
Learnable Sampler Distillation for Discrete Diffusion Models

Feiyang Fu, Tongxian Guo, Zhaoqiang Liu*
University of Electronic Science and Technology of China

Abstract

Discrete diffusion models (DDMs) have shown powerful generation ability for discrete data modalities like text and molecules. However, their practical application is hindered by inefficient sampling, requiring a large number of sampling steps. Accelerating DDMs by using larger step sizes typically introduces significant problems in generation quality, as it amplifies the impact of both the compounding decoding error due to factorized predictions and discretization error from numerical approximations, leading to a significant decrease in sampling quality. To address these challenges, we propose learnable sampler distillation (LSD), a novel approach to train fast and high-fidelity samplers for DDMs. LSD employs a distillation approach where a student sampler with a few steps learns to align its intermediate score trajectory with that of a high-quality teacher sampler with numerous steps. This alignment is achieved by optimizing learnable sampler coefficients that adaptively adjust sampling dynamics. Additionally, we further propose LSD+, which also learns time schedules that allocate steps non-uniformly. Experiments across text generation, image generation, and synthetic tasks demonstrate that our proposed approaches outperform existing samplers for DDMs, achieving substantially higher sampling quality with significantly fewer sampling steps. Our code is available at <https://github.com/feiyangfu/LSD>.

1 Introduction

Diffusion models have demonstrated remarkable success across various generative tasks, particularly excelling in the synthesis of data within continuous domains like images, audio, and videos [1, 2, 3, 4, 5]. These models frame the data generation process as a gradual denoising procedure in a continuous latent space. However, many other important data modalities, such as natural language, molecular sequences, and categorical data, inherently possess discrete structures. Applying diffusion models directly to these discrete spaces is challenging, as the standard formulation relies on continuous state transitions. Recently, discrete diffusion models (DDMs) [6, 7, 8, 9, 10, 11, 12] have been developed to address this issue. DDMs are specifically designed to operate on discrete data, adapting the core diffusion idea to categorical variables and enabling principled generation. Recent advances in DDMs have shown promising results, achieving competitive performance in generating high-fidelity discrete data. Despite their promising applicability, DDMs face an important challenge in sampling efficiency, and they typically require a substantial number of function evaluations (NFEs), e.g., 1024 or more, making inference computationally expensive.

Current sampling methods for DDMs are mainly divided into two categories: 1) Exact simulation methods [13, 14] provide unbiased samples from the target distribution but suffer from high sampling times and expensive computational costs due to numerous model evaluations, leading to poor scaling with dimensionality. 2) Approximate methods like τ -leaping [15, 16] are designed for parallelization

*Corresponding author.

and potentially faster sampling. However, such methods are first-order accurate and require small step sizes to ensure sampling quality.

Directly accelerating the sampling of DDMs through reducing NFEs typically produces unsatisfactory results, since this amplifies the impact of the compounding decoding error [17] and discretization error. Compounding decoding error arises since DDMs employ a factorized parameterization for computational efficiency, predicting the denoised state of each token independently, and ignoring inherent dependencies between tokens in the sequence. Consequently, the learned factorized denoising distribution differs from the true reversal process. This discrepancy is exacerbated when reducing NFEs, as the approximation quality degrades over larger intervals. Discretization error occurs since large step sizes make it inaccurate for numerical methods like Euler [18] and τ -leaping [15] to approximate the reverse dynamics. Moreover, these two errors accumulate over the sampling trajectory, severely degrading sampling quality when using small NFEs. Throughout the following, we call the combination of compounding decoding error and discretization error as accumulated error for brevity. To address the issue incurred by large accumulated error, we propose learnable sampler distillation (LSD) and its improved version for efficient sampling of DDMs.

1.1 Related Work

Efficient sampling in continuous diffusion models Recent efforts to accelerate sampling in continuous diffusion models largely focus on reducing the NFEs for solving the reverse-time ordinary differential equation (ODE) or stochastic differential equation (SDE) [19, 4, 20, 21, 22, 23, 24].

One major direction involves designing advanced ODE solvers. Some works [20, 25, 26, 27, 4] provide efficient sampling methods by establishing high-order numerical ODE solvers for continuous diffusion models.

There are also approaches that learn or optimize various components of the sampling process. AYS [28] seeks non-uniform time step schedules specific to given models and datasets, though their optimization can be computationally intensive. DMN [29] proposes a general framework for designing an optimization problem that seeks more appropriate time steps by minimizing the distance between the ground-truth solution to the ODE and an approximate solution corresponding to the numerical solver. AMED-Solver [30] learns adaptive mean estimation directions based on the observation that trajectories often reside in low-dimensional subspaces, which typically involves training an auxiliary network with high costs.

The most relevant works to us are perhaps LD3 [31] and S4S [32], both proposing learning diffusion model solvers via distillation in the continuous domain. LD3 efficiently learns the time discretization by backpropagating through the ODE-solving procedure using the proposed surrogate loss. S4S further learns the coefficients of the student solver by minimizing the distance between the final samples generated by the student and teacher solvers using learned non-uniform time schedules. However, these approaches face challenges when applied to DDMs. We highlight several distinctions in our learnable sampler distillation (LSD) approach (and its improved version) designed to address these challenges. 1) DDM sampling involves non-differentiable categorical sampling at each step, obstructing direct gradient flow from the final discrete output back to the sampler parameters. The reliance of S4S on final sample comparison is thus infeasible. We address this issue by aligning the intermediate score trajectories between the student and teacher samplers. This provides a viable path for gradient-based optimization of the learnable coefficients within the discrete sampling methods. 2) The work for S4S uses the final sample matching error to learn the time schedules, which may ignore dynamic changes of the accumulated error in the intermediate steps. We instead learn the time steps by aligning the effective transition term at intermediate stages during the reverse process. The effective transition term in the reverse process incorporates step sizes and concrete scores, which are tailored for DDMs. 3) The work for S4S optimizes the continuous initial noise using projected stochastic gradient descent (SGD) within an L_2 ball. This is inapplicable to DDMs where the initial state is often a discrete sequence, e.g., all masked tokens, which lacks a continuous gradient. To address this issue, we adapt the approach by measuring proximity using Hamming distance, which is suitable for discrete spaces and does not perform gradient updates on itself.

Distillation in Diffusion Models The distillation of continuous diffusion models is a rapidly advancing field. A prominent direction is related to the consistency model [33], which aims to learn a function that maps any point on an ODE trajectory to its origin, enabling one-step or few-

step generation. This paradigm has been extended to multi-step variants [34, 35] for improved performance. Other significant works focus on directly matching student and teacher distributions, such as distilling guided diffusion models [36], proposing simplified and faster matching objectives [37], recursively distilling a deterministic diffusion sampler into a new model [38], or concentrating on one-step distillation [39]. While these methods are highly effective for continuous models, they usually rely on continuous paths in the sense that the sampling process of each step is differentiable. Our work diverges by proposing a distillation framework specifically for the discrete diffusion model, which does not assume such a continuous path, and addressing a different set of challenges like the non-differentiability of the outputs. A recent work for Di[M]O [40] also involves distilling discrete diffusion models. It distills a multi-step masked diffusion model into a one-step generator. This is achieved by training a new student model from scratch, using a sophisticated proxy objective that involves creating "pseudo-intermediate states" and training an auxiliary model to match conditional output distributions. Our approaches are significantly different with Di[M]O in both goal and mechanism. Similarly to LD3 and S4S, we focus on a few-step sampler distillation. We tackle the challenge of non-differentiability of sampling from categorical distributions, and we enhance an existing sampler rather than replacing the model, which avoids the complexity of training a new generator and an auxiliary model.

Discrete diffusion models DDMs have emerged and undergone substantial development recently. SEDD [6] proposes score entropy, a novel loss that naturally extends score matching to discrete spaces and integrates seamlessly to build DDMs. RADD [9] reveals that the concrete score in absorbing diffusion can be expressed as conditional probabilities of clean data, multiplied by a time-dependent scalar in an analytic form and it unifies absorbing DDMs and any-order autoregressive models.

Various strategies have been proposed to accelerate the sampling of DDMs while maintaining quality. Among these, approximate simulation methods [18, 16, 15, 8] are widely used due to their potential for parallelization. A prominent example is the τ -leaping algorithm [8] that is adapted for DDMs. τ -leaping simulates the process by taking an approximate Euler-like step at each data dimension simultaneously and independently. Tweedie τ -leaping [6, 41] is an extension to τ -leaping and is proposed to improve accuracy by specifically considering how the rate matrix changes according to the noise schedule throughout the reverse process. While these τ -leaping variants offer the advantage of parallelization, the inherent approximation error still necessitates using many small steps to achieve high sampling quality.

The recent work for JYS [17] attempts to accelerate the sampling process of DDMs by focusing on optimizing the time steps of the sampling schedule. It minimizes a Kullback–Leibler divergence upper bound (KLUB) that implicitly captures the overall impact of the compounding decoding error and strategically allocates sampling steps. However, JYS operates by optimizing when to sample, rather than how to sample. At each chosen time, it still relies on the intrinsically biased model and employs standard large-step approximations that suffer from a significant discretization error.

1.2 Contributions

To address limitations mentioned above, we move beyond fixed or hand-tuned inference strategies. We introduce a novel learnable sampler distillation (LSD) approach specifically for DDMs. We employ a teacher sampler using small step sizes to approximate a high-quality trajectory. A student sampler is then trained with larger step sizes. Instead of mimicking only the final output of the teacher sampler, which is challenging due to non-differentiability in the discrete pipeline, the student sampler learns to align its intermediate score trajectory with that of the teacher sampler. This alignment is achieved by optimizing the learnable sampler coefficients, which provide the ability to adaptively adjust the sampling dynamics at each step and potentially compensate for the accumulated error given larger step sizes. Furthermore, we propose LSD+ that also learns sampling time schedules. This is done by comparing the effective transition term in the reverse process at intermediate stages and empirically works better compared with uniform sampling schedules used by LSD. We also utilize a relaxed objective during the learning process to alleviate the difficulty of hard alignment between the teacher and student samplers.

Overall, our contribution can be summarized as follows:

- We propose the LSD approach. Inspired by the insight of aligning intermediate score trajectories, LSD trains an efficient student sampler via distillation by optimizing learnable sampler coefficients and incorporating a relaxed training objective for improved feasibility.
- We further introduce LSD+, an extension to LSD that additionally learns non-uniform time schedules. This allows for adaptive allocation of sampling steps, offering a mechanism to potentially better capture varying dynamics and further reduce accumulated errors compared to using uniform time schedules.
- Extensive experiments across text generation, image generation, and synthetic data tasks demonstrate that our proposed approaches achieve significantly higher sampling quality compared to existing baselines at reduced NFEs.

2 Preliminaries

2.1 Continuous time discrete diffusion models

DDMs model the generative process that can be expressed as a continuous time Markov chain (CTMC) on a finite state space $\mathcal{X} = \{1, \dots, N\}$ [8, 42]. The forward process describes how data is corrupted. Specifically, the probability of transitioning from state x at time t to state y after a small time interval Δt is denoted by $p_{t+\Delta t|t}(y|x)$. This is characterized by [9]:

$$p_{t+\Delta t|t}(y|x) = \begin{cases} Q_t(x, y)\Delta t + o(\Delta t), & y \neq x, \\ 1 + Q_t(x, x)\Delta t + o(\Delta t), & y = x, \end{cases} \quad (1)$$

where $Q_t(x, y)$ is the (x, y) element of the transition rate matrix Q_t . The transition rate matrix Q_t is usually formed as $\sigma(t)Q$ [8], where $\sigma(t)$ is a scalar factor, Q is a pre-defined standard matrix with special structures [8]. Let p_t denote the marginal distribution of states at time t . In particular, $p_0 = p_{\text{data}}$ is the true distribution of the data. Additionally, for the terminal time T , p_T approaches a distribution π . Depending on Q , π can mainly be modeled as two distributions, namely a uniform distribution or a distribution that converts samples into masked tokens. For the reverse process that transfers p_T back to p_0 , the inverse CTMC can be characterized as follows [9]:

$$p_{t-\Delta t|t}(y|x) = \begin{cases} \tilde{Q}_t(x, y)\Delta t + o(\Delta t), & y \neq x, \\ 1 + \tilde{Q}_t(x, x)\Delta t + o(\Delta t), & y = x, \end{cases} \quad (2)$$

where $\tilde{Q}_t$ is the reverse transition rate matrix [41], which can be parameterized by:

$$\tilde{Q}(x, y) = \begin{cases} \frac{p_t(y)}{p_t(x)} Q_t(y, x), & y \neq x, \\ -\sum_{z \neq x} \tilde{Q}_t(x, z), & y = x. \end{cases} \quad (3)$$

The concrete score term $\frac{p_t(y)}{p_t(x)}$ needs to be estimated, as p_t is generally unknown. Therefore, the goal of training a score network $s_\theta : \mathcal{X} \times \mathbb{R} \rightarrow \mathbb{R}^{|\mathcal{X}|}$ is to approximate these score values. For instance, SEDD [6] provides an effective method that learns s_θ such that it satisfies $s_\theta(x, t) \approx \left[\frac{p_t(y)}{p_t(x)} \right]_{y \neq x}$.

