

Diffusion Models: A Comprehensive Survey of Methods and Applications

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Diffusion models have emerged as a powerful new family of deep generative models with record-breaking performance in many applications, including image synthesis, video generation, and molecule design. In this survey, we provide an overview of the rapidly expanding body of work on diffusion models, categorizing the research into three key areas: efficient sampling, improved likelihood estimation, and handling data with special structures. We also discuss the potential for combining diffusion models with other generative models for enhanced results. We further review the wide-ranging applications of diffusion models in fields spanning from computer vision, natural language processing, temporal data modeling, to interdisciplinary applications in other scientific disciplines. This survey aims to provide a contextualized, in-depth look at the state of diffusion models, identifying the key areas of focus and pointing to potential areas for further exploration. Github: <https://github.com/YangLing0818/Diffusion-Models-Papers-Survey-Taxonomy>.

CCS Concepts: • Computing methodologies → Computer vision tasks; Natural language generation; Machine learning approaches.

Additional Key Words and Phrases: Generative Models, Diffusion Models, Score-Based Generative Models, Stochastic Differential Equations

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1 INTRODUCTION

Diffusion models [90, 215, 220, 225] have emerged as the new state-of-the-art family of deep generative models. They have broken the long-time dominance of generative adversarial networks (GANs) [73] in the challenging task of image synthesis [49, 90, 220, 225] and have also shown potential in a variety of domains, ranging from computer vision [2, 9, 17, 20, 91, 93, 115, 118, 137, 151, 161, 174, 200, 202, 245, 267, 268, 282, 289], natural language processing [6, 96, 141, 207, 272], temporal data modeling [1, 29, 126, 192, 230, 260], multi-modal modeling [7, 186, 198, 201, 287], robust machine learning [16, 24, 114, 240, 270], to interdisciplinary applications in fields such as computational chemistry [3, 94, 107, 133, 135, 153, 256] and medical image reconstruction [22, 36–38, 42, 159, 178, 224, 257].

Numerous methods have been developed to improve diffusion models, either by enhancing empirical performance [166, 217, 221] or by extending the model’s capacity from a theoretical perspective [145, 146, 219, 225, 277]. Over the past two years, the body of research on diffusion models has grown significantly, making it increasingly challenging for new researchers to stay abreast of the recent developments in the field. Additionally, the sheer volume of work can obscure major trends and hinder further research progress. This survey aims to address these problems by providing a comprehensive overview of the state of diffusion model research, categorizing various approaches, and highlighting key advances. We hope this survey to serve as a helpful entry point for researchers new to the field while providing a broader perspective for experienced researchers.

In this paper, we first explain the foundations of diffusion models (Section 2), providing a brief but self-contained introduction to three predominant formulations: denoising diffusion probabilistic models (DDPMs) [90, 215], score-based generative models (SGMs) [220, 221], and stochastic differential equations (Score SDEs) [113, 219, 225]. Key to all these approaches is to progressively perturb data with intensifying random noise (called the “diffusion” process),



Fig. 1. Taxonomy of diffusion models variants (in Sections 3 to 5), connections with other generative models (in Section 6), applications of diffusion models (in Section 7), and future directions (in Section 8).

then successively remove noise to generate new data samples. We clarify how they work under the same principle of diffusion and explain how these three models are connected and can be reduced to one another.

Next, we present a taxonomy of recent research that maps out the field of diffusion models, categorizing it into three key areas: efficient sampling (Section 3), improved likelihood estimation (Section 4), and methods for handling data with special structures (Section 5), such as relational data, data with permutation/rotational invariance, and data residing on manifolds. We further examine the models by breaking each category into more detailed sub-categories, as illustrated

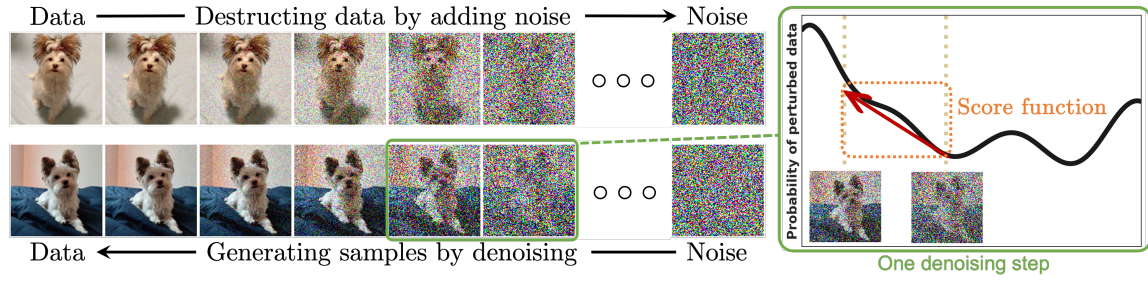


Fig. 2. Diffusion models smoothly perturb data by adding noise, then reverse this process to generate new data from noise. Each denoising step in the reverse process typically requires estimating the score function (see the illustrative figure on the right), which is a gradient pointing to the directions of data with higher likelihood and less noise.

in Fig. 1. In addition, we discuss the connections of diffusion models to other deep generative models (Section 6), including variational autoencoders (VAEs) [123, 195], generative adversarial networks (GANs) [73], normalizing flows [50, 52, 175, 197], autoregressive models [237], and energy-based models (EBMs) [132, 223]. By combining these models with diffusion models, researchers have the potential to achieve even stronger performance.

Following that, our survey reviews six major categories of application that diffusion models have been applied to in the existing research (Section 7): computer vision, natural language process, temporal data modeling, multi-modal learning, robust learning, and interdisciplinary applications. For each task, we provide a definition, describe how diffusion models can be employed to address it and summarize relevant previous work. We conclude our paper (Sections 8 and 9) by providing an outlook on possible future directions for this exciting new area of research.

2 FOUNDATIONS OF DIFFUSION MODELS

Diffusion models are a family of probabilistic generative models that progressively destruct data by injecting noise, then learn to reverse this process for sample generation. We present the intuition of diffusion models in Fig. 2. Current research on diffusion models is mostly based on three predominant formulations: denoising diffusion probabilistic models (DDPMs) [90, 166, 215], score-based generative models (SGMs) [220, 221], and stochastic differential equations (Score SDEs) [219, 225]. We give a self-contained introduction to these three formulations in this section, while discussing their connections with each other along the way.

2.1 Denoising Diffusion Probabilistic Models (DDPMs)

A *denoising diffusion probabilistic model* (DDPM) [90, 215] makes use of two Markov chains: a forward chain that perturbs data to noise, and a reverse chain that converts noise back to data. The former is typically hand-designed with the goal to transform any data distribution into a simple prior distribution (e.g., standard Gaussian), while the latter Markov chain reverses the former by learning transition kernels parameterized by deep neural networks. New data points are subsequently generated by first sampling a random vector from the prior distribution, followed by ancestral sampling through the reverse Markov chain [125].

