\text{max}}^{3/2} \leq 2(d^{3/2} + \kappa_{\text{max}})$. By substituting this bound into (24), we get the following bound:
  \[
  \tilde{O} \left( \frac{(d^{3/2} + \kappa_{\text{max}}) \kappa^{1/2}}{\mu^{1/2}} \right). \tag{26}
  \]

The reduction over ULMC is significant when either $d$ or $\kappa$ is large.

We also compare RC-ULMC with RC-OLMC discussed in [Shen et al. 2019, Ding et al. 2020]. To achieve $\epsilon$-accuracy, the total cost of RC-OLMC is $\tilde{O} \left( \frac{\sum_{i=1}^{d} \kappa_i^2 / \phi_i}{\mu^2} \right)$. Noting that $\sqrt{\sum_{i=1}^{d} \kappa_i^2 / \phi_i} \leq \sum_{i=1}^{d} \kappa_i / \phi_i \leq \sum_{i=1}^{d} \kappa_i^2 / \phi_i$, (22) is always smaller, meaning RC-ULMC is always cheaper than RC-OLMC when $\epsilon < 1/L^{1/2}$. Furthermore, if we choose uniform sampling ($\phi_i = 1/d$) and assume $\gamma, \mu, \kappa$ are $O(1)$, then the cost of RC-ULMC is $\tilde{O}(d^{3/2}/\epsilon)$ while the cost of RC-OLMC is $\tilde{O}(d^{5/2}/\epsilon)$.

Finally, we note that in [Shen and Lee 2019], randomized midpoint method (RMM) is used to discretize SDE (8). According to [He et al. 2020], RMM needs $O(d^{3/2}/(\kappa^{2/3}))$ iteration steps to achieve $\epsilon$-accuracy, which equates to a cost of $O(d^{3/2}/(\kappa^{2/3}))$. By comparison, (26) is smaller in some extreme regimes, such as $d^{3/2} \kappa^{1/2} \gamma < \kappa < \epsilon^{2/3} d^{3/2}$. We note that the comparison between RC-ULMC and RMM is not entirely fair since RMM uses a better discretization scheme than the Euler approximation (11) used in RC-ULMC. It is possible to include the RCD idea to RMM on ULMC as well, for a potentially better convergence rate. We leave that topic to future investigation.

### 6.3 Tightness of the bound

Corollary 6.3 shows that the numerical cost is roughly $O(d^{3/2}/\epsilon)$ when the problem is well conditioned. We show by use of an example that this bound is tight with respect to $d$ and $\epsilon$. 
Proposition 6.4. Let the target distribution be a standard Gaussian \( p(x) = \frac{1}{(2\pi)^{d/2}} \exp(-|x|^2/2) \), which is the marginal distribution of the target distribution \( p(x, v) = \frac{1}{(2\pi)^{d/2}} \exp(-|x|^2/2 - |v|^2/2) \). Suppose the initial distribution \( q_0 \) is chosen to be \( q_0(x, v) = \frac{1}{(2\pi)^{d/2}} \exp(-|x - u|^2/2 - |v|^2/2) \) with \( u \in \mathbb{R}^d \) and \( u_i = 1/400 \) for all \( i \). Then if we choose \( h \) so that \( h < 10^{-8}/d \), we have

\[
W_m \geq \frac{\exp(-4hm) d}{800^2} \frac{d^{3/2}h}{8} + \frac{320 - 464dh}{8},
\]

where \( W_m := W_2(q_m, p) \), and \( q_m \) is the probability distribution of \((x^m, v^m)\) generated by Algorithm 3 with \( \gamma = 1/L = 1 \) using uniform coordinate sampling. Furthermore, to have \( W_m \leq \epsilon \), one needs at least \( M = O(d^{3/2}/\epsilon) \) iterations.