3 Methods

Achieving efficient inference in DDMs with high sampling quality necessitates accurate approximation of the reverse CTMC using significantly fewer steps than traditional high-fidelity samplers. Numerical samplers like the Euler sampler approximate this process by taking discrete steps guided by the concrete score from the model. However, increasing the step size for faster inference alters the discrete transition dynamics, leading to increased accumulated errors and degraded sampling quality. To enable accurate sampling with large step sizes, we propose learnable sampler distillation (LSD) to make some components of the numerical sampler learnable. Specifically, LSD employs learnable coefficients to dynamically adjust the influence of the concrete score at each time step, allowing the sampler to compensate for large-step discretization errors. Furthermore, while LSD could lead to significant improvement in the generation quality, we further propose LSD+. Instead of learning the coefficients using a fixed uniform schedule, LSD+ further learns a sequence of non-uniform time steps. By training these parameters through distillation from a high-quality teacher sampler, our method learns an optimized discrete-time trajectory. Figure 1 shows the pipeline of our method, and the details of the method are described in the following subsections.

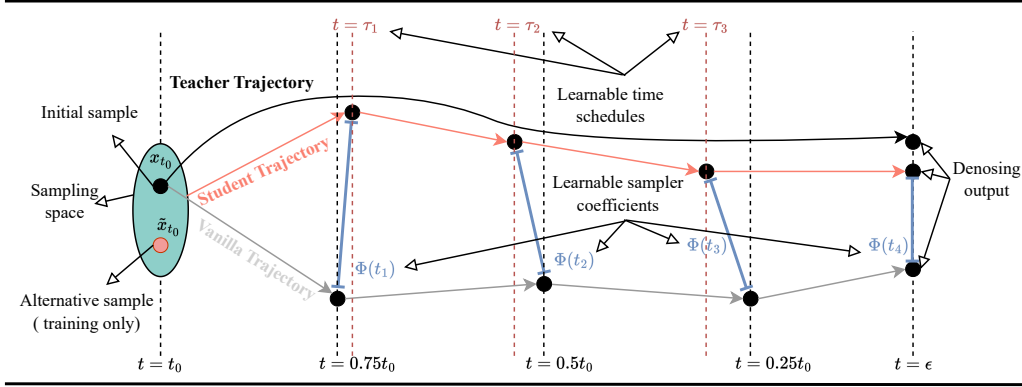


Figure 1: Illustration of our LSD+ approach. During training, the student sampler starts from an alternative initial sample $\tilde{x}_{t_0}$ within the sample space (close to the original x_{t_0}). During inference, sampling starts from the initial sample x_{t_0} . We can see that the vanilla sampling trajectory often introduces significant discretization errors with large step sizes. In contrast, LSD+ employs learnable coefficients $\Phi(t_k)$ and learnable time schedules τ_k to adaptively adjust its trajectory. This enables the LSD sampler to more accurately mimic the trajectory of the teacher sampler, effectively compensating for errors inherent in accelerated sampling.

3.1 LSD: Coefficients

In this subsection, we illustrate our LSD approach that learns time-dependent coefficients for accelerating DDMs. Given a pre-trained score network $s_\theta(\cdot, \cdot)$, transition rate matrix Q_t , and an initial state x_T sampled from p_T at the initial time T , the reverse sampling process for DDMs generates a sample by iteratively applying an update rule. For an Euler-type sampler, the transition probability for the i -th token from its current state x_t at time t to the next state $x_{t-\Delta t}$ at time $t - \Delta t$ can be parameterized as:

$$p(x_{t-\Delta t}^i | x_t^i) = \delta_{x_t^i}(x_{t-\Delta t}^i) + \Delta t Q_t(x_t^i, x_{t-\Delta t}^i) s_\theta(x_t, t)_{i, x_{t-\Delta t}^i}. \quad (4)$$

Here, x_t^i denotes the i -th token of the current state sequence x_t , $\delta_{x_t^i}(x_{t-\Delta t}^i)$ is the Kronecker delta function, Δt represents the time step size, $Q_t(x_t^i, x_{t-\Delta t}^i)$ denotes the $(x_t^i, x_{t-\Delta t}^i)$ element of the transition rate matrix Q_t , and $s_\theta(x_t, t)_{i, x_{t-\Delta t}^i}$ is the $(i, x_{t-\Delta t}^i)$ element of the concrete score $s_\theta(x_t, t)$.

We apply a fixed teacher sampler that approximates the true reverse process with high fidelity using time schedules $\{t_j^*\}_{j=0}^N$ comprising N steps, with $T = t_0^* > t_1^* > \dots > t_N^* = \epsilon > 0$.² The state generated by the teacher sampler at time t_j^* along its trajectory is denoted as $x_{t_j^*}$. The sampling process of a teacher sampler Ψ^* yields a high-quality final sample $x_\epsilon^* = \Psi^*(x_T, \{t_j^*\}_{j=0}^N, s_\theta, \{Q_{t_j^*}\}_{j=0}^N)$, which is abbreviated as $x_\epsilon^* = \Psi^*(x_T)$ for simplicity in notation.

We apply a student sampler that operates with a time schedule $\{t_k\}_{k=0}^M$ comprising M steps, where $M \ll N$ and $T = t_0 > t_1 > \dots > t_M = \epsilon > 0$. $\{t_k\}_{k=0}^M$ is a subsequence of $\{t_j^*\}_{j=0}^N$. Our goal is to learn a set of time-dependent coefficients $\Phi = \{\Phi(t_k)\}_{k=1}^M$ to improve the quality of the output of the student sampler.³ The state generated by the student sampler at time t_k along its trajectory is denoted as x_{t_k} . The sampling process of a student sampler Ψ yields a final sample $x_\epsilon = \Psi(x_T, \{t_k\}_{k=0}^M, s_\theta, \{Q_{t_k}\}_{k=0}^M, \{\Phi(t_k)\}_{k=1}^M)$, which is abbreviated as $x_\epsilon = \Psi_\Phi(x_T)$ to highlight the dependence on the coefficients $\{\Phi(t_k)\}_{k=1}^M$.

The update rule for the i -th token within the student sampler incorporating Φ becomes:

$$p(x_{t_{k+1}}^i | x_{t_k}^i) = \delta_{x_{t_k}^i}(x_{t_{k+1}}^i) + \Delta t Q_{t_k}(x_{t_k}^i, x_{t_{k+1}}^i) (\Phi(t_k) s_\theta(x_{t_k}, t_k))_{i, x_{t_{k+1}}^i}. \quad (5)$$

Similar to strategies in some learning methods for continuous ODE solvers [32], a direct objective is to minimize the distance $d(x_\epsilon, x_\epsilon^*)$, where $d(\cdot, \cdot)$ is a certain distance metric. However, it is

²Here, ϵ is a positive value close to 0 to avoid stability issues as discussed in [4].

³Here, $\Phi(t_0)$ is fixed to 1.

generally infeasible for DDMs to directly minimize $d(x_\epsilon, x_\epsilon^*)$ since the non-differentiable categorical sampling at each step obstructs gradient propagation. Instead, we propose to align intermediate score predictions. At each time step t_k , the student sampler computes its score $s_k = s_\theta(x_{t_k}, t_k)$. The teacher sampler evolves its state to the same time step t_k (i.e., for certain j such that $t_j^* = t_k$) using its more accurate sampling process and caches its score $s_k^* = s_\theta(x_{t_k}^*, t_k)$. The states x_{t_k} and $x_{t_k}^*$ differ due to the distinct sampling paths taken to reach t_k . Then, our objective is to minimize the discrepancy between s_k^* and $\Phi(t_k)s_k$ for all $k \in \{1, 2, \dots, M\}$. This can be expressed as:

$$\mathcal{L}_k(\Phi(t_k)) = \mathbb{E}_{x_{t_0} \sim \pi} [d(s_k^*, \Phi(t_k)s_k)]. \quad (6)$$

This intermediate score trajectory alignment provides a differentiable path for optimizing $\{\Phi(t_k)\}_{k=1}^M$ and ensures the student sampler mimics the trajectory of the teacher sampler across the full denoising path, not just at the final output. We present details of the sampling and training processes for LSD in Algorithms 1 and 2 respectively.

3.2 LSD+: Coefficients with learnable time schedules

While LSD improves the sampler by learning the sequence of coefficients $\{\Phi(t_k)\}_{k=1}^M$ under a fixed time schedule, the reverse diffusion dynamics vary significantly across time. We additionally propose LSD+ to also learn non-uniform time schedules. The intuition is that, by learning from a high-fidelity teacher sampler, the student sampler implicitly learns to allocate its limited steps in a manner that best approximates the trajectory of the teacher. Specifically, given time steps for the student sampler $\{t_k\}_{k=0}^M$, the uniform time schedule uses a step size at $\Delta t = \frac{T-\epsilon}{M}$. Our goal is to learn customized step sizes $\{\kappa_k\}_{k=1}^M$, which are initialized as Δt . The learnable time steps are calculated by:

$$\tau_k = T - \sum_{\ell=1}^k \kappa_\ell. \quad (7)$$

At each learned time step τ_k , the student sampler computes its score $s_\theta(x_{\tau_k}, \tau_k)$. To learn the step size κ_k , we utilize the so-called effective transition term in the reverse process. Specifically, for the student sampler, this is proportional to $\kappa_k s_\theta(x_{\tau_k}, \tau_k)$, for the teacher sampler, this is proportional to $\frac{T-\epsilon}{N} s_\theta(x_{t_k}^*, t_k)$, where $\frac{T-\epsilon}{N}$ is the step size for the teacher sampler and $s_\theta(x_{t_k}^*, t_k)$ is the cached score of teacher sampler. By calculating the distance of the effective transition terms between the student sampler and teacher sampler, we effectively update κ_k considering the unique characteristics of DDMs. This allows the time schedule to adaptively allocate step sizes based on the specific transition structures in DDMs. The updating process can be parameterized as follows:

$$\tilde{\mathcal{L}}_k(\kappa_k) = \mathbb{E}_{x_{t_0} \sim \pi} \left[d \left(\kappa_k s_\theta(x_{\tau_k}, \tau_k), \frac{T-\epsilon}{N} s_\theta(x_{t_k}^*, t_k) \right) \right]. \quad (8)$$

We present details of the training and sampling processes for LSD+ in the supplementary material.