Formally, given a data distribution $\mathbf{x}_0 \sim q(\mathbf{x}_0)$, the forward Markov process generates a sequence of random variables $\mathbf{x}_1, \mathbf{x}_2 \dots \mathbf{x}_T$ with transition kernel $q(\mathbf{x}_t \mid \mathbf{x}_{t-1})$. Using the chain rule of probability and the Markov property, we can

factorize the joint distribution of $\mathbf{x}_1, \mathbf{x}_2 \dots \mathbf{x}_T$ conditioned on \mathbf{x}_0 , denoted as $q(\mathbf{x}_1, \dots, \mathbf{x}_T | \mathbf{x}_0)$, into

$$q(\mathbf{x}_1, \dots, \mathbf{x}_T | \mathbf{x}_0) = \prod_{t=1}^T q(\mathbf{x}_t | \mathbf{x}_{t-1}). \quad (1)$$

In DDPMs, we handcraft the transition kernel $q(\mathbf{x}_t | \mathbf{x}_{t-1})$ to incrementally transform the data distribution $q(\mathbf{x}_0)$ into a tractable prior distribution. One typical design for the transition kernel is Gaussian perturbation, and the most common choice for the transition kernel is

$$q(\mathbf{x}_t | \mathbf{x}_{t-1}) = \mathcal{N}(\mathbf{x}_t; \sqrt{1 - \beta_t} \mathbf{x}_{t-1}, \beta_t \mathbf{I}), \quad (2)$$

where $\beta_t \in (0, 1)$ is a hyperparameter chosen ahead of model training. We use this kernel to simply our discussion here, although other types of kernels are also applicable in the same vein. As observed by Sohl-Dickstein et al. (2015) [215], this Gaussian transition kernel allows us to marginalize the joint distribution in Eq. (1) to obtain the analytical form of $q(\mathbf{x}_t | \mathbf{x}_0)$ for all $t \in \{0, 1, \dots, T\}$. Specifically, with $\alpha_t := 1 - \beta_t$ and $\bar{\alpha}_t := \prod_{s=0}^t \alpha_s$, we have

$$q(\mathbf{x}_t | \mathbf{x}_0) = \mathcal{N}(\mathbf{x}_t; \sqrt{\bar{\alpha}_t} \mathbf{x}_0, (1 - \bar{\alpha}_t) \mathbf{I}). \quad (3)$$

Given \mathbf{x}_0 , we can easily obtain a sample of \mathbf{x}_t by sampling a Gaussian vector $\boldsymbol{\epsilon} \sim \mathcal{N}(\mathbf{0}, \mathbf{I})$ and applying the transformation

$$\mathbf{x}_t = \sqrt{\bar{\alpha}_t} \mathbf{x}_0 + \sqrt{1 - \bar{\alpha}_t} \boldsymbol{\epsilon}. \quad (4)$$

When $\bar{\alpha}_T \approx 0$, \mathbf{x}_T is almost Gaussian in distribution, so we have $q(\mathbf{x}_T) := \int q(\mathbf{x}_T | \mathbf{x}_0) q(\mathbf{x}_0) d\mathbf{x}_0 \approx \mathcal{N}(\mathbf{x}_T; \mathbf{0}, \mathbf{I})$.

Intuitively speaking, this forward process slowly injects noise to data until all structures are lost. For generating new data samples, DDPMs start by first generating an unstructured noise vector from the prior distribution (which is typically trivial to obtain), then gradually remove noise therein by running a learnable Markov chain in the reverse time direction. Specifically, the reverse Markov chain is parameterized by a prior distribution $p(\mathbf{x}_T) = \mathcal{N}(\mathbf{x}_T; \mathbf{0}, \mathbf{I})$ and a learnable transition kernel $p_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t)$. We choose the prior distribution $p(\mathbf{x}_T) = \mathcal{N}(\mathbf{x}_T; \mathbf{0}, \mathbf{I})$ because the forward process is constructed such that $q(\mathbf{x}_T) \approx \mathcal{N}(\mathbf{x}_T; \mathbf{0}, \mathbf{I})$. The learnable transition kernel $p_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t)$ takes the form of

$$p_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t) = \mathcal{N}(\mathbf{x}_{t-1}; \mu_\theta(\mathbf{x}_t, t), \Sigma_\theta(\mathbf{x}_t, t)) \quad (5)$$

where θ denotes model parameters, and the mean $\mu_\theta(\mathbf{x}_t, t)$ and variance $\Sigma_\theta(\mathbf{x}_t, t)$ are parameterized by deep neural networks. With this reverse Markov chain in hand, we can generate a data sample \mathbf{x}_0 by first sampling a noise vector $\mathbf{x}_T \sim p(\mathbf{x}_T)$, then iteratively sampling from the learnable transition kernel $\mathbf{x}_{t-1} \sim p_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t)$ until $t = 1$.

Key to the success of this sampling process is training the reverse Markov chain to match the actual time reversal of the forward Markov chain. That is, we have to adjust the parameter θ so that the joint distribution of the reverse Markov chain $p_\theta(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T) := p(\mathbf{x}_T) \prod_{t=1}^T p_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t)$ closely approximates that of the forward process $q(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T) := q(\mathbf{x}_0) \prod_{t=1}^T q(\mathbf{x}_t | \mathbf{x}_{t-1})$ (Eq. (1)). This is achieved by minimizing the Kullback-Leibler (KL)

divergence between these two:

$$\text{KL}(q(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T) \parallel p_\theta(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T)) \quad (6)$$

$$\stackrel{(i)}{=} -\mathbb{E}_{q(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T)} [\log p_\theta(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T)] + \text{const} \quad (7)$$

$$\stackrel{(ii)}{=} \mathbb{E}_{q(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T)} \underbrace{\left[-\log p(\mathbf{x}_T) - \sum_{t=1}^T \log \frac{p_\theta(\mathbf{x}_{t-1} \mid \mathbf{x}_t)}{q(\mathbf{x}_t \mid \mathbf{x}_{t-1})} \right]}_{:= -L_{\text{VLB}}(\mathbf{x}_0)} + \text{const} \quad (8)$$

$$\stackrel{(iii)}{\geq} \mathbb{E} [-\log p_\theta(\mathbf{x}_0)] + \text{const}, \quad (9)$$

where (i) is from the definition of KL divergence, (ii) is from the fact that $q(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T)$ and $p_\theta(\mathbf{x}_0, \mathbf{x}_1, \dots, \mathbf{x}_T)$ are both products of distributions, and (iii) is from Jensen's inequality. The first term in Eq. (8) is the variational lower bound (VLB) of the log-likelihood of the data \mathbf{x}_0 , a common objective for training probabilistic generative models. We use "const" to symbolize a constant that does not depend on the model parameter θ and hence does not affect optimization. The objective of DDPM training is to maximize the VLB (or equivalently, minimizing the negative VLB), which is particularly easy to optimize because it is a sum of independent terms, and can thus be estimated efficiently by Monte Carlo sampling [164] and optimized effectively by stochastic optimization [226].