The proof can be found in Supp. [2]. For this particular initialization, we have \( W_0 = \sqrt{d}/400 \). According to (27), we can guarantee \( W_m \leq \epsilon \) only if both terms are smaller than \( \epsilon \), meaning that (ignoring a log term) \( h \lesssim \frac{320}{d^2 \epsilon} \), \( m \gtrsim \frac{1}{4h} \), which implies a cost of \( O(d^{3/2}/\epsilon) \).

7 Numerical experiments

We present numerical evidence to demonstrate the improvement of RC-ULMC over the classical ULMC. In this section we only present a toy problem, and we leave a more practical example to Supp. [2].

In the examples, we repeat the Markov chain for \( N \) independent trials and denote \( \{x^{(i)}, M\}_{i=1}^N \) the list of \( N \) samples at \( M \)-th iteration. Since Wasserstein distance is difficult to measure directly numerically, especially when the underlying distribution function is presented by a list of particles, we evaluate the following error as a surrogate:

\[
\text{Error} = \frac{1}{N} \left\| \frac{1}{N} \sum_{i=1}^N \psi(x^{(i)}, M) - E_{p_X}(\psi) \right\|_2,
\]

where \( \psi(x) \) is a matrix-valued function and referred to as the test function, \( \| \cdot \|_2 \) means the spectral norm of the matrix, and \( E_{p_X}(\psi) \) is the expected value of \( \psi \) with respect to the target distribution \( p_X \).

In the first example, we set the target distribution function to be \( p_X(x) \propto p_1(x)p_2(x) \) with \( p_2(x) = \exp\left(-\sum_{i=1}^d |x_i|^2/2\right) \) and \( p_1(x) = \exp(-\frac{1}{2}x^\top \Gamma^\top \Gamma x) \), where \( x = (x_1, x_2, \ldots, x_{10})^\top \) is the list of first 10 entries, and \( \Gamma = T + \frac{d}{M} I \). Here \( I \) is the 10 \times 10 identity matrix and \( T \) is a random matrix whose entries are i.i.d. standard Gaussian random variables. This example has an ill-conditioned \( f \). The Lipschitz constants are \( E(L_{1 \leq i \leq 10}) = \frac{d^2}{100} \) and \( L_{i \geq 10} = 1 \). Thus \( \mu = 1 \), and \( E(k_{1 \leq i \leq 10}) = \frac{d^2}{100} \). When \( d \gg 1 \), we have \( \sum_{j=1}^d k_{i} \ll d \kappa_p \) for \( p \geq 1 \).

In the simulation we set \( d = 100, N = 10^5 \), and let \( \psi(x) = xx^\top \in \mathbb{R}^{10 \times 10} \).

Initially, all particles are drawn from the density distribution \( p_0(x, v) \propto p_1(x - 0.5e_{10})p_2(x) \exp(-|v|^2/2) \), where \( e_{10} \) is a vector in \( \mathbb{R}^{10} \) and all entries equal to 1. It is expected that the density of the target distribution is \( p_X(x) \exp(-|v|^2/2) \), making \( p_X \) the marginal probability density.

The result is plotted in Figure 1. To run RC-ULMC, we use time stepsize \( h = 10^{-4} \). For comparison we also run ULMC, however, due to the cost difference per iteration, there is no standard choice of \( h \) for ULMC for a fair comparison. Since \( d = 100 \) in this example, per iteration, the cost of ULMC is 100 times of that of RC-ULMC, we first experiment ULMC with \( h = 10^{-2} \). It is clear that RC-ULMC, presented by the purple line achieves a lower error than ULMC with the same amount of cost.

We then test ULMC with different choices of \( h \), hoping to find its best performance. As one decreases \( h \), the error plateau is also lower, but the decay rate of error with respect to the cost decreases, as one can see by comparing the yellow, red and blue lines in Figure 1 all produced by ULMC with different values of \( h \). None of them, however, can compete with RC-ULMC regarding the level of error at the same cost.

![Figure 1: The decay of error with respect to the cost (the number of \( \partial f \) calculations).](image)

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