Algorithm 1 Sampling process of LSD

Require: Score network s_θ , time schedule $\{t_k\}_{k=0}^M$ for the student sampler, learned coefficients of the student sampler $\{\Phi(t_k)\}_{k=1}^M$, transition rate matrices $\{Q_{t_k}\}_{k=0}^M$

- 1: Sample $x_{t_0} \sim \pi$
- 2: **for** $k = 0$ to $M - 1$ **do**
- 3: Sample $x_{t_{k+1}}$ based on x_{t_k} and $\Phi(t_k)$:
- 4: $p(x_{t_{k+1}}^i | x_{t_k}^i) = \delta_{x_{t_k}^i}(x_{t_{k+1}}^i) + (t_k - t_{k+1}) Q_{t_k}(x_{t_k}^i, x_{t_{k+1}}^i) (\Phi(t_k)s_\theta(x_{t_k}, t_k))_{i, x_{t_{k+1}}^i}$
- 5: $x_{t_{k+1}}^i \sim p(x_{t_{k+1}}^i | x_{t_k}^i)$ for all i
- 6: **end for**
- 7: **return** x_ϵ

Algorithm 2 Training process of LSD

Require: Score network s_θ , frozen teacher sampler Ψ^* with N steps, learnable student sampler Ψ_Φ with M steps, learning rate η , distance metric d , time schedule $\{t_j^*\}_{j=0}^N$ for the teacher sampler, time schedule $\{t_k\}_{k=0}^M$ for the student sampler (a subsequence of $\{t_j^*\}_{j=0}^N$), transition rate matrices $\{Q_{t_j^*}\}_{j=0}^N$

- 1: Initialize $\Phi(t_k) = 1$ for $k = 1, \dots, M$
- 2: **while** not converged **do**
- 3: Sample $x_{t_0} \sim \pi$, set $x_{t_0}^* \leftarrow x_{t_0}$
- 4: **for** $k = 1$ to M **do**
- 5: Calculate the state x_{t_k} generated by the student sampler at time t_k and calculate the score $s_k = s_\theta(x_{t_k}, t_k)$
- 6: Calculate the state $x_{t_k}^*$ generated by the teacher sampler at time t_k and calculate the score $s_k^* = s_\theta(x_{t_k}^*, t_k)$
- 7: **end for**
- 8: **for** $k = 1$ to M **do**
- 9: $L_k \leftarrow d(\Phi(t_k)s_k, s_k^*)$
- 10: $\Phi(t_k) \leftarrow \Phi(t_k) - \eta \nabla_{\Phi(t_k)} L_k$
- 11: **end for**
- 12: **end while**
- 13: **return** $\{\Phi(t_k)\}_{k=1}^M$

3.3 Relaxed objective

For a student sampler which typically has lower NFEs compared to a teacher sampler, it is non-trivial to force it to accurately match the output of the teacher sampler given the same initial input x_{t_0} . Thus, we adopt a relaxed training objective for both LSD and LSD+. We take LSD as an example for further presentation. Instead of strictly requiring the score of the student sampler $s_\theta(x_{t_0}, t_0)$ to match the score of the teacher sampler $s_\theta(x_{t_0}^*, t_0)$, we only require that there exists an alternative input $\tilde{x}_{t_0}$ sufficiently close to the original x_{t_0} (within a small Hamming distance [43] in our discrete token space). Specifically, $\tilde{x}_{t_0}$ satisfies:

$$d_H(x_{t_0}, \tilde{x}_{t_0}) \leq \zeta, \quad (9)$$

where $d_H(\cdot, \cdot)$ denotes the Hamming distance between two sequences, ζ represents positive integer threshold that defines the maximum allowed Hamming distance between $\tilde{x}_{t_0}$ and x_{t_0} , where we set it as around 5% of the sequence length.

Therefore, the output of the student sampler at this perturbed score should approximately match the score of the teacher sampler at the original input such that $s_\theta(\tilde{x}_{t_0}, t_0) \approx s_\theta(x_{t_0}^*, t_0)$. Moreover, the relaxed objective function for LSD can be expressed as:

$$\mathcal{L}_{\text{relaxed},k}(\Phi(t_k)) = \mathbb{E}_{x_{t_0}, \tilde{x}_{t_0}} [d(s_\theta(x_{t_k}^*, t_k), \Phi(t_k)s_\theta(\tilde{x}_{t_k}, t_k))], \quad (10)$$

where $\tilde{x}_{t_0}$ and x_{t_0} satisfies Eq. (9) and $\tilde{x}_{t_k}$ is sampled starting from $\tilde{x}_{t_0}$. This relaxation makes the optimization task more feasible for the capacity-constrained student sampler by alleviating the rigorous matching requirement. Notably, this input perturbation $\tilde{x}_{t_0}$ is only used during training, at inference time, the student sampler receives the original and unperturbed input x_{t_0} . We provide further discussion on the reasonableness of the relaxed objective in the supplementary material.

4 Experiments

In this section, we empirically evaluate the performance of our proposed LSD approach and its improved version LSD+. Our goal is to validate their ability to generate high-quality samples at low NFEs. We conduct evaluations across diverse settings, including text generation, image generation, and a synthetic sequence task, comparing against various baselines. We highlight that our LSD+ provides an efficient learning process for the coefficients and time schedules, typically requiring 5 minutes on an NVIDIA RTX4090 GPU, compared to around 10 minutes of training time for JYS under the same environment. And the learned student sampler introduces no additional computational burden during sampling.

4.1 Text generation

For the text generation task, we employed three pre-trained DDM backbones for validation, namely SEDD-small [6], SEDD-medium [6], and RADD [9]. These are absorbing DDMs of GPT-2 level for text generation, trained on the OpenWebText dataset [44]. For the uniform DDMs, please refer to the supplementary material. We compare LSD and LSD+ against standard Euler and Tweedie samplers [6] and the JYS method [17]. For the RADD baseline, we also compare with higher-order samplers, the θ -RK-2 and θ -trapezoidal [45].⁴ We evaluate the generative perplexity of unconditionally generated text using a GPT2-large model. We generated 1024 samples, each containing 1024 tokens. The results are presented in Tables 1, 2, and 3. LSD (LSD+)-Euler (Tweedie) denotes that we implement LSD (LSD+) based on the Euler (Tweedie) sampler [6]. The empirical results show that our methods significantly outperform the baseline methods across all three backbones and all tested NFEs. Moreover, we find that LSD+ generally outperforms LSD, which indicates that the learned non-uniform time schedules help to further reduce accumulated errors. Therefore, we only present the results for LSD+ for the experiments in Sections 4.2 and 4.3.

Table 1: Comparison of generative perplexity ($\leq$) on the SEDD-small backbone. Best performances are bolded.

Sampler \ NFEs	8	16	32	64
Euler	423.109	215.472	72.820	56.218
Tweedie	404.881	177.539	64.347	50.151
JYS-Euler	308.123	125.283	55.842	32.943
JYS-Tweedie	307.382	127.232	56.382	31.192
LSD-Euler	145.490	88.564	31.235	21.956
LSD-Tweedie	168.846	86.282	35.786	21.981
LSD+-Euler	128.413	51.769	36.800	20.728
LSD+-Tweedie	137.862	60.970	38.157	20.473

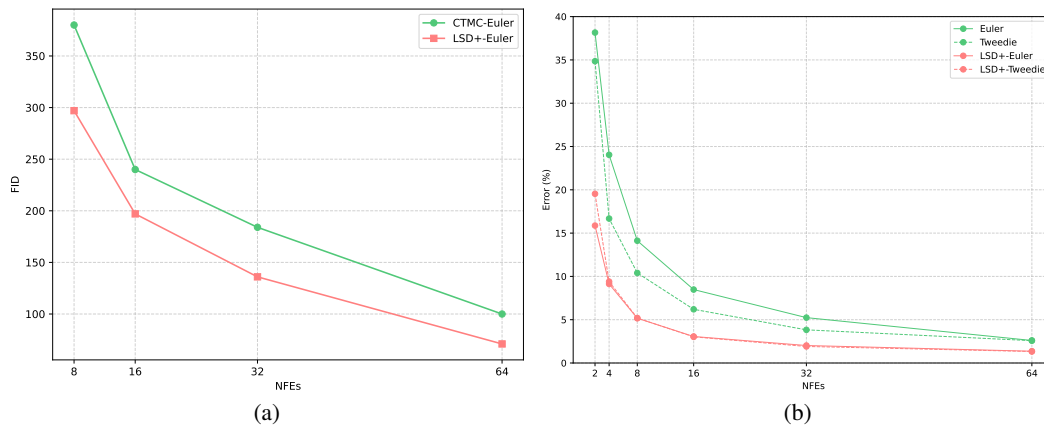


Figure 2: Validation on (a) the image generation task and (b) the synthetic countdown task. Our LSD+ method shows superior performance.

4.2 Image generation

We also validate our LSD+ approach on the image generation task for the CIFAR-10 dataset [46]. We utilize CTMC [8] as the baseline, which employs a Gaussian transition matrix and denoising parameterization. Each data sample is a flattened image with a size of $3 \times 32 \times 32$, composed of

⁴Since the source code of [45] is inaccessible, we could only compare our method with these high-order samplers on the RADD backbone. The results are reported from the paper. "/" in Table 3 denotes that this paper does not report the results when NFEs is 8.

Table 2: Comparison of generative perplexity ($\leq$) on the SEDD-medium backbone. Best performances are bolded.

Sampler \ NFEs	8	16	32	64
Euler	399.315	184.603	77.925	44.370
Tweedie	394.470	178.485	67.114	40.487
JYS-Euler	299.394	115.853	43.958	25.545
JYS-Tweedie	300.492	118.218	48.430	28.539
LSD-Euler	125.769	54.865	28.001	19.886
LSD-Tweedie	98.209	51.223	26.668	20.794
LSD+-Euler	121.583	46.145	25.144	15.929
LSD+-Tweedie	90.033	50.765	26.799	16.239

Table 3: Comparison of generative perplexity ($\leq$) on the RADD backbone. Best performances are bolded.

Sampler \ NFEs	8	16	32	64
Euler	670.977	282.115	152.403	113.913
Tweedie	648.736	285.471	155.472	98.879
θ -RK2	/	127.363	109.351	66.549
θ -Trapezoidal	/	123.585	89.912	66.549
LSD-Euler	121.420	60.219	46.268	32.817
LSD-Tweedie	122.167	67.176	40.274	28.644
LSD+-Euler	89.830	36.106	33.234	29.115
LSD+-Tweedie	90.364	40.263	36.129	24.312

tokens with values ranging from 0 to 255. We evaluate the FID score using 50k samples with the NFEs selected from $\{8, 16, 32, 64\}$. Figure 2(a) shows the results and we can observe that our method provides better FID scores compared to the baseline method.

4.3 Synthetic countdown task

We follow [47] to evaluate our LSD+ approach on a synthetic sequence task with strong dependencies. The dataset features 256-token sequences (with values in 0-31) where non-zero tokens must strictly decrease by one. We trained an absorbing SEDD [6] model and measured performance by the error rate, which is the proportion of generated samples violating this countdown rule. As shown in Figure 2(b), our method achieves lower error rates across various NFEs compared to baselines.

5 Ablation Study

The learnable coefficients form the core of our LSD approach and have demonstrated significant performance improvements. Also, we observe that LSD+ generally outperforms LSD as seen in Tables 1, 2, and 3, which indicates the benefit of the learned non-uniform time schedules. Therefore, our ablation study aims to assess the contributions of the relaxed objective during training.

Benefit of the relaxed objective We proposed a relaxed objective, allowing the student sampler to match the trajectory of the teacher sampler originating from x_{t_0} by using a perturbed starting point $\tilde{x}_{t_0}$ that is close to x_{t_0} during the training process. Table 4 compares the performance of LSD+ trained with and without this relaxation. The results clearly indicate that employing the relaxed objective generally yields better performance than training with the strict objective. This validates the benefit of the relaxation, confirming that it makes the trajectory alignment task more feasible and leads to better convergence.

Impact of Hamming distance threshold To investigate the robustness of the algorithm to the Hamming distance threshold, we conduct the ablation on the SEDD-small backbone using the Euler

sampler with 32 inference steps. We train our LSD+ method using several different values for the Hamming distance threshold, specifically 0%, 1%, 5%, 10%, 20% of the sequence length, while keeping all other hyperparameters unchanged. The performance, measured by Perplexity, is reported below in Table 5.

Table 4: Ablation study on the RADD backbone validating the importance of the Relaxed Objective (RO). "w/o RO" indicates parameters learned without relaxed objective, while "w/ RO" denotes parameters learned using our proposed relaxed objective. Both settings are evaluated using either Euler or Tweedie as the base sampler for our LSD+ method. Best performances are bolded.