Ho et al. (2020) [90] propose to reweight various terms in L_{VLB} for better sample quality and noticed an important equivalence between the resulting loss function and the training objective for noise-conditional score networks (NCSNs), one type of *score-based generative models*, in Song and Ermon [220]. The loss in [90] takes the form of

$$\mathbb{E}_{t \sim \mathcal{U}[\![1, T]\!], \mathbf{x}_0 \sim q(\mathbf{x}_0), \boldsymbol{\epsilon} \sim \mathcal{N}(\mathbf{0}, \mathbf{I})} [\lambda(t) \|\boldsymbol{\epsilon} - \boldsymbol{\epsilon}_\theta(\mathbf{x}_t, t)\|^2] \quad (10)$$

where $\lambda(t)$ is a positive weighting function, \mathbf{x}_t is computed from \mathbf{x}_0 and $\boldsymbol{\epsilon}$ by Eq. (4), $\mathcal{U}[\![1, T]\!]$ is a uniform distribution over the set $\{1, 2, \dots, T\}$, and $\boldsymbol{\epsilon}_\theta$ is a deep neural network with parameter θ that predicts the noise vector $\boldsymbol{\epsilon}$ given \mathbf{x}_t and t . This objective reduces to Eq. (8) for a particular choice of the weighting function $\lambda(t)$, and has the same form as the loss of denoising score matching over multiple noise scales for training score-based generative models [220], another formulation of diffusion models to be discussed in the next section.

2.2 Score-Based Generative Models (SGMs)

At the core of score-based generative models [220, 221] is the concept of (*Stein*) score (a.k.a., score or score function) [101]. Given a probability density function $p(\mathbf{x})$, its score function is defined as the gradient of the log probability density $\nabla_{\mathbf{x}} \log p(\mathbf{x})$. Unlike the commonly used *Fisher score* $\nabla_\theta \log p_\theta(\mathbf{x})$ in statistics, the Stein score considered here is a function of the data \mathbf{x} rather than the model parameter θ . It is a vector field that points to directions along which the probability density function has the largest growth rate.

The key idea of score-based generative models (SGMs) [220] is to perturb data with a sequence of intensifying Gaussian noise and jointly estimate the score functions for all noisy data distributions by training a deep neural network model conditioned on noise levels (called a noise-conditional score network, NCSN, in [220]). Samples are generated by chaining the score functions at decreasing noise levels with score-based sampling approaches, including Langevin Monte Carlo [81, 110, 176, 220, 225], stochastic differential equations [109, 225], ordinary differential equations [113, 146, 219, 225, 277], and their various combinations [225]. Training and sampling are completely decoupled

in the formulation of score-based generative models, so one can use a multitude of sampling techniques after the estimation of score functions.

With similar notations in Section 2.1, we let $q(\mathbf{x}_0)$ be the data distribution, and $0 < \sigma_1 < \sigma_2 < \dots < \sigma_t < \dots < \sigma_T$ be a sequence of noise levels. A typical example of SGMs involves perturbing a data point \mathbf{x}_0 to \mathbf{x}_t by the Gaussian noise distribution $q(\mathbf{x}_t | \mathbf{x}_0) = \mathcal{N}(\mathbf{x}_t; \mathbf{x}_0, \sigma_t^2 \mathbf{I})$. This yields a sequence of noisy data densities $q(\mathbf{x}_1), q(\mathbf{x}_2), \dots, q(\mathbf{x}_T)$, where $q(\mathbf{x}_t) := \int q(\mathbf{x}_t)q(\mathbf{x}_0)d\mathbf{x}_0$. A noise-conditional score network is a deep neural network $s_\theta(\mathbf{x}, t)$ trained to estimate the score function $\nabla_{\mathbf{x}_t} \log q(\mathbf{x}_t)$. Learning score functions from data (a.k.a., score estimate) has established techniques such as score matching [101], denoising score matching [188, 189, 238], and sliced score matching [222], so we can directly employ one of them to train our noise-conditional score networks from perturbed data points. For example, with denoising score matching and similar notations in Eq. (10), the training objective is given by

$$\mathbb{E}_{t \sim \mathcal{U}[\![1, T]\!], \mathbf{x}_0 \sim q(\mathbf{x}_0), \mathbf{x}_t \sim q(\mathbf{x}_t | \mathbf{x}_0)} \left[\lambda(t) \sigma_t^2 \|\nabla_{\mathbf{x}_t} \log q(\mathbf{x}_t) - s_\theta(\mathbf{x}_t, t)\|^2 \right] \quad (11)$$

$$\stackrel{(i)}{=} \mathbb{E}_{t \sim \mathcal{U}[\![1, T]\!], \mathbf{x}_0 \sim q(\mathbf{x}_0), \mathbf{x}_t \sim q(\mathbf{x}_t | \mathbf{x}_0)} \left[\lambda(t) \sigma_t^2 \|\nabla_{\mathbf{x}_t} \log q(\mathbf{x}_t | \mathbf{x}_0) - s_\theta(\mathbf{x}_t, t)\|^2 \right] + \text{const} \quad (12)$$

$$\stackrel{(ii)}{=} \mathbb{E}_{t \sim \mathcal{U}[\![1, T]\!], \mathbf{x}_0 \sim q(\mathbf{x}_0), \mathbf{x}_t \sim q(\mathbf{x}_t | \mathbf{x}_0)} \left[\lambda(t) \left\| -\frac{\mathbf{x}_t - \mathbf{x}_0}{\sigma_t} - \sigma_t s_\theta(\mathbf{x}_t, t) \right\|^2 \right] + \text{const} \quad (13)$$

$$\stackrel{(iii)}{=} \mathbb{E}_{t \sim \mathcal{U}[\![1, T]\!], \mathbf{x}_0 \sim q(\mathbf{x}_0), \boldsymbol{\epsilon} \sim \mathcal{N}(\mathbf{0}, \mathbf{I})} \left[\lambda(t) \|\boldsymbol{\epsilon} + \sigma_t s_\theta(\mathbf{x}_t, t)\|^2 \right] + \text{const}, \quad (14)$$

where (i) is derived by [238], (ii) is from the assumption that $q(\mathbf{x}_t | \mathbf{x}_0) = \mathcal{N}(\mathbf{x}_t; \mathbf{x}_0, \sigma_t^2 \mathbf{I})$, and (iii) is from the fact that $\mathbf{x}_t = \mathbf{x}_0 + \sigma_t \boldsymbol{\epsilon}$. Again, we denote by $\lambda(t)$ a positive weighting function, and “const” a constant that does not depend on the trainable parameter θ . Comparing Eq. (14) with Eq. (10), it is clear that the training objectives of DDPMs and SGMs are equivalent, once we set $\boldsymbol{\epsilon}_\theta(\mathbf{x}, t) = -\sigma_t s_\theta(\mathbf{x}, t)$.

For sample generation, SGMs leverage iterative approaches to produce samples from $s_\theta(\mathbf{x}, T), s_\theta(\mathbf{x}, T-1), \dots, s_\theta(\mathbf{x}, 0)$ in succession. Many sampling approaches exist due to the decoupling of training and inference in SGMs, some of which are discussed in the next section. Here we introduce the first sampling method for SGMs, called annealed Langevin dynamics (ALD) [220]. Let N be the number of iterations per time step and $s_t > 0$ be the step size. We first initialize ALD with $\mathbf{x}_T^{(N)} \sim \mathcal{N}(\mathbf{0}, \mathbf{I})$, then apply Langevin Monte Carlo for $t = T, T-1, \dots, 1$ one after the other. At each time step $0 \leq t < T$, we start with $\mathbf{x}_t^{(0)} = \mathbf{x}_{t+1}^{(N)}$, before iterating according to the following update rule for $i = 0, 1, \dots, N-1$:

$$\begin{aligned} \boldsymbol{\epsilon}^{(i)} &\leftarrow \mathcal{N}(\mathbf{0}, \mathbf{I}) \\ \mathbf{x}_t^{(i+1)} &\leftarrow \mathbf{x}_t^{(i)} + \frac{1}{2} s_t s_\theta(\mathbf{x}_t^{(i)}, t) + \sqrt{s_t} \boldsymbol{\epsilon}^{(i)}. \end{aligned}$$

The theory of Langevin Monte Carlo [176] guarantees that as $s_t \rightarrow 0$ and $N \rightarrow \infty$, $\mathbf{x}_0^{(N)}$ becomes a valid sample from the data distribution $q(\mathbf{x}_0)$.