Sampler \ NFEs				
	8	16	32	64
LSD+ w/o RO-Euler	95.943	38.192	34.983	31.392
LSD+ w/ RO-Euler	89.830	36.106	33.234	29.115

Table 5: Ablation study on the Hamming distance threshold for the relaxed objective.

Threshold(%)	0	1	5 (Our choice)	10	20
Perplexity(↓)	35.98	32.15	31.24	39.97	51.52

6 Conclusion

This paper aims to address the challenge of inefficient sampling in DDMs, a major obstacle to their practical deployment. While reducing the NFEs accelerates inference, previous accelerating methods suffer from accumulated compounding decoding error and discretization errors, significantly degrading sampling quality. We introduce LSD, a novel approach that leverages distillation from a high-fidelity teacher sampler. Instead of merely matching final outputs, LSD trains a student sampler with a few steps to align its entire intermediate score trajectory with that of the teacher sampler. This is achieved by optimizing learnable, time-dependent coefficients. And we additionally propose LSD+ that also learns non-uniform sampling schedules and this allows the sampler to adaptively compensate for errors induced by larger step sizes. Extensive experiments demonstrate that our methods significantly outperform the baseline samplers across diverse tasks, achieving high sampling fidelity at low NFEs.

A promising direction for future research is to provide theoretical guarantees regarding the distributional discrepancy between the outputs of teacher and student samplers, potentially building on existing theoretical findings related to discrete diffusion models [48, 49, 50, 51, 52, 45, 53, 54].

Acknowledgment. We sincerely thank the five anonymous reviewers and the area chair for their meticulous reading of our work and their constructive comments, which have substantially enhanced the quality and rigor of our study.

References

- [1] Ting Chen, Ruixiang Zhang, and Geoffrey Hinton. Analog bits: Generating discrete data using diffusion models with self-conditioning. In *International Conference on Learning Representations*, 2022.
- [2] Jonathan Ho, Ajay Jain, and Pieter Abbeel. Denoising diffusion probabilistic models. In *Advances in Neural Information Processing Systems*, 2020.
- [3] Zhifeng Kong, Wei Ping, Jiaji Huang, Kexin Zhao, and Bryan Catanzaro. DiffWave: A versatile diffusion model for audio synthesis. In *International Conference on Learning Representations*, 2021.
- [4] Cheng Lu, Yuhao Zhou, Fan Bao, Jianfei Chen, Chongxuan Li, and Jun Zhu. DPM-Solver: A Fast ODE Solver for Diffusion Probabilistic Model Sampling in Around 10 Steps. In *Advances in Neural Information Processing Systems*, 2022.
- [5] Alexander Quinn Nichol and Prafulla Dhariwal. Improved denoising diffusion probabilistic models. In *International Conference on Machine Learning*, 2021.
- [6] Aaron Lou, Chenlin Meng, and Stefano Ermon. Discrete diffusion modeling by estimating the ratios of the data distribution. In *International Conference on Machine Learning*, 2024.
- [7] Jacob Austin, Daniel D Johnson, Jonathan Ho, Daniel Tarlow, and Rianne Van Den Berg. Structured denoising diffusion models in discrete state-spaces. In *Advances in Neural Information Processing Systems*, 2021.
- [8] Andrew Campbell, Joe Benton, Valentin De Bortoli, Thomas Rainforth, George Deligiannidis, and Arnaud Doucet. A continuous time framework for discrete denoising models. In *Advances in Neural Information Processing Systems*, 2022.
- [9] Jingyang Ou, Shen Nie, Kaiwen Xue, Fengqi Zhu, Jiacheng Sun, Zhenguo Li, and Chongxuan Li. Your absorbing discrete diffusion secretly models the conditional distributions of clean data. In *International Conference on Learning Representations*, 2025.
- [10] Chenlin Meng, Kristy Choi, Jiaming Song, and Stefano Ermon. Concrete score matching: Generalized score matching for discrete data. In *Advances in Neural Information Processing Systems*, 2022.
- [11] Itai Gat, Tal Remez, Neta Shaul, Felix Kreuk, Ricky TQ Chen, Gabriel Synnaeve, Yossi Adi, and Yaron Lipman. Discrete flow matching. In *Advances in Neural Information Processing Systems*, 2024.
- [12] Zixiang Chen, Huizhuo Yuan, Yongqian Li, Yiwen Kou, Junkai Zhang, and Quanquan Gu. Fast sampling via discrete non-Markov diffusion models with predetermined transition time. In *Advances in Neural Information Processing Systems*, 2024.
- [13] Kaiwen Zheng, Yongxin Chen, Hanzi Mao, Ming-Yu Liu, Jun Zhu, and Qinsheng Zhang. Masked diffusion models are secretly time-agnostic masked models and exploit inaccurate categorical sampling. In *International Conference on Learning Representations*, 2025.
- [14] Jiaxin Shi, Kehang Han, Zhe Wang, Arnaud Doucet, and Michalis Titsias. Simplified and generalized masked diffusion for discrete data. In *Advances in Neural Information Processing Systems*, 2024.
- [15] Daniel T Gillespie. Approximate accelerated stochastic simulation of chemically reacting systems. *The Journal of Chemical Physics*, 2001.
- [16] Bradley Efron. Tweedie’s formula and selection bias. *Journal of the American Statistical Association*, 2011.
- [17] Yong-Hyun Park, Chieh-Hsin Lai, Satoshi Hayakawa, Yuhta Takida, and Yuki Mitsufuji. Jump your steps: Optimizing sampling schedule of discrete diffusion models. In *International Conference on Learning Representations*, 2025.

- [18] David F Anderson, Arnab Ganguly, and Thomas G Kurtz. Error analysis of tau-leap simulation methods. *Annals of Applied Probability*, 2011.
- [19] Yang Song, Jascha Sohl-Dickstein, Diederik P Kingma, Abhishek Kumar, Stefano Ermon, and Ben Poole. Score-based generative modeling through stochastic differential equations. In *International Conference on Learning Representations*, 2021.
- [20] Kaiwen Zheng, Cheng Lu, Jianfei Chen, and Jun Zhu. DPM-Solver-v3: Improved diffusion ODE solver with empirical model statistics. In *Advances in Neural Information Processing Systems*, 2023.
- [21] Shuchen Xue, Mingyang Yi, Weijian Luo, Shifeng Zhang, Jiacheng Sun, Zhenguo Li, and Zhi-Ming Ma. SA-Solver: Stochastic Adams solver for fast sampling of diffusion models. In *Advances in Neural Information Processing Systems*, 2023.
- [22] Gen Li, Yu Huang, Timofey Efimov, Yuting Wei, Yuejie Chi, and Yuxin Chen. Accelerating convergence of score-based diffusion models, provably. In *International Conference on Machine Learning*, 2024.
- [23] Jiajun Ma, Shuchen Xue, Tianyang Hu, Wenjia Wang, Zhaoqiang Liu, Zhenguo Li, Zhi-Ming Ma, and Kenji Kawaguchi. The surprising effectiveness of skip-tuning in diffusion sampling. In *International Conference on Machine Learning*, 2024.
- [24] Mingyuan Zhou, Huangjie Zheng, Zhendong Wang, Mingzhang Yin, and Hai Huang. Score identity distillation: Exponentially fast distillation of pretrained diffusion models for one-step generation. In *International Conference on Machine Learning*, 2024.
- [25] Cheng Lu, Yuhao Zhou, Fan Bao, Jianfei Chen, Chongxuan Li, and Jun Zhu. DPM-Solver++: Fast solver for guided sampling of diffusion probabilistic models. *arXiv preprint arXiv:2211.01095*, 2022.
- [26] Qinsheng Zhang and Yongxin Chen. Fast sampling of diffusion models with exponential integrator. In *International Conference on Learning Representations*, 2023.
- [27] Tim Dockhorn, Arash Vahdat, and Karsten Kreis. GENIE: Higher-order denoising diffusion solvers. In *Advances in Neural Information Processing Systems*, 2022.
- [28] Amirmojtaba Sabour, Sanja Fidler, and Karsten Kreis. Align your steps: Optimizing sampling schedules in diffusion models. In *International Conference on Machine Learning*, 2024.
- [29] Shuchen Xue, Zhaoqiang Liu, Fei Chen, Shifeng Zhang, Tianyang Hu, Enze Xie, and Zhenguo Li. Accelerating diffusion sampling with optimized time steps. In *IEEE/CVF Conference on Computer Vision and Pattern Recognition*, 2024.
- [30] Zhenyu Zhou, Defang Chen, Can Wang, and Chun Chen. Fast ODE-based sampling for diffusion models in around 5 steps. In *IEEE/CVF Conference on Computer Vision and Pattern Recognition*, 2024.
- [31] Vinh Tong, Trung-Dung Hoang, Anji Liu, Guy Van den Broeck, and Mathias Niepert. Learning to discretize denoising diffusion ODEs. In *International Conference on Learning Representations*, 2025.
- [32] Eric Frankel, Sitan Chen, Jerry Li, Pang Wei Koh, Lillian J Ratliff, and Sewoong Oh. S4S: Solving for a diffusion model solver. In *International Conference on Machine Learning*, 2025.
- [33] Yang Song, Prafulla Dhariwal, Mark Chen, and Ilya Sutskever. Consistency models. In *International Conference on Machine Learning*, 2023.
- [34] Nicholas M Boffi, Michael S Albergo, and Eric Vanden-Eijnden. How to build a consistency model: Learning flow maps via self-distillation. *arXiv preprint arXiv:2505.18825*, 2025.
- [35] Jonathan Heek, Emiel Hoogeboom, and Tim Salimans. Multistep consistency models. *arXiv preprint arXiv:2403.06807*, 2024.

- [36] Chenlin Meng, Robin Rombach, Ruiqi Gao, Diederik Kingma, Stefano Ermon, Jonathan Ho, and Tim Salimans. On distillation of guided diffusion models. In *IEEE/CVF Conference on Computer Vision and Pattern Recognition*, 2023.
- [37] Zhenyu Zhou, Defang Chen, Can Wang, Chun Chen, and Siwei Lyu. Simple and fast distillation of diffusion models. In *Advances in Neural Information Processing Systems*, 2024.
- [38] Tim Salimans and Jonathan Ho. Progressive distillation for fast sampling of diffusion models. In *International Conference on Learning Representations*, 2022.
- [39] Sirui Xie, Zhisheng Xiao, Diederik Kingma, Tingbo Hou, Ying Nian Wu, Kevin P Murphy, Tim Salimans, Ben Poole, and Ruiqi Gao. EM distillation for one-step diffusion models. In *Advances in Neural Information Processing Systems*, 2024.
- [40] Yuanzhi Zhu, Xi Wang, Stéphane Lathuilière, and Vicky Kalogeiton. Di[M]O: Distilling masked diffusion models into one-step generator. In *IEEE International Conference on Computer Vision*, 2025.
- [41] Haoran Sun, Lijun Yu, Bo Dai, Dale Schuurmans, and Hanjun Dai. Score-based continuous-time discrete diffusion models. In *International Conference on Learning Representations*, 2023.
- [42] Andrew Campbell, Jason Yim, Regina Barzilay, Tom Rainforth, and Tommi Jaakkola. Generative flows on discrete state-spaces: Enabling multimodal flows with applications to protein co-design. In *International Conference on Machine Learning*, 2024.
- [43] Mohammad Norouzi, David J Fleet, and Russ R Salakhutdinov. Hamming distance metric learning. In *Advances in Neural Information Processing Systems*, 2012.
- [44] Aaron Gokaslan and Vanya Cohen. OpenWebText Corpus. <http://Skylion007.github.io/OpenWebTextCorpus>, 2019.
- [45] Yinuo Ren, Haoxuan Chen, Yuchen Zhu, Wei Guo, Yongxin Chen, Grant M Rotskoff, Molei Tao, and Lexing Ying. Fast solvers for discrete diffusion models: Theory and applications of high-order algorithms. *arXiv preprint arXiv:2502.00234*, 2025.
- [46] Alex Krizhevsky. Learning multiple layers of features from tiny images. Technical report, 2009.
- [47] Yixiu Zhao, Jiaxin Shi, Feng Chen, Shaul Druckmann, Lester Mackey, and Scott Linderman. Informed correctors for discrete diffusion models. *arXiv preprint arXiv:2407.21243*, 2024.
- [48] Yinuo Ren, Haoxuan Chen, Grant M Rotskoff, and Lexing Ying. How discrete and continuous diffusion meet: Comprehensive analysis of discrete diffusion models via a stochastic integral framework. In *International Conference on Learning Representations*, 2025.
- [49] Hongrui Chen and Lexing Ying. Convergence analysis of discrete diffusion model: Exact implementation through uniformization. In *International Conference on Learning Representations*, 2025.
- [50] Zikun Zhang, Zixiang Chen, and Quanquan Gu. Convergence of score-based discrete diffusion models: A discrete-time analysis. In *International Conference on Learning Representations*, 2025.
- [51] Guhao Feng, Yihan Geng, Jian Guan, Wei Wu, Liwei Wang, and Di He. Theoretical benefit and limitation of diffusion language model. *arXiv preprint arXiv:2502.09622*, 2025.
- [52] Gen Li and Changxiao Cai. A convergence theory for diffusion language models: An information-theoretic perspective. *arXiv preprint arXiv:2505.21400*, 2025.
- [53] Leo Zhang. The cosine schedule is Fisher-Rao-optimal for masked discrete diffusion models. *arXiv preprint arXiv:2508.04884*, 2025.
- [54] Ruixiang Zhang, Shuangfei Zhai, Yizhe Zhang, James Thornton, Zijiang Ou, Joshua M Susskind, and Navdeep Jaitly. Target concrete score matching: A holistic framework for discrete diffusion. In *International Conference on Machine Learning*, 2025.