2.3 Stochastic Differential Equations (Score SDEs)

DDPMs and SGMs can be further generalized to the case of infinite time steps or noise levels, where the perturbation and denoising processes are solutions to stochastic differential equations (SDEs). We call this formulation *Score SDE* [225], as it leverages SDEs for noise perturbation and sample generation, and the denoising process requires estimating score functions of noisy data distributions.

Score SDEs perturb data to noise with a diffusion process governed by the following stochastic differential equation (SDE) [225]:

$$dx = f(x, t)dt + g(t)d\mathbf{w} \quad (15)$$

where $f(x, t)$ and $g(t)$ are diffusion and drift functions of the SDE, and \mathbf{w} is a standard Wiener process (a.k.a., Brownian motion). The forward processes in DDPMs and SGMs are both discretizations of this SDE. As demonstrated in Song et al. (2020) [225], for DDPMs, the corresponding SDE is:

$$dx = -\frac{1}{2}\beta(t)xdt + \sqrt{\beta(t)}d\mathbf{w} \quad (16)$$

where $\beta(\frac{t}{T}) = T\beta_t$ as T goes to infinity; and for SGMs, the corresponding SDE is given by

$$dx = \sqrt{\frac{d[\sigma(t)^2]}{dt}}d\mathbf{w}, \quad (17)$$

where $\sigma(\frac{t}{T}) = \sigma_t$ as T goes to infinity. Here we use $q_t(x)$ to denote the distribution of x_t in the forward process.

Crucially, for any diffusion process in the form of Eq. (15), Anderson [4] shows that it can be reversed by solving the following reverse-time SDE:

$$dx = [f(x, t) - g(t)^2 \nabla_x \log q_t(x)] dt + g(t)d\bar{\mathbf{w}} \quad (18)$$

where $\bar{\mathbf{w}}$ is a standard Wiener process when time flows backwards, and dt denotes an infinitesimal negative time step. The solution trajectories of this reverse SDE share the same marginal densities as those of the forward SDE, except that they evolve in the opposite time direction [225]. Intuitively, solutions to the reverse-time SDE are diffusion processes that gradually convert noise to data. Moreover, Song et al. (2020) [225] prove the existence of an ordinary differential equation (ODE), namely the *probability flow ODE*, whose trajectories have the same marginals as the reverse-time SDE. The probability flow ODE is given by:

$$dx = \left[f(x, t) - \frac{1}{2}g(t)^2 \nabla_x \log q_t(x) \right] dt. \quad (19)$$

Both the reverse-time SDE and the probability flow ODE allow sampling from the same data distribution as their trajectories have the same marginals.

Once the score function at each time step t , $\nabla_x \log q_t(x)$, is known, we unlock both the reverse-time SDE (Eq. (18)) and the probability flow ODE (Eq. (19)) and can subsequently generate samples by solving them with various numerical techniques, such as annealed Langevin dynamics [220] (cf., Section 2.2), numerical SDE solvers [109, 225], numerical ODE solvers [113, 146, 217, 225, 277], and predictor-corrector methods (combination of MCMC and numerical ODE/SDE solvers) [225]. Like in SGMs, we parameterize a time-dependent score model $s_\theta(x_t, t)$ to estimate the score function by generalizing the score matching objective in Eq. (14) to continuous time, leading to the following objective:

$$\mathbb{E}_{t \sim \mathcal{U}[0, T], x_0 \sim q(x_0), x_t \sim q(x_t | x_0)} \left[\lambda(t) \| s_\theta(x_t, t) - \nabla_{x_t} \log q_{0t}(x_t | x_0) \|^2 \right], \quad (20)$$

where $\mathcal{U}[0, T]$ denotes the uniform distribution over $[0, T]$, and the remaining notations follow Eq. (14).

Subsequent research on diffusion models focuses on improving these classical approaches (DDPMs, SGMs, and Score SDEs) from three major directions: faster and more efficient sampling, more accurate likelihood and density estimation, and handling data with special structures (such as permutation invariance, manifold structures, and discrete data). We survey each direction extensively in the next three sections (Sections 3 to 5). In Table 1, we list the three types of

diffusion models with more detailed categorization, corresponding articles and years, under continuous and discrete time settings.

Table 1. Three types of diffusion models are listed with corresponding articles and years, under continuous and discrete settings.

Primary	Secondary	Tertiary	Article	Year	Setting
Learning-Free Sampling	SDE Solvers		Song et al. [225]	2020	Continuous
			Jolicoeur et al. [110]	2021	Continuous
			Jolicoeur et al. [109]	2021	Continuous
			Chuang et al. [37]	2022	Continuous
			Song et al. [220]	2019	Continuous
			Karras et al. [113]	2022	Continuous
	ODE Solvers		Dockhorn et al. [54]	2021	Continuous
			Song et al. [217]	2020	Continuous
			Zhang et al. [278]	2022	Continuous
			Karras et al. [113]	2022	Continuous
Efficient Sampling	Optimized Discretization		Lu et al. [146]	2022	Continuous
			Zhang et al. [277]	2022	Continuous
			Liu et al. [142]	2021	Continuous
			Watson et al. [243]	2021	Discrete
			Watson et al. [242]	2021	Discrete
	Learning-Based Sampling		Dockhorn et al. [55]	2021	Continuous
			Salimans et al. [203]	2021	Discrete
			Luhman et al. [148]	2021	Discrete
			Lyu et al. [156]	2022	Discrete
			Zheng et al. [284]	2022	Discrete
Improved Likelihood	Noise Schedule Optimization	Noise Schedule Optimization	Nichol et al. [166]	2021	Discrete
			Kingma et al. [121]	2021	Discrete
			Bao et al. [8]	2021	Discrete
			Nichol et al. [166]	2021	Discrete
			Song et al. [219]	2021	Continuous
	Exact Likelihood Computation	Exact Likelihood Computation	Huang et al. [98]	2021	Continuous
			Song et al. [225]	2020	Continuous
			Lu et al. [145]	2022	Continuous
			Vahdat et al. [234]	2021	Continuous
			Wehenkel et al. [244]	2021	Discrete
Data with Special Structures	Manifold Structures	Learned Manifolds	Ramesh et al. [186]	2022	Discrete
			Rombach et al. [198]	2022	Discrete
			Bortoli et al. [45]	2022	Continuous
			Huang et al. [97]	2022	Continuous
			Niu et al. [171]	2020	Discrete
	Discrete Data	Discrete Data	Jo et al. [108]	2022	Continuous
			Shi et al. [210]	2022	Continuous
			Xu et al. [259]	2021	Discrete
			Sohl et al. [215]	2015	Discrete
			Austin et al. [6]	2021	Discrete

3 DIFFUSION MODELS WITH EFFICIENT SAMPLING

Generating samples from diffusion models typically demands iterative approaches that involve a large number of evaluation steps. A great deal of recent work has focused on speeding up the sampling process while also improving quality of the resulting samples. We classify these efficient sampling methods into two main categories: those that do not involve learning (learning-free sampling) and those that require an additional learning process after the diffusion model has been trained (learning-based sampling).