- [55] Yuchen Liang, Renxiang Huang, Lifeng Lai, Ness Shroff, and Yingbin Liang. Absorb and converge: Provable convergence guarantee for absorbing discrete diffusion models. *arXiv preprint arXiv:2506.02318*, 2025.
- [56] Zikun Zhang, Zixiang Chen, and Quanquan Gu. Convergence of score-based discrete diffusion models: A discrete-time analysis. In *International Conference on Learning Representations*, 2025.
- [57] Huiwen Chang, Han Zhang, Lu Jiang, Ce Liu, and William T Freeman. MaskGIT: Masked generative image transformer. In *IEEE/CVF Conference on Computer Vision and Pattern Recognition*, 2022.
- [58] Victor Besnier, Mickael Chen, David Hurych, Eduardo Valle, and Matthieu Cord. Halton scheduler for masked generative image transformer. *arXiv preprint arXiv:2503.17076*, 2025.
- [59] Justin Deschenaux and Caglar Gulcehre. Beyond autoregression: Fast LLMs via self-distillation through time. In *International Conference on Learning Representations*, 2025.
- [60] Guanghan Wang, Yair Schiff, Subham Sekhar Sahoo, and Volodymyr Kuleshov. Remasking discrete diffusion models with inference-time scaling. *arXiv preprint arXiv:2503.00307*, 2025.
- [61] Subham Sekhar Sahoo, Marianne Arriola, Aaron Gokaslan, Edgar Mariano Marroquin, Alexander M Rush, Yair Schiff, Justin T Chiu, and Volodymyr Kuleshov. Simple and effective masked diffusion language models. In *Conference on Neural Information Processing Systems*, 2024.
- [62] Abhimanyu Dubey, Abhinav Jauhri, Abhinav Pandey, Abhishek Kadian, Ahmad Al-Dahle, Aiesha Letman, Akhil Mathur, Alan Schelten, Amy Yang, Angela Fan, et al. The Llama 3 herd of models. *arXiv preprint arXiv:2407.21783*, 2024.
- [63] Chengyue Wu, Hao Zhang, Shuchen Xue, Zhijian Liu, Shizhe Diao, Ligeng Zhu, Ping Luo, Song Han, and Enze Xie. Fast-dLLM: Training-free acceleration of diffusion LLM by enabling KV cache and parallel decoding. *arXiv preprint arXiv:2505.22618*, 2025.
- [64] Shansan Gong, Shivam Agarwal, Yizhe Zhang, Jiacheng Ye, Lin Zheng, Mukai Li, Chenxin An, Peilin Zhao, Wei Bi, Jiawei Han, et al. Scaling diffusion language models via adaptation from autoregressive models. In *International Conference on Learning Representations*, 2025.
- [65] Myle Ott, Sergey Edunov, Alexei Baevski, Angela Fan, Sam Gross, Nathan Ng, David Grangier, and Michael Auli. FAIRSEQ: A fast, extensible toolkit for sequence modeling. *arXiv preprint arXiv:1904.01038*, 2019.
- [66] Andrzej Maćkiewicz and Waldemar Ratajczak. Principal components analysis. *Computers & Geosciences*, 1993.
- [67] Zhaoqiang Liu and Vincent YF Tan. The informativeness of k -means for learning mixture models. *IEEE Transactions on Information Theory*, 65(11):7460–7479, 2019.

NeurIPS Paper Checklist

1. Claims

Question: Do the main claims made in the abstract and introduction accurately reflect the paper's contributions and scope?

Answer: [Yes]

Justification: We provide brief and clear main claims in the abstract and introduction.

Guidelines:

- The answer NA means that the abstract and introduction do not include the claims made in the paper.
- The abstract and/or introduction should clearly state the claims made, including the contributions made in the paper and important assumptions and limitations. A No or NA answer to this question will not be perceived well by the reviewers.
- The claims made should match theoretical and experimental results, and reflect how much the results can be expected to generalize to other settings.
- It is fine to include aspirational goals as motivation as long as it is clear that these goals are not attained by the paper.

2. Limitations

Question: Does the paper discuss the limitations of the work performed by the authors?

Answer: [Yes]

Justification: We discuss the limitations in the supplementary material.

Guidelines:

- The answer NA means that the paper has no limitation while the answer No means that the paper has limitations, but those are not discussed in the paper.
- The authors are encouraged to create a separate "Limitations" section in their paper.
- The paper should point out any strong assumptions and how robust the results are to violations of these assumptions (e.g., independence assumptions, noiseless settings, model well-specification, asymptotic approximations only holding locally). The authors should reflect on how these assumptions might be violated in practice and what the implications would be.
- The authors should reflect on the scope of the claims made, e.g., if the approach was only tested on a few datasets or with a few runs. In general, empirical results often depend on implicit assumptions, which should be articulated.
- The authors should reflect on the factors that influence the performance of the approach. For example, a facial recognition algorithm may perform poorly when image resolution is low or images are taken in low lighting. Or a speech-to-text system might not be used reliably to provide closed captions for online lectures because it fails to handle technical jargon.
- The authors should discuss the computational efficiency of the proposed algorithms and how they scale with dataset size.
- If applicable, the authors should discuss possible limitations of their approach to address problems of privacy and fairness.
- While the authors might fear that complete honesty about limitations might be used by reviewers as grounds for rejection, a worse outcome might be that reviewers discover limitations that aren't acknowledged in the paper. The authors should use their best judgment and recognize that individual actions in favor of transparency play an important role in developing norms that preserve the integrity of the community. Reviewers will be specifically instructed to not penalize honesty concerning limitations.

3. Theory assumptions and proofs

Question: For each theoretical result, does the paper provide the full set of assumptions and a complete (and correct) proof?

Answer: [NA]

Justification: Our paper does not involve theoretical proofs.

Guidelines:

- The answer NA means that the paper does not include theoretical results.
- All the theorems, formulas, and proofs in the paper should be numbered and cross-referenced.
- All assumptions should be clearly stated or referenced in the statement of any theorems.
- The proofs can either appear in the main paper or the supplemental material, but if they appear in the supplemental material, the authors are encouraged to provide a short proof sketch to provide intuition.
- Inversely, any informal proof provided in the core of the paper should be complemented by formal proofs provided in appendix or supplemental material.
- Theorems and Lemmas that the proof relies upon should be properly referenced.

4. Experimental result reproducibility

Question: Does the paper fully disclose all the information needed to reproduce the main experimental results of the paper to the extent that it affects the main claims and/or conclusions of the paper (regardless of whether the code and data are provided or not)?

Answer: [\[Yes\]](#)

Justification: We provide the implementation details in Section 4 and the supplementary material.

Guidelines:

- The answer NA means that the paper does not include experiments.
- If the paper includes experiments, a No answer to this question will not be perceived well by the reviewers: Making the paper reproducible is important, regardless of whether the code and data are provided or not.
- If the contribution is a dataset and/or model, the authors should describe the steps taken to make their results reproducible or verifiable.
- Depending on the contribution, reproducibility can be accomplished in various ways. For example, if the contribution is a novel architecture, describing the architecture fully might suffice, or if the contribution is a specific model and empirical evaluation, it may be necessary to either make it possible for others to replicate the model with the same dataset, or provide access to the model. In general, releasing code and data is often one good way to accomplish this, but reproducibility can also be provided via detailed instructions for how to replicate the results, access to a hosted model (e.g., in the case of a large language model), releasing of a model checkpoint, or other means that are appropriate to the research performed.
- While NeurIPS does not require releasing code, the conference does require all submissions to provide some reasonable avenue for reproducibility, which may depend on the nature of the contribution. For example
 - (a) If the contribution is primarily a new algorithm, the paper should make it clear how to reproduce that algorithm.
 - (b) If the contribution is primarily a new model architecture, the paper should describe the architecture clearly and fully.
 - (c) If the contribution is a new model (e.g., a large language model), then there should either be a way to access this model for reproducing the results or a way to reproduce the model (e.g., with an open-source dataset or instructions for how to construct the dataset).
 - (d) We recognize that reproducibility may be tricky in some cases, in which case authors are welcome to describe the particular way they provide for reproducibility. In the case of closed-source models, it may be that access to the model is limited in some way (e.g., to registered users), but it should be possible for other researchers to have some path to reproducing or verifying the results.

5. Open access to data and code

Question: Does the paper provide open access to the data and code, with sufficient instructions to faithfully reproduce the main experimental results, as described in supplemental material?

Answer: [No]

Justification: The data used in the experiments are open-sourced, and our code will be released upon acceptance.

Guidelines:

- The answer NA means that paper does not include experiments requiring code.
- Please see the NeurIPS code and data submission guidelines (<https://nips.cc/public/guides/CodeSubmissionPolicy>) for more details.
- While we encourage the release of code and data, we understand that this might not be possible, so “No” is an acceptable answer. Papers cannot be rejected simply for not including code, unless this is central to the contribution (e.g., for a new open-source benchmark).
- The instructions should contain the exact command and environment needed to run to reproduce the results. See the NeurIPS code and data submission guidelines (<https://nips.cc/public/guides/CodeSubmissionPolicy>) for more details.
- The authors should provide instructions on data access and preparation, including how to access the raw data, preprocessed data, intermediate data, and generated data, etc.
- The authors should provide scripts to reproduce all experimental results for the new proposed method and baselines. If only a subset of experiments are reproducible, they should state which ones are omitted from the script and why.
- At submission time, to preserve anonymity, the authors should release anonymized versions (if applicable).
- Providing as much information as possible in supplemental material (appended to the paper) is recommended, but including URLs to data and code is permitted.

6. Experimental setting/details

Question: Does the paper specify all the training and test details (e.g., data splits, hyper-parameters, how they were chosen, type of optimizer, etc.) necessary to understand the results?

Answer: [Yes]

Justification: Section 4 and the supplementary material provide the experimental settings.

Guidelines:

- The answer NA means that the paper does not include experiments.
- The experimental setting should be presented in the core of the paper to a level of detail that is necessary to appreciate the results and make sense of them.
- The full details can be provided either with the code, in appendix, or as supplemental material.

7. Experiment statistical significance

Question: Does the paper report error bars suitably and correctly defined or other appropriate information about the statistical significance of the experiments?

Answer: [No]

Justification: We follow closely relevant works such as SEDD [6] and RADD [9] to calculate the quantitative results without reporting error bars.

Guidelines:

- The answer NA means that the paper does not include experiments.
- The authors should answer "Yes" if the results are accompanied by error bars, confidence intervals, or statistical significance tests, at least for the experiments that support the main claims of the paper.
- The factors of variability that the error bars are capturing should be clearly stated (for example, train/test split, initialization, random drawing of some parameter, or overall run with given experimental conditions).
- The method for calculating the error bars should be explained (closed form formula, call to a library function, bootstrap, etc.)
- The assumptions made should be given (e.g., Normally distributed errors).

- It should be clear whether the error bar is the standard deviation or the standard error of the mean.
- It is OK to report 1-sigma error bars, but one should state it. The authors should preferably report a 2-sigma error bar than state that they have a 96% CI, if the hypothesis of Normality of errors is not verified.
- For asymmetric distributions, the authors should be careful not to show in tables or figures symmetric error bars that would yield results that are out of range (e.g. negative error rates).
- If error bars are reported in tables or plots, The authors should explain in the text how they were calculated and reference the corresponding figures or tables in the text.

8. Experiments compute resources

Question: For each experiment, does the paper provide sufficient information on the computer resources (type of compute workers, memory, time of execution) needed to reproduce the experiments?

Answer: [Yes]

Justification: We provide sufficient information on the computer resources in the supplementary material.

Guidelines:

- The answer NA means that the paper does not include experiments.
- The paper should indicate the type of compute workers CPU or GPU, internal cluster, or cloud provider, including relevant memory and storage.
- The paper should provide the amount of compute required for each of the individual experimental runs as well as estimate the total compute.
- The paper should disclose whether the full research project required more compute than the experiments reported in the paper (e.g., preliminary or failed experiments that didn't make it into the paper).

9. Code of ethics

Question: Does the research conducted in the paper conform, in every respect, with the NeurIPS Code of Ethics <https://neurips.cc/public/EthicsGuidelines>?

Answer: [Yes]

Justification: We have read and conformed with the NeurIPS Code of Ethics in every respect.

Guidelines:

- The answer NA means that the authors have not reviewed the NeurIPS Code of Ethics.
- If the authors answer No, they should explain the special circumstances that require a deviation from the Code of Ethics.
- The authors should make sure to preserve anonymity (e.g., if there is a special consideration due to laws or regulations in their jurisdiction).

10. Broader impacts

Question: Does the paper discuss both potential positive societal impacts and negative societal impacts of the work performed?

Answer: [Yes]

Justification: We provide discussions on potential societal impacts in the supplementary material.

Guidelines:

- The answer NA means that there is no societal impact of the work performed.
- If the authors answer NA or No, they should explain why their work has no societal impact or why the paper does not address societal impact.
- Examples of negative societal impacts include potential malicious or unintended uses (e.g., disinformation, generating fake profiles, surveillance), fairness considerations (e.g., deployment of technologies that could make decisions that unfairly impact specific groups), privacy considerations, and security considerations.

- The conference expects that many papers will be foundational research and not tied to particular applications, let alone deployments. However, if there is a direct path to any negative applications, the authors should point it out. For example, it is legitimate to point out that an improvement in the quality of generative models could be used to generate deepfakes for disinformation. On the other hand, it is not needed to point out that a generic algorithm for optimizing neural networks could enable people to train models that generate Deepfakes faster.
- The authors should consider possible harms that could arise when the technology is being used as intended and functioning correctly, harms that could arise when the technology is being used as intended but gives incorrect results, and harms following from (intentional or unintentional) misuse of the technology.
- If there are negative societal impacts, the authors could also discuss possible mitigation strategies (e.g., gated release of models, providing defenses in addition to attacks, mechanisms for monitoring misuse, mechanisms to monitor how a system learns from feedback over time, improving the efficiency and accessibility of ML).

11. Safeguards

Question: Does the paper describe safeguards that have been put in place for responsible release of data or models that have a high risk for misuse (e.g., pretrained language models, image generators, or scraped datasets)?

Answer: [NA]

Justification: This paper poses no such risks.

Guidelines:

- The answer NA means that the paper poses no such risks.
- Released models that have a high risk for misuse or dual-use should be released with necessary safeguards to allow for controlled use of the model, for example by requiring that users adhere to usage guidelines or restrictions to access the model or implementing safety filters.
- Datasets that have been scraped from the Internet could pose safety risks. The authors should describe how they avoided releasing unsafe images.
- We recognize that providing effective safeguards is challenging, and many papers do not require this, but we encourage authors to take this into account and make a best faith effort.

12. Licenses for existing assets

Question: Are the creators or original owners of assets (e.g., code, data, models), used in the paper, properly credited and are the license and terms of use explicitly mentioned and properly respected?

Answer: [Yes]

Justification: We have cited the original paper that produced the code package or dataset.

Guidelines:

- The answer NA means that the paper does not use existing assets.
- The authors should cite the original paper that produced the code package or dataset.
- The authors should state which version of the asset is used and, if possible, include a URL.
- The name of the license (e.g., CC-BY 4.0) should be included for each asset.
- For scraped data from a particular source (e.g., website), the copyright and terms of service of that source should be provided.
- If assets are released, the license, copyright information, and terms of use in the package should be provided. For popular datasets, paperswithcode.com/datasets has curated licenses for some datasets. Their licensing guide can help determine the license of a dataset.
- For existing datasets that are re-packaged, both the original license and the license of the derived asset (if it has changed) should be provided.

- If this information is not available online, the authors are encouraged to reach out to the asset's creators.

13. **New assets**

Question: Are new assets introduced in the paper well documented and is the documentation provided alongside the assets?

Answer: [NA]

Justification: This paper does not release new assets.

Guidelines:

- The answer NA means that the paper does not release new assets.
- Researchers should communicate the details of the dataset/code/model as part of their submissions via structured templates. This includes details about training, license, limitations, etc.
- The paper should discuss whether and how consent was obtained from people whose asset is used.
- At submission time, remember to anonymize your assets (if applicable). You can either create an anonymized URL or include an anonymized zip file.

14. **Crowdsourcing and research with human subjects**

Question: For crowdsourcing experiments and research with human subjects, does the paper include the full text of instructions given to participants and screenshots, if applicable, as well as details about compensation (if any)?

Answer: [NA]

Justification: This paper does not involve crowdsourcing nor research with human subjects.

Guidelines:

- The answer NA means that the paper does not involve crowdsourcing nor research with human subjects.
- Including this information in the supplemental material is fine, but if the main contribution of the paper involves human subjects, then as much detail as possible should be included in the main paper.
- According to the NeurIPS Code of Ethics, workers involved in data collection, curation, or other labor should be paid at least the minimum wage in the country of the data collector.

15. **Institutional review board (IRB) approvals or equivalent for research with human subjects**

Question: Does the paper describe potential risks incurred by study participants, whether such risks were disclosed to the subjects, and whether Institutional Review Board (IRB) approvals (or an equivalent approval/review based on the requirements of your country or institution) were obtained?

Answer: [NA]

Justification: This paper does not involve crowdsourcing nor research with human subjects.

Guidelines:

- The answer NA means that the paper does not involve crowdsourcing nor research with human subjects.
- Depending on the country in which research is conducted, IRB approval (or equivalent) may be required for any human subjects research. If you obtained IRB approval, you should clearly state this in the paper.
- We recognize that the procedures for this may vary significantly between institutions and locations, and we expect authors to adhere to the NeurIPS Code of Ethics and the guidelines for their institution.
- For initial submissions, do not include any information that would break anonymity (if applicable), such as the institution conducting the review.

16. **Declaration of LLM usage**

Question: Does the paper describe the usage of LLMs if it is an important, original, or non-standard component of the core methods in this research? Note that if the LLM is used only for writing, editing, or formatting purposes and does not impact the core methodology, scientific rigorousness, or originality of the research, declaration is not required.

Answer: [NA]

Justification: The core method development in this research does not involve LLMs as any important, original, or non-standard components.

Guidelines:

- The answer NA means that the core method development in this research does not involve LLMs as any important, original, or non-standard components.
- Please refer to our LLM policy (<https://neurips.cc/Conferences/2025/LLM>) for what should or should not be described.

Supplementary Material

Learnable Sampler Distillation for Discrete Diffusion Models (NeurIPS 2025)

Feiyang Fu, Tongxian Guo, Zhaoqiang Liu

A Limitations and Broader Impact

While our proposed learnable sampler distillation (LSD) approach significantly enhances the sampling quality of discrete diffusion models (DDMs) at low NFEs, there exists a limitation that warrants discussion. Specifically, the performance ceiling of the student sampler is inherently tied to the quality of the teacher sampler. Consequently, while our student sampler can efficiently achieve the performance of the teacher sampler with very few NFEs, surpassing it is challenging. However, this dependence on the teacher is an intrinsic characteristic of knowledge distillation paradigms and LSD is compatible with future advancements, since more sophisticated samplers can serve as improved teacher samplers.

Our approaches can be integrated with various DDM samplers, offering a general path to enhance their efficiency, and hold significant promise for accelerating scientific discovery, such as by facilitating the design of candidate DNA and protein sequences. However, the increased power of generative AI also necessitates a commitment to responsible development. This includes proactive efforts to mitigate societal risks, notably the potential for generating misinformation or amplifying existing biases, and underscores the importance of ethical guidelines and detection research.

B Discussion

B.1 Intuitive explanation

Given that several theoretical works on DDMs have been conducted [49, 55, 56, 48], we provide an intuitive explanation for the effectiveness of our method, which may help lay the groundwork for subsequent theoretical framing. Specifically, the final discrepancy between the outputs of the student and teacher samplers stems from the accumulation of small local errors made at each step. Each of these local errors, in turn, is tied to the specific score predicted by the model at that step. Our training objective directly enforces alignment between the score predictions of the student sampler and those of the teacher sampler at every step. By implicitly correcting these small local errors throughout the process, our approach guides the student sampler to produce final outputs that closely match the high-quality results of the teacher sampler.

B.2 Relaxed objective

Our use of a relaxed training objective enhances training feasibility compared to strict alignment. However, the theoretical guarantees that the resulting student distribution closely matches the teacher distribution might be less rigorous than what could potentially be argued for in continuous spaces. To the best of our knowledge, establishing such rigorous theoretical bounds in discrete spaces faces several challenges not present in their continuous counterparts. Specifically, (1) In continuous diffusion, initial state perturbations can often be controlled and analyzed via differentiable operations (e.g., L_2 -constrained gradient steps) [32]. In contrast, discrete initial states (e.g., token sequences) and their perturbations that are measured by Hamming distance lack this differentiability. This precludes similar continuous optimization and analysis pathways. (2) The analytical tools used for relaxed objective in continuous diffusion ODE/SDE, such as perturbation analysis for smooth dynamical systems [31], do not directly translate to the discrete dynamics of CTMCs and their approximations. Therefore, our proposed relaxed objective serves as an empirically effective solution for training feasibility in LSD.

C Details for discrete diffusion models

In the main text, we illustrate the mechanism by which the forward process of CTMC introduces corruption into the data. Within Eq. (1), the term $Q_t(x, y)$ is delineated as the transition rate from state x to state y at time t . Consistent with this definition, $Q_t(x, y)$ can be articulated as follows:

$$Q_t(x, y) = \begin{cases} \lim_{\Delta t \rightarrow 0} \frac{p_{t+\Delta t|t}(y|x) - p_t(y|x)}{\Delta t}, & y \neq x, \\ \lim_{\Delta t \rightarrow 0} \frac{p_{t+\Delta t|t}(x|x) - 1}{\Delta t}, & y = x. \end{cases} \quad (11)$$

For instances where $t > s$, we define $P_{t|s}(x, y) := p_{t|s}(y|x)$. Drawing upon Kolmogorov's forward equation [8] and the definition of Q_t , we derive

$$\frac{d}{dt} P_{t|s} = P_{t|s} Q_t. \quad (12)$$

The analytical solution to Eq. (12) is given by $P_{t|s} = \exp((\bar{\sigma}(t) - \bar{\sigma}(s))Q)$, with $\bar{\sigma}(t)$ denoting the integral $\int_0^t \sigma(s)ds$ and $\exp$ denoting the matrix exponential function. This solution facilitates the direct sampling of x_t from x_s in a single step for all $t > s$ scenarios.