3.1 Learning-Free Sampling

Many samplers for diffusion models rely on discretizing either the reverse-time SDE present in Eq. (18) or the probability flow ODE from Eq. (19). Since the cost of sampling increases proportionally with the number of discretized time steps, many researchers have focused on developing discretization schemes that reduce the number of time steps while also minimizing discretization errors.

3.1.1 SDE Solvers. The generation process of DDPM [90, 215] can be viewed as a particular discretization of the reverse-time SDE. As discussed in Section 2.3, the forward process of DDPM discretizes the SDE in Eq. (16), whose corresponding reverse SDE takes the form of

$$dx = -\frac{1}{2}\beta(t)(x_t - \nabla_{x_t} \log q_t(x_t))dt + \sqrt{\beta(t)}dw \quad (21)$$

Song et al. (2020) [225] show that the reverse Markov chain defined by Eq. (5) amounts to a numerical SDE solver for Eq. (21).

Noise-Conditional Score Networks (NCSNs) [220] and Critically-Damped Langevin Diffusion (CLD) [54] both solve the reverse-time SDE with inspirations from Langevin dynamics. In particular, NCSNs leverage annealed Langevin dynamics (ALD, *cf.*, Section 2.2) to iteratively generate data while smoothly reducing noise level until the generated data distribution converges to the original data distribution. Although the sampling trajectories of ALD are not exact solutions to the reverse-time SDE, they have the correct marginals and hence produce correct samples under the assumption that Langevin dynamics converges to its equilibrium at every noise level. The method of ALD is further improved by Consistent Annealed Sampling (CAS) [110], a score-based MCMC approach with better scaling of time steps and added noise. Inspired by statistical mechanics, CLD proposes an augmented SDE with an auxiliary velocity term resembling underdamped Langevin diffusion. To obtain the time reversal of the extended SDE, CLD only needs to learn the score function of the conditional distribution of velocity given data, arguably easier than learning scores of data directly. The added velocity term is reported to improve sampling speed as well as quality.

The reverse diffusion method proposed in [225] discretizes the reverse-time SDE in the same way as the forward one. For any one-step discretization of the forward SDE, one may write the general form below:

$$x_{i+1} = x_i + f_i(x_i) + g_i z_i, \quad i = 0, 1, \dots, N-1 \quad (22)$$

where $z_i \sim \mathcal{N}(0, I)$, f_i and g_i are determined by drift/diffusion coefficients of the SDE and the discretization scheme. Reverse diffusion proposes to discretize the reverse-time SDE similarly to the forward SDE, *i.e.*,

$$x_i = x_{i+1} - f_{i+1}(x_{i+1}) + g_{i+1} g_{i+1}^t s_{\theta^*}(x_{i+1}, t_{i+1}) + g_{i+1} z_i \quad i = 0, 1, \dots, N-1 \quad (23)$$

where $s_{\theta^*}(x_i, t_i)$ is the trained noise-conditional score model. Song et al. (2020) [225] prove that the reverse diffusion method is a numerical SDE solver for the reverse-time SDE in Eq. (18). This process can be applied to any types of forward SDEs, and empirical results indicate this sampler performs slightly better than DDPM [225] for a particular type of SDEs called the VP-SDE.

Jolicoeur-Martineau et al. (2021) [109] develop an SDE solver with adaptive step sizes for faster generation. The step size is controlled by comparing the output of a high-order SDE solver versus the output of a low-order SDE solver. At each time step, the high- and low-order solvers generate new sample x'_{high} and x'_{low} from the previous sample x'_{prev} respectively. The step size is then adjusted by comparing the difference between the two samples. If x'_{high} and x'_{low} are similar, the algorithm will return x'_{high} and then increase the step size. The similarity between x'_{high} and x'_{low} is

measured by:

$$E_q = \left\| \frac{\mathbf{x}'_{\text{low}} - \mathbf{x}'_{\text{high}}}{\delta(\mathbf{x}', \mathbf{x}'_{\text{prev}})} \right\|^2 \quad (24)$$

where $\delta(\mathbf{x}'_{\text{low}}, \mathbf{x}'_{\text{prev}}) := \max(\epsilon_{abs}, \epsilon_{rel} \max(|\mathbf{x}'_{\text{low}}|, |\mathbf{x}'_{\text{prev}}|))$, and ϵ_{abs} and ϵ_{rel} are absolute and relative tolerances.

The predictor-corrector method proposed in [225] solves the reverse SDE by combining numerical SDE solvers (“predictor”) and iterative Markov chain Monte Carlo (MCMC) approaches (“corrector”). At each time step, the predictor-corrector method first employs a numerical SDE solver to produce a coarse sample, followed by a “corrector” that corrects the sample’s marginal distribution with score-based MCMC. The resulting samples have the same time-marginals as solution trajectories of the reverse-time SDE, *i.e.*, they are equivalent in distribution at all time steps. Empirical results demonstrate that adding a corrector based on Langevin Monte Carlo is more efficient than using an additional predictor without correctors [225]. Karras et al. (2022) [113] further improve the Langevin dynamics corrector in [225] by proposing a Langevin-like “churn” step of adding and removing noise, achieving new state-of-the-art sample quality on datasets like CIFAR-10 [128] and ImageNet-64 [47].

3.1.2 ODE solvers. A large body of works on faster diffusion samplers are based on solving the probability flow ODE (Eq. (19)) introduced in Section 2.3. In contrast to SDE solvers, the trajectories of ODE solvers are deterministic and thus not affected by stochastic fluctuations. These deterministic ODE solvers typically converge much faster than their stochastic counterparts at the cost of slightly inferior sample quality.

Denoising Diffusion Implicit Models (DDIM) [217] is one of the earliest work on accelerating diffusion model sampling. The original motivation was to extend the original DDPM to non-Markovian case with the following Markov chain

$$q(\mathbf{x}_1, \dots, \mathbf{x}_T \mid \mathbf{x}_0) = \prod_{t=1}^T q(\mathbf{x}_t \mid \mathbf{x}_{t-1}, \mathbf{x}_0) \quad (25)$$

$$q_\sigma(\mathbf{x}_{t-1} \mid \mathbf{x}_t, \mathbf{x}_0) = \mathcal{N}(\mathbf{x}_{t-1} \mid \tilde{\mu}_t(\mathbf{x}_t, \mathbf{x}_0), \sigma_t^2 \mathbf{I}) \quad (26)$$

$$\tilde{\mu}_t(\mathbf{x}_t, \mathbf{x}_0) := \sqrt{\alpha_{t-1}} \mathbf{x}_0 + \sqrt{1 - \bar{\alpha}_{t-1} - \sigma_t^2} \cdot \frac{\mathbf{x}_t - \sqrt{\bar{\alpha}_t} \mathbf{x}_0}{\sqrt{1 - \bar{\alpha}_t}} \quad (27)$$

This formulation encapsulates DDPM and DDIM as special cases, where DDPM corresponds to setting $\sigma_t^2 = \frac{\hat{\beta}_{t-1}}{\hat{\beta}_t} \beta_t$ and DDIM corresponds to setting $\sigma_t^2 = 0$. DDIM learns a Markov chain to reverse this non-Markov perturbation process, which is fully deterministic when $\sigma_t^2 = 0$. It is observed in [113, 146, 203, 217] that the DDIM sampling process amounts to a special discretization scheme of the probability flow ODE. Inspired by an analysis of DDIM on a singleton dataset, generalized Denoising Diffusion Implicit Models (gDDIM) [278] proposes a modified parameterization of the score network that enables deterministic sampling for more general diffusion processes, such as the one in Critically-Damped Langevin Diffusion (CLD) [54]. PNDM [142] proposes a pseudo numerical method to generate sample along a specific manifold in \mathcal{R}^N . It uses numerical solver with nonlinear transfer part to solve differential equation on manifolds and then generates sample, which encapsulates DDIM as a special case.