The transition rate matrix Q_t is formulated as $\sigma(t)Q$, where Q represents a pre-specified standard matrix. In the context of defining matrix Q , two principal alternatives are presented: A uniform distribution or a MASK absorbing state. When the base transition matrix Q is selected to be a uniform matrix, it simulates a fully connected graph structure. Within this framework, each state is interconnected with all other states, implying that transitions from any given state to any other state are possible. The definition of this transition matrix ensures extensive exploratory capabilities of the state space, permitting the model to account for all potential state transitions during simulation. Specifically, the construction of Q^{uniform} is as follows:

$$Q^{\text{uniform}} = \begin{bmatrix} 1-N & 1 & \cdots & 1 & 1 \\ 1 & 1-N & \cdots & 1 & 1 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 1 & 1 & \cdots & 1-N & 1 \\ 1 & 1 & \cdots & 1 & 1-N \end{bmatrix}. \quad (13)$$

An alternative approach in the construction of the transition matrix Q includes the formulation of a matrix that encompasses absorbing states. An absorbing state refers to a state, where the system will no longer undergo state transitions once it enters this state, and such a design facilitates the rapid convergence of the model to a stable state. Recent works [6, 9] have also demonstrated that the adoption of an absorbing matrix is associated with better performance and serves to accelerate the sampling process. The construction of Q^{absorb} is defined as:

$$Q^{\text{absorb}} = \begin{bmatrix} -1 & 0 & \cdots & 0 & 1 \\ 0 & -1 & \cdots & 0 & 1 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 0 & 0 & \cdots & -1 & 1 \\ 0 & 0 & \cdots & 0 & 0 \end{bmatrix}. \quad (14)$$

To effectively simulate the reverse process, the common practice is to train a neural network $s_\theta(x, t)$ to approximate the required concrete score $\frac{p_t(y)}{p_t(x)}$. In this process, to optimize the training of network $s_\theta(x, t)$, SEDD [6] introduces an effective loss function:

$$\int_0^T \mathbb{E}_{x \sim p_{t|0}(x|x_0)} \sum_{y \neq x} Q_t(y, x) \left(s_\theta(x, t)_y - \frac{p_{t|0}(y|x_0)}{p_{t|0}(x|x_0)} \log s_\theta(x, t)_y + C \right) dt, \quad (15)$$

where C is a constant with $C = K \left(\frac{p_{t|0}(y|x_0)}{p_{t|0}(x|x_0)} \right)$ and $K(a) := a \log a - a$, $s_\theta(x, t)_y$ represents an estimate by the neural network of the probability of transitioning from state x to state y at time t .

D Additional method details

D.1 LSD using Tweedie τ -leaping

In the main text, we detailed the application of our LSD approach to an Euler-type discrete sampler. Notably, LSD can also be readily integrated with other sampling methods, such as Tweedie τ -leaping.

The Tweedie τ -leaping update rule for DDMs leverages Tweedie's formula to relate the conditional expectation of a cleaner state to the score function. Given time schedules $\{t_k\}_{k=0}^M$ for the student sampler. The transition probability $p(x_{t_{k+1}}^i | x_{t_k}^i)$ for the i -th token from state x_{t_k} at time t_k to state $x_{t_{k+1}}$ at the subsequent time step t_{k+1} is constructed as follows:

$$p(x_{t_{k+1}}^i | x_{t_k}^i) = (\exp((\bar{\sigma}(t_{k+1}) - \bar{\sigma}(t_k))Q) s_{\theta}(x_{t_k}, t_k)_i)_{x_{t_{k+1}}^i} \times (\exp((\bar{\sigma}(t_k) - \bar{\sigma}(t_{k+1}))Q))_{x_{t_k}^i, x_{t_{k+1}}^i}, \quad (16)$$

where Q is the predefined standard matrix with special structure as mentioned in Section C, $\bar{\sigma}(t)$ be the cumulative noise schedule, which is a non-decreasing function of t , $s_{\theta}(x_{t_k}, t_k)_i$ is the i -th element of the score $s_{\theta}(x_{t_k}, t_k)$. The subscript $x_{t_{k+1}}^i$ on the first main term denotes the $x_{t_{k+1}}^i$ -th element of the resulting vector. Similarly, the subscript $(x_{t_k}^i, x_{t_{k+1}}^i)$ on the second main term denotes selecting the corresponding element from the second main term.

To incorporate our LSD approach, we introduce the learnable time-dependent coefficient $\{\Phi(t_k)\}_{k=1}^M$ to modulate the score network s_{θ} within the Tweedie update. The LSD-modified Tweedie τ -leaping update rule is as follows:

$$p(x_{t_{k+1}}^i | x_{t_k}^i) = (\exp((\bar{\sigma}(t_{k+1}) - \bar{\sigma}(t_k))Q) (\Phi(t_k) s_{\theta}(x_{t_k}, t_k))_i)_{x_{t_{k+1}}^i} \times (\exp((\bar{\sigma}(t_k) - \bar{\sigma}(t_{k+1}))Q))_{x_{t_k}^i, x_{t_{k+1}}^i}. \quad (17)$$

This formulation allows the student sampler using Tweedie τ -leaping to learn an adaptive scaling $\{\Phi(t_k)\}_{k=1}^M$ of the score guidance.

D.2 Pseudocode for LSD+

In this section, we present the sampling and training processes of LSD+ in Algorithms 3 and 4. Empirically, we do not learn the coefficients and time schedules in the same training epoch, which empirically leads to training instability issues. Instead, we learn them separately in different training epochs. For brevity, we only list the algorithm that learns the time schedules, rather than the whole learning process.

Algorithm 3 Sampling process of LSD+

Require: Score network s_{θ} , learned time schedule $\{\tau_k\}_{k=0}^M$ for the student sampler with M steps, learned coefficients of the student sampler $\{\Phi(\tau_k)\}_{k=1}^M$, transition rate matrices $\{Q_{\tau_k}\}_{k=0}^M$

- 1: Sample $x_{t_0} \sim \pi$
 - 2: Sample $x_{\tau_{k+1}}$ based on x_{τ_k} and $\Phi(\tau_k)$:
 - 3: $p(x_{\tau_{k+1}}^i | x_{\tau_k}^i) = \delta_{x_{\tau_k}^i}(x_{\tau_{k+1}}^i) + (\tau_k - \tau_{k+1}) Q_{\tau_k}(x_{\tau_k}^i, x_{\tau_{k+1}}^i) \Phi(\tau_k)(s_{\theta}(x_{\tau_k}, \tau_k))_{i, x_{\tau_{k+1}}^i}$
 - 4: $x_{\tau_{k+1}}^i \sim p(x_{\tau_{k+1}}^i | x_{\tau_k}^i)$ for all i
 - 5: **return** x_{ϵ}
-

Algorithm 4 Training process of LSD+ that learns the time schedules

Require: Score network s_θ , frozen teacher sampler Ψ^* with N steps, learnable student sampler Ψ_Φ with M steps, learning rate η , distance metric d , time schedule $\{t_j^*\}_{j=0}^N$ for the teacher sampler, original time schedule $\{t_k\}_{k=0}^M$ for the student sampler (a subsequence of $\{t_j^*\}_{j=0}^N$), transition rate matrices $\{Q_{t_j^*}\}_{j=0}^N$

```
1: Initialize step sizes  $\kappa_k = \frac{t_0 - t_M}{M}$  for  $k = 1, 2, \dots, M$ 
2: Initialize learnable time schedule  $\tau_k = t_0 - \sum_{\ell=1}^k \kappa_\ell$  for  $k = 1, 2, \dots, M$ 
3: while not converged do
4:   Sample  $x_{t_0} \sim \pi$ , set  $x_{t_0}^* \leftarrow x_{t_0}$ 
5:   for  $k = 1$  to  $M$  do
6:     Calculate the state  $x_{\tau_k}$  generated by the student sampler at time  $\tau_k$  and calculate  $s_k = s_\theta(x_{\tau_k}, \tau_k)$ 
7:     Calculate the state  $x_{t_k}^*$  generated by the teacher sampler at time  $t_k$  and calculate the score  $s_k^* = s_\theta(x_{t_k}^*, t_k)$ 
8:   end for
9:   for  $k = 1$  to  $M$  do
10:     $\tilde{L}_k \leftarrow d(\kappa_k s_k, \frac{t_0 - t_N}{N} s_k^*)$ 
11:     $\kappa_k \leftarrow \kappa_k - \eta \nabla_{\kappa_k} \tilde{L}_k$ 
12:     $\tau_k = t_0 - \sum_{l=1}^k \kappa_l$ 
13:   end for
14: end while
15: return  $\{\kappa_k\}_{k=1}^M$ 
```

E Additional empirical details

E.1 Training settings

In this subsection, we provide training settings and implementation details for reproducing our empirical results. Specifically, we set the terminate time ϵ as 0.0001, the total sampling steps N of the teacher sampler as 1024, and the distance metric d as the KL divergence. We set the number of training samples as 64, the training epoch as 20, and the learning rate η as 0.001. All experiments are conducted on an NVIDIA RTX4090 GPU.

E.2 More results on large-scale image datasets

We conduct experiments on ImageNet (256x256) using the MaskGIT [57] architecture as the backbone, incorporating the recently proposed advanced Halton sampler [58] as the sampling method. The results are reported in Table 6, which demonstrates that our LSD+ method yields improved generation performance as measured by the FID metric. We also study the effect of softmax temperature in MaskGIT on ImageNet (256x256) for the image generation task, with the results listed in Table 7.

Table 6: Comparison on ImageNet 256x256 (in terms of FID↓)

Sampler \ NFES				
	4	8	16	32
Halton	14.16	10.15	8.89	6.92
LSD+-Halton	12.78	8.66	7.17	6.32

Table 7: Comparisons on ImageNet 256x256 (in terms of FID↓ when NFE=4).

Sampler \ Temperature			
	$\tau = 0.6$	$\tau = 0.8$	$\tau = 1.0$
Halton	54.05	26.45	14.16
LSD+-Halton	48.29	24.51	12.78

E.3 More results on text generation.

We provide more results on the task of text generation.

We first conduct experiments on predictor-corrector solvers in τ LDR-10 [8] sampler on the SEDD-small backbone, with the results presented in Table 8.

To further break through the limitation of only relying on perplexity to assess generation quality and provide a more thorough evaluation, we conduct experiments on the SDTT-KLD [59] backbone, which utilizes Ancestral as the sampler. We report MAUVE, Perplexity and Entropy scores in Table 9, and these results demonstrate that our LSD+ method usually yields improved generation performance.

Meanwhile, we also integrate LSD+ with ReMasking (ReMDM) [60] on the MDLM [61] backbone, with relevant findings reported in Table 10. The results show that LSD+ can effectively learn to work in conjunction with this method, further improving performance.

To ensure the generalization of our main results, we further re-evaluate our findings on the SEDD and RADD backbones, using Llama-3-8B [62] to assess perplexity. The re-evaluation results are reported in Tables 11 and 12. Moreover, we also integrate our LSD+ with multi-token unmasking heuristics in FastDLLMs [63] and report the corresponding accuracy and throughput metrics in Table 13. These metrics indicate that our methods can improve both sampling quality and speed.

To demonstrate that our method is not limited to smaller models, we conduct experiments on applying LSD+ to larger-scale models including DiffuLLaMA and DiffuGPT [64]. Specifically, we perform sampler distillation on pre-trained DiffuGPT-S and DiffuLLaMA checkpoints, and the results in terms of Perplexity are presented in Tables 14 and 15.

Finally, we conduct experiments on the sampler of DNDM [12], which uses the FairSeq [65] as the backbone, the corresponding perplexity results are reported in Table 16.

Table 8: Comparisons on the SEDD-small backbone.

Sampler	Perplexity($\downarrow$)				Entropy($\uparrow$)			
	16	32	64	128	16	32	64	128
τ LDR-10	443.17	318.44	277.16	199.51	5.63	5.69	5.57	5.24
LSD+- τ LDR-10	205.42	143.58	114.93	90.43	5.58	5.49	5.59	5.44

Table 9: Comparisons on the SDTT-KLD backbone.