Through extensive experimental investigations, Karras et al. (2022) [113] show that Heun’s 2nd order method [5] provides an excellent trade off between sample quality and sampling speed. The higher-order solver leads to smaller discretization error at the cost of one additional evaluation of the learned score function per time step. Heun’s method generates samples of comparable, if not better quality than Euler’s method with fewer sampling steps.

Diffusion Exponential Integrator Sampler [277] and DPM-solver [146] leverage the semi-linear structure of probability flow ODE to develop customized ODE solvers that are more efficient than general-purpose Runge-Kutta methods. Specifically, the linear part of probability flow ODE can be analytically computed, while the non-linear part can be solved with techniques similar to exponential integrators in the field of ODE solvers. These methods contain DDIM as a first-order approximation. However, they also allow for higher order integrators, which can produce high-quality samples in just 10 to 20 iterations—far fewer than the hundreds of iterations typically required by diffusion models without accelerated sampling.

3.2 Learning-Based Sampling

Learning-based sampling is another efficient approach for diffusion models. By using partial steps or training a sampler for the reverse process, this method achieves faster sampling speeds at the expense of slight degradation in sample quality. Unlike learning-free approaches that use handcrafted steps, learning-based sampling typically involves selecting steps by optimizing certain learning objectives.

3.2.1 Optimized Discretization. Given a pre-trained diffusion model, Watson et al. (2021) [243] put forth a strategy for finding the optimal discretization scheme by selecting the best K time steps to maximize the training objective for DDPMs. Key to this approach is the observation that the DDPM objective can be broken down into a sum of individual terms, making it well suited for dynamic programming. However, it is well known that the variational lower bound used for DDPM training does not correlate directly with sample quality [232]. A subsequent work, called Differentiable Diffusion Sampler Search [242], addresses this issue by directly optimizing a common metric for sample quality called the Kernel Inception Distance (KID) [15]. This optimization is feasible with the help of reparameterization [123, 195] and gradient rematerialization. Based on truncated Taylor methods, Dockhorn et al. (2022) [55] derive a second-order solver for accelerating synthesis by training a additional head on top of the first-order score network.

3.2.2 Truncated Diffusion. One can improve sampling speed by truncating the forward and reverse diffusion processes [156, 284]. The key idea is to halt the forward diffusion process early on, after just a few steps, and to begin the reverse denoising process with a non-Gaussian distribution. Samples from this distribution can be obtained efficiently by diffusing samples from pre-trained generative models, such as variational autoencoders [123, 195] or generative adversarial networks [73].

3.2.3 Knowledge Distillation. Approaches that use knowledge distillation [148, 203] can significantly improve the sampling speed of diffusion models. Specifically, in Progressive Distillation [203], the authors propose distilling the full sampling process into a faster sampler that requires only half as many steps. By parameterizing the new sampler as a deep neural network, authors are able to train the sampler to match the input and output of the DDIM sampling process. Repeating this procedure can further reduce sampling steps, although fewer steps can result in reduced sample quality. To address this issue, the authors suggest new parameterizations for diffusion models and new weighting schemes for the objective function.

4 DIFFUSION MODELS WITH IMPROVED LIKELIHOOD

As discussed in Section 2.1, the training objective for diffusion models is a (negative) variational lower bound (VLB) on the log-likelihood. This bound, however, may not be tight in many cases [121], leading to potentially suboptimal log-likelihoods from diffusion models. In this section, we survey recent works on likelihood maximization for diffusion

models. We focus on three types of methods: noise schedule optimization, reverse variance learning, and exact log-likelihood evaluation.

4.1 Noise Schedule Optimization

In the classical formulation of diffusion models, noise schedules in the forward process are handcrafted without trainable parameters. By optimizing the forward noise schedule jointly with other parameters of diffusion models, one can further maximize the VLB in order to achieve higher log-likelihood values [121, 166].

The work of iDDPM [166] demonstrates that a certain cosine noise schedule can improve log-likelihoods. Specifically, the cosine noise schedule in their work takes the form of

$$\bar{\alpha}_t = \frac{h(t)}{h(0)}, \quad h(t) = \cos\left(\frac{t/T + m}{1+m} \cdot \frac{\pi}{2}\right)^2 \quad (28)$$

where $\bar{\alpha}_t$ and β_t are defined in Eqs. (2) and (3), and m is a hyperparameter to control the noise scale at $t = 0$. They also propose a parameterization of the reverse variance with an interpolation between β_t and $1 - \bar{\alpha}_t$ in the log domain.

In Variational Diffusion Models (VDMs) [121], authors propose to improve the likelihood of continuous-time diffusion models by jointly training the noise schedule and other diffusion model parameters to maximize the VLB. They parameterize the noise schedule using a monotonic neural network $\gamma_\eta(t)$, and build the forward perturbation process according to $\sigma_t^2 = \text{sigmoid}(\gamma_\eta(t))$, $q(\mathbf{x}_t | \mathbf{x}_0) = \mathcal{N}(\bar{\alpha}_t \mathbf{x}_0, \sigma_t^2 \mathbf{I})$, and $\bar{\alpha}_t = \sqrt{(1 - \sigma_t^2)}$. Moreover, authors prove that the VLB for data point \mathbf{x} can be simplified to a form that only depends on the signal-to-noise ratio $R(t) := \frac{\bar{\alpha}_t^2}{\sigma_t^2}$. In particular, the L_{VLB} can be decomposed to

$$L_{VLB} = -\mathbb{E}_{\mathbf{x}_0} \text{KL}(q(\mathbf{x}_T | \mathbf{x}_0) || p(\mathbf{x}_T)) + \mathbb{E}_{\mathbf{x}_0, \mathbf{x}_1} \log p(\mathbf{x}_0 | \mathbf{x}_1) - L_D, \quad (29)$$

where the first and second terms can be optimized directly in analogy to training variational autoencoders. The third term can be further simplified to the following:

$$L_D = \frac{1}{2} \mathbb{E}_{\mathbf{x}_0, \epsilon \sim \mathcal{N}(0, \mathbf{I})} \int_{R_{\min}}^{R_{\max}} \|\mathbf{x}_0 - \tilde{\mathbf{x}}_\theta(\mathbf{x}_v, v)\|_2^2 dv, \quad (30)$$

where $R_{\max} = R(1)$, $R_{\min} = R(T)$, $\mathbf{x}_v = \bar{\alpha}_v \mathbf{x}_0 + \sigma_v \epsilon$ denotes a noisy data point obtained by diffusing \mathbf{x}_0 with the forward perturbation process until $t = R^{-1}(v)$, and $\tilde{\mathbf{x}}_\theta$ denotes the predicted noise-free data point by the diffusion model. As a result, noise schedules do not affect the VLB as long as they share the same values at R_{\min} and R_{\max} , and will only affect the variance of Monte Carlo estimators for VLB.

4.2 Reverse Variance Learning

The classical formulation of diffusion models assumes that Gaussian transition kernels in the reverse Markov chain have fixed variance parameters. Recall that we formulated the reverse kernel as $q_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t) = \mathcal{N}(\mu_\theta(\mathbf{x}_t, t), \Sigma_\theta(\mathbf{x}_t, t))$ in Eq. (5) but often fixed the reverse variance $\Sigma_\theta(\mathbf{x}_t, t)$ to $\beta_t \mathbf{I}$. Many methods propose to train the reverse variances as well to further maximize VLB and log-likelihood values.

In iDDPM [166], Nichol and Dhariwal propose to learn the reverse variances by parameterizing them with a form of linear interpolation and training them using a hybrid objective. This results in higher log-likelihoods and faster sampling without losing sample quality. In particular, they parameterize the reverse variance in Eq. (5) as:

$$\Sigma_\theta(\mathbf{x}_t, t) = \exp(\theta \cdot \log \beta_t + (1 - \theta) \cdot \log \tilde{\beta}_t), \quad (31)$$

where $\tilde{\beta}_t := \frac{1-\tilde{\alpha}_{t-1}}{1-\tilde{\alpha}_t} \cdot \beta_t$ and θ is jointly trained to maximize VLB. This simple parameterization avoids the instability of estimating more complicated forms of $\Sigma_\theta(\mathbf{x}_t, t)$ and is reported to improve likelihood values.

Analytic-DPM [8] shows a remarkable result that the optimal reverse variance can be obtained from a pre-trained score function, with the analytic form below:

$$\Sigma_\theta(\mathbf{x}_t, t) = \sigma_t^2 + \left(\sqrt{\frac{\tilde{\beta}_t}{\alpha_t}} - \sqrt{\tilde{\beta}_{t-1} - \sigma_t^2} \right)^2 \cdot \left(1 - \tilde{\beta}_t \mathbb{E}_{q_t(\mathbf{x}_t)} \frac{\|\nabla_{\mathbf{x}_t} \log q_t(\mathbf{x}_t)\|^2}{d} \right) \quad (32)$$

As a result, given a pre-trained score model, we can estimate its first- and second-order moments to obtain the optimal reverse variances. Plugging them into the VLB can lead to tighter VLBs and higher likelihood values.

4.3 Exact Likelihood Computation

In the Score SDE [225] formulation, samples are generated by solving the following reverse SDE, where $\nabla_{\mathbf{x}_t} \log p_\theta(\mathbf{x}_t, t)$ in Eq. (18) is replaced by the learned noise-conditional score model $\mathbf{s}_\theta(\mathbf{x}_t, t)$:

$$d\mathbf{x} = f(\mathbf{x}_t, t) - g(t)^2 \mathbf{s}_\theta(\mathbf{x}_t, t) dt + g(t) d\mathbf{w}. \quad (33)$$

Here we use p_θ^{sde} to denote the distribution of samples generated by solving the above SDE. One can also generate data by plugging the score model into the probability flow ODE in Eq. (19), which gives:

$$\begin{aligned} \frac{d\mathbf{x}_t}{dt} &= \underbrace{f(\mathbf{x}_t, t) - \frac{1}{2} g^2(t) \mathbf{s}_\theta(\mathbf{x}_t, t)}_{:= \tilde{f}_\theta(\mathbf{x}_t, t)} \end{aligned} \quad (34)$$

Similarly, we use p_θ^{oode} to denote the distribution of samples generated via solving this ODE. The theory of neural ODEs [30] and continuous normalizing flows [77] indicates that p_θ^{oode} can be computed accurately albeit with high computational cost. For p_θ^{sde} , several concurrent works [98, 145, 219] demonstrate that there exists an efficiently computable variational lower bound, and we can directly train our diffusion models to maximize p_θ^{sde} using modified diffusion losses.

Specifically, Song et al. (2021) [219] prove that with a special weighting function (likelihood weighting), the objective used for training score SDEs implicitly maximizes the expected value of p_θ^{sde} on data. It is shown that

$$\mathbf{D}_{KL}(q_0 \parallel p_\theta^{\text{sde}}) \leq \mathcal{L}(\theta; g(\cdot)^2) + \mathbf{D}_{KL}(q_T \parallel \pi), \quad (35)$$

where $\mathcal{L}(\theta; g(\cdot)^2)$ is the Score SDE objective in Eq. (20) with $\lambda(t) = g(t)^2$. Since $\mathbf{D}_{KL}(q_0 \parallel p_\theta^{\text{sde}}) = -\mathbb{E}_{q_0} \log(p_\theta^{\text{sde}}) + \text{const}$, and $\mathbf{D}_{KL}(q_T \parallel \pi)$ is a constant, training with $\mathcal{L}(\theta; g(\cdot)^2)$ amounts to minimizing $-\mathbb{E}_{q_0} \log(p_\theta^{\text{sde}})$, the expected negative log-likelihood on data. Moreover, Song et al. (2021) and Huang et al. (2021) [98, 219] provide the following bound for $p_\theta^{\text{sde}}(\mathbf{x})$:

$$-\log p_\theta^{\text{sde}}(\mathbf{x}) \leq \mathcal{L}'(\mathbf{x}), \quad (36)$$

where $\mathcal{L}'(\mathbf{x})$ is defined by

$$\mathcal{L}'(\mathbf{x}) := \int_0^T \mathbb{E} \left[\frac{1}{2} \|g(t) \mathbf{s}_\theta(\mathbf{x}_t, t)\|^2 + \nabla \cdot (g(t)^2 \mathbf{s}_\theta(\mathbf{x}_t, t) - f(\mathbf{x}_t, t)) \mid \mathbf{x}_0 = \mathbf{x} \right] dt - \mathbb{E}_{\mathbf{x}_T} [\log p_\theta^{\text{sde}}(\mathbf{x}_T) \mid \mathbf{x}_0 = \mathbf{x}] \quad (37)$$

The first part of Eq. (37) is reminiscent of implicit score matching [101] and the whole bound can be efficiently estimated with Monte Carlo methods.

Since the probability flow ODE is a special case of neural ODEs or continuous normalizing flows, we can use well-established approaches in those fields to compute $\log p_\theta^{\text{oode}}$ accurately. Specifically, we have

$$\log p_\theta^{\text{oode}}(\mathbf{x}_0) = \log p_T(\mathbf{x}_T) + \int_{t=0}^T \nabla \cdot \tilde{f}_\theta(\mathbf{x}_t, t) dt. \quad (38)$$

One can compute the one-dimensional integral above with numerical ODE solvers and the Skilling-Hutchinson trace estimator [100, 214]. Unfortunately, this formula cannot be directly optimized to maximize p_θ^{oode} on data, as it requires calling expensive ODE solvers for each data point \mathbf{x}_0 . To reduce the cost of directly maximizing p_θ^{oode} with the above formula, Song et al. (2021) [219] propose to maximize the variational lower bound of p_θ^{sde} as a proxy for maximizing p_θ^{oode} , giving rise to a family of diffusion models called *ScoreFlows*.