Sampler	MAUVE($\uparrow$)			Perplexity($\downarrow$)			Entropy($\uparrow$)		
	8	16	32	8	16	32	8	16	32
Ancestral	0.884	0.912	0.943	110.391	56.652	42.128	5.331	5.285	5.222
LSD+-Ancestral	0.905	0.928	0.951	68.130	36.577	31.597	5.298	5.239	5.226

Table 10: Comparisons on the MDLM backbone.

Sampler	Perplexity($\downarrow$)				Entropy($\uparrow$)			
	16	32	64	128	16	32	64	128
ReMDM	434.08	174.72	85.15	62.33	5.73	5.66	5.48	5.55
LSD+-ReMDM	201.52	102.02	62.97	49.33	5.41	5.42	5.52	5.33

Table 11: Comparisons on SEDD-small backbone (in terms of Perplexity $\downarrow$), judged by LLaMA-3-8B.

Sampler	NFEs			
	8	16	32	64
Euler	116.93	67.43	49.81	46.88
LSD+-Euler	74.65	33.25	30.70	21.64

Table 12: Comparisons on RADD backbone (in terms of Perplexity↓), judged by LLaMA-3-8B.

	NFEs	8	16	32	64
Sampler					
Euler		337.04	216.01	119.25	95.06
LSD+-Euler		130.00	60.90	42.53	35.65

Table 13: Integration with Fast-dLLM on LLaDA. We report accuracy (throughput, token/s).

Benchmark	Samplers	LLaDA	+Cache	+Parallel	Fast-dLLM	LSD+-Fast-dLLM
GSM8K(5-shot)		79.3(6.7)	79.5(21.2)	79.2(16.5)	78.5(54.4)	79.0(62.5)
MATH(4-shot)		33.5(9.1)	33.3(23.7)	33.4(24.8)	33.2(51.7)	33.4(58.1)

Table 14: Comparisons on the DiffuGPT-S backbone (in terms of Perplexity↓).

	NFEs	16	32	64	128
Method					
DiffuGPT-S		117.32	75.19	58.34	37.16
LSD+-DiffuGPT-S		53.95	41.37	32.10	22.25

Table 15: Comparisons on the DiffuLLaMA backbone (in terms of Perplexity↓).

	NFEs	16	32	64	128
Method					
DiffuLLaMA		100.04	69.11	42.17	30.55
LSD+-LLaMA		49.83	34.32	29.18	24.72

Table 16: Comparisons on the FairSeq backbone (in terms of Perplexity↓).

	NFEs	8	16	32	64
Method					
DNDM		919.23	774.92	748.41	622.14
LSD+-DNDM		601.22	554.19	477.10	403.13

E.4 More results on uniform discrete diffusion models

In the main text, we provide comparisons with existing samplers on the absorbing DDMs. In this subsection, we provide additional empirical results on uniform DDMs in Tables 17, 18 and 19. We observe that our methods also outperform existing sampling methods, validating the superiority and robustness of our methods.

Table 17: Comparison of generative perplexity ($\leq$) on the uniform SEDD-small backbone. Best performances are bolded.

	NFEs	8	16	32	64
Sampler					
Euler		467.832	224.954	76.364	54.293
Tweedie		433.590	215.233	70.361	52.364
JYS-Euler		310.329	130.034	60.574	33.951
JYS-Tweedie		308.732	129.843	55.293	30.675
LSD-Euler		160.811	103.346	38.059	23.365
LSD-Tweedie		177.728	101.213	37.682	25.675
LSD+-Euler		157.535	46.448	39.850	23.938
LSD+-Tweedie		128.910	47.080	38.058	21.724

Table 18: Comparison of generative perplexity ($\leq$) on the uniform SEDD-medium backbone. Best performances are bolded.

Sampler \ NFEs	8	16	32	64
Euler	403.654	190.784	80.094	47.885
Tweedie	387.743	182.312	65.853	44.754
JYS-Euler	311.427	121.954	49.912	32.934
JYS-Tweedie	306.089	116.233	45.287	28.192
LSD-Euler	132.209	60.823	28.283	22.956
LSD-Tweedie	118.138	53.591	27.162	21.308
LSD+-Euler	111.870	49.427	25.732	19.623
LSD+-Tweedie	93.196	47.932	27.381	18.119

Table 19: Comparison of generative perplexity ($\leq$) on the uniform RADD backbone. Best performances are bolded.

Sampler \ NFEs	8	16	32	64
Euler	657.732	280.743	157.825	115.743
Tweedie	652.381	277.195	160.045	108.730
LSD-Euler	126.394	65.923	51.197	34.764
LSD-Tweedie	127.098	71.034	44.936	29.834
LSD+-Euler	94.554	41.986	32.832	27.283
LSD+-Tweedie	95.029	45.823	37.900	26.883

F PCA analysis of coefficients

To analyze the the training evolution of learned sampler coefficients $\{\Phi(t_k)\}_{k=1}^M$, we performed Principal Component Analysis (PCA) [66, 67] on $\mathbb{R}^M$ -dimensional coefficient vectors collected at each epoch. Figure 3 visualizes projections onto the top two principal components. The trajectories of learned coefficients show substantial divergence from the vanilla baseline point, empirically demonstrating that optimization yields configurations distinct from fixed scaling. This dynamic learning process, while sensitive to initialization across runs, explores configurations enabling high-fidelity sampling at low NFEs, and the difference between different training runs is relatively small.

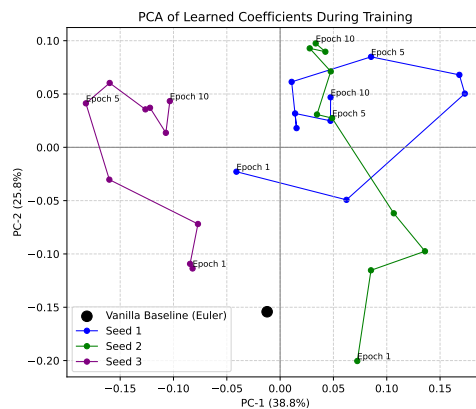


Figure 3: PCA analysis of learned coefficients at each epoch of training.

G Text generation results

In this section, we provide text generation results from LSD+. All results are generated using the Euler sampler on the SEDD-medium backbone.

swagqell on Twitter and for reading! Heath and Hello, that's this information in Continue reading. This's a good idea, I've going to have some ask you who. On I asked you to come in a powerful is, with for then, here were about some of the reasons for that. By any time in Baltimore, in Camden, or Washington, I'll sure you if some sources would and, well, I'd be in I was in Soltas, of Philadelphia (Washington and elsewhere) about the game that's process and his way on the football team. For instance, cornerback Scott Snyder on Washington called "", the the 49ers' 2012 coach (if "" of those season said more byand (that). I'm curious if you would ""—to have a pragmatic thing that actually assures a "" the right of the .election.article" level activists and that it's not, to to the system of American columns, to this qualified as follows in a If I've the. Before the election for the days where the didn't ask, and the Republican candidates to lead at all—without clear direction and without first getting a record in light of the constituents' from which I've taken my time to think about such thing and I'd not be too unapologetic about spending a few days (Don't know, the rise of the headed system, I's in curious whether he wasn't in. Here thought was in response case. This is the same is of one's attempt to , so funny enough, long as it remains without a". The 55 lot of the plans, so as more of an in-house plan to than ervative", a plan to interest increase and saying, that should function as a thought. As for whether Obama won. , he's currently third in the, in Texas, nominee. I wrote about the new bill in an article-ed. Even if the "If" you believe in him" is. you be the the, which was in favor of repealing it. In terms of the game being can self-destruct of Romney's spokesperson, Mitt Romney. I was you who did it. case, it was passed last which was the new health insurance law. The law was Obamacare, the American Health Care Act. I think was about to pre the Obamacar

Figure 4: The generated text for NFE=8.

and that the federal programs have been pay out for additional much. It was agreed at the summit that there will be increases in the mean time and that they will need to look back and make full pay."We have got to stop before you get started," Boehner said. "And not a few of us are going to have a budget in the some, because what we have't started looking at that's just beginning. We need to pay for all those programs -- our normal budget plus we will make work toward the end of the year."In order to be trusted in conjunction with the austerity of the entitlement will begin almost immediately. We are the nation -- President Obama will have said it and the Republicans, for that matter, won't have said it -- to have to face.Obama has done everything in his power not to acknowledge this possibility. There maintains a veneer of honesty to his administration, but else he displayed is almost entirely to blame on the cliff.On a united loan from the House Party, Obama has paid—Both parties in ownership the national debt.Specifically: the companies that profit from, and through whose contracts, the money government owes, and it is owed more. It has given a flying strip limit"s. It also has debt small companies, the United States on this earth.In America, the government has had enormous responsibility to America. I can tell Washington DC really has done the least for America over the world.Perhaps it Washington, it will be able to assist and alter the government whether or not they have created something to the taxpayer the financial representation of this debt feels symbolically self sufficient. But their future goes far beyond the effects of this austerity in many ways. For one, of the debt will create an opportunity for a new generation to get out of the burden of the debt. If a de-biz for all tax plan is publicly announced.Beyond that is however, self-governance will take, and the consequences mean be -- and is obvious. from here to America, as I wrote here, recently, debt

Figure 5: The generated text for NFE=16.

it's just me, it's bad feelings. He tells aware of sadistic behavior that is a joy to watch as a missionary and me when I see some right-handed people validating Mormonism to fall in love, and some I know are not, as I tell him. I teach me the left- hands, so does he. I love him for having his son. "He's often running riot," he says. "Like a kid bit in a get-out-of-way mode." I help him study a lot about that and I teach him about them. As a missionary I'm all out in search of sacred books, literally and figuratively. Every. I help him learn and I learn of in the way we prepare for mission. Again, these are complements of his ministry. "Man is man and she is fairly simple too, because I can see the power of that saying. when I'm all over him to evangel I really struggle to deal with it. But, I feel grounded in my own life. Though I may know what to read, I study scriptures, read sacred books and now when your eyes see this as clearly as I do, that counts as a difference. that you feel he is lonely. It, beautiful and is that, when you never, you realize he is brokenhearted. But from the best of his, it's just a beautiful reflection of his desire to experience in his daily life and the desires and needs of those he condemns and tries to project them onto young people. Either way, I welcome it. He is a human being and—I love him just as well as I can. I will not deny his character as a kind, humble, and hospitable man who loves us in many different ways, tender-hearted and will be some the greatest on the feet of the earth. He loves us all, but as he discerns more and more about people. That said, I'll love him a part of his missionary call. He knows I shouldn't change my mind and when he has come to be the Savior, every response creates rays of change in my heart. He loves us about people every day in the work of God, through his mission to "correct the whole world." The better he can, the more he can come to understand people. Sometimes, he says that he casts a shadow for th

Figure 6: The generated text for NFE=32.

we're going to let the players know that some new players have a responsibility to the fans. Q: Do you have a idea of it like this, how to the teams with the fans? No, before that, went into this season, we were at the moment we were heading for the playoffs in this game. The only difference is that, at the beginning, we had seven to go. in that moment, you put on the match with the players that popped up from the field with the players only, which is hard to do, because there are so many to deal with that we had to mess around with. And there, it makes it over to the training room" the training room. I think everyone on the team is in the training room. In the training room, the players come out here, the players come to the standing on the ground, the pass the ball. When the head of the player passes, the player has to push-up their head on the ground, if it from the pass. then all the players, who are going to pass when the touch the ball, have to push-up their heads on this when they receive the ball before they receive the ball. The organization knows that the match is not won not just by the fans' support, so that first half the fans' fans decided to become fans in the training room, where the players can help provide the input by looking at the defender's tips. The defense is going to score the goal's at the match's end. That will be based on the first half. This is when we know it's going to be difficult for them. The team's title is going to be for the people. That's the title, for the people. Q: That's the talk that you gave the players this past week for in Tuesday. A number of times they didn't really set up routines for the game. Yes. They didn't know. I mean: I don't know, in the team group, I just think that they all the whatever it is, they just got to prepare themselves for the easiest any time, and when we play the game for a week, then we'll go to something to the rest. For the players, we are the ones to have the preparation routine, the ones to

Figure 7: The generated text for NFE=64.