Lu et al. (2022) [145] further improve ScoreFlows by proposing to minimize not just the vanilla score matching loss function, but also its higher order generalizations. They prove that $\log p_\theta^{\text{oode}}$ can be bounded with the first, second, and third-order score matching errors. Building upon this theoretical result, authors further propose efficient training algorithms for minimizing high order score matching losses and reported improved p_θ^{oode} on data.

5 DIFFUSION MODELS FOR DATA WITH SPECIAL STRUCTURES

While diffusion models have achieved great success for data domains like images and audio, they do not necessarily translate seamlessly to other modalities. Many important data domains have special structures that must be taken into account for diffusion models to function effectively. Difficulties may arise, for example, when models rely on score functions that are only defined on continuous data domains, or when data reside on low dimensional manifolds. To cope with these challenges, diffusion models have to be adapted in various ways.

5.1 Discrete Data

Most diffusion models are geared towards continuous data domains, because Gaussian noise perturbation as used in DDPMs is not a natural fit for discrete data, and the score functions required by SGMs and Score SDEs are only defined on continuous data domains. To overcome this difficulty, several works [6, 83, 96, 255] build on Sohl-Dickstein et al. (2015) [215] to generate discrete data of high dimensions. Specifically, VQ-Diffusion [83] replaces Gaussian noise with a random walk on the discrete data space, or a random masking operation. The resulting transition kernel for the forward process takes the form of

$$q(\mathbf{x}_t \mid \mathbf{x}_{t-1}) = \mathbf{v}^\top(\mathbf{x}_t) \mathbf{Q}_t \mathbf{v}(\mathbf{x}_{t-1}) \quad (39)$$

where $\mathbf{v}(\mathbf{x})$ is a one-hot column vector, and \mathbf{Q}_t is the transition kernel of a lazy random walk. D3PM [6] accommodates discrete data in diffusion models by constructing the forward noising process with absorbing state kernels or discretized Gaussian kernels. Campbell et al. (2022) [21] present the first continuous-time framework for discrete diffusion models. Leveraging Continuous Time Markov Chains, they are able to derive efficient samplers that outperform discrete counterparts, while providing a theoretical analysis on the error between the sample distribution and the true data distribution.

5.2 Data with Invariant Structures

Data in many important domains have invariant structures. For example, graphs are permutation invariant, and point clouds are both translation and rotation invariant. In diffusion models, these invariances are often ignored, which can

lead to suboptimal performance. To address this issue, several works [45, 171] propose to endow diffusion models with the ability to account for invariance in data.

Niu et al. (2020) [171] first tackle the problem of permutation invariant graph generation with diffusion models. They achieve this by using a permutation equivariant graph neural network [74, 208, 251], called EDP-GNN, to parameterize the noise-conditional score model. GDSS [108] further develops this idea by proposing a continuous-time graph diffusion process. This process models both the joint distribution of nodes and edges through a system of stochastic differential equations (SDEs), where message-passing operations are used to guarantee permutation invariance.

Similarly, Shi et al. (2021) [210] and Xu et al. (2022) [259] enable diffusion models to generate molecular conformations that are invariant to both translation and rotation. For example, Xu et al. (2022) [259] shows that Markov chains starting with an invariant prior and evolving with equivariant Markov kernels can induce an invariant marginal distribution, which can be used to enforce appropriate data invariance in molecular conformation generation. Formally, let \mathcal{T} be a rotation or translation operation. Given that $p(\mathbf{x}_T) = p(\mathcal{T}(\mathbf{x}_T))$, $p_\theta(\mathbf{x}_{t-1} | \mathbf{x}_t) = p_\theta(\mathcal{T}(\mathbf{x}_{t-1}) | \mathcal{T}(\mathbf{x}_t))$, Xu et al. (2022) [259] prove that the distribution of samples is guaranteed to be invariant to \mathcal{T} , that is, $p_0(\mathbf{x}) = p_0(\mathcal{T}(\mathbf{x}))$. As a result, one can build a diffusion model that generates rotation and translation invariant molecular conformations as long as the prior and transition kernels enjoy the same invariance.

5.3 Data with Manifold Structures

Data with manifold structures are ubiquitous in machine learning. As the manifold hypothesis [63] posits, natural data often reside on manifolds with lower intrinsic dimensionality. In addition, many data domains have well-known manifold structures. For instance, climate and earth data naturally lie on the sphere because that is the shape of our planet. Many works have focused on developing diffusion models for data on manifolds. We categorize them based on whether the manifolds are known or learned, and introduce some representative works below.

5.3.1 Known Manifolds. Recent studies have extended the Score SDE formulation to various known manifolds. This adaptation parallels the generalization of neural ODEs [30] and continuous normalizing flows [77] to Riemannian manifolds [144, 158]. To train these models, researchers have also adapted score matching and score functions to Riemannian manifolds.

The Riemannian Score-Based Generative Model (RSGM) [45] accommodates a wide range of manifolds, including spheres and toruses, provided they satisfy mild conditions. The RSGM demonstrates that it is possible to extend diffusion models to compact Riemannian manifolds. The model also provides a formula for reversing diffusion on a manifold. Taking an intrinsic view, the RSGM approximates the sampling process on Riemannian manifolds using a Geodesic Random Walk. It is trained with a generalized denoising score matching objective.

In contrast, the Riemannian Diffusion Model (RDM) [97] employs a variational framework to generalize the continuous-time diffusion model to Riemannian manifolds. The RDM uses a variational lower bound (VLB) of the log-likelihood as its loss function. The authors of the RDM model have shown that maximizing this VLB is equivalent to minimizing a Riemannian score-matching loss. Unlike the RSGM, the RDM takes an extrinsic view, assuming that the relevant Riemannian manifold is embedded in a higher dimensional Euclidean space.

5.3.2 Learned Manifolds. According to the manifold hypothesis [63], most natural data lies on manifolds with significantly reduced intrinsic dimensionality. Consequently, identifying these manifolds and training diffusion models directly on them can be advantageous due to the lower data dimensionality. Many recent works have built on this idea, starting by using an autoencoder to condense the data into a lower dimensional manifold, followed by training

diffusion models in this latent space. In these cases, the manifold is implicitly defined by the autoencoder and learned through the reconstruction loss. In order to be successful, it is crucial to design a loss function that allows for the joint training of the autoencoder and the diffusion models.

The Latent Score-Based Generative Model (LSGM) [234] seeks to address the problem of joint training by pairing a Score SDE diffusion model with a variational autoencoder (VAE) [123, 195]. In this configuration, the diffusion model is responsible for learning the prior distribution. The authors of the LSGM propose a joint training objective that merges the VAE’s evidence lower bound with the diffusion model’s score matching objective. This results in a new lower bound for the data log-likelihood. By situating the diffusion model within the latent space, the LSGM achieves faster sample generation than conventional diffusion models. Additionally, the LSGM can manage discrete data by converting it into continuous latent codes.

Rather than jointly training the autoencoder and diffusion model, the Latent Diffusion Model (LDM) [198] addresses each component separately. First, an autoencoder is trained to produce a low-dimensional latent space. Then, the diffusion model is trained to generate latent codes. DALLE-2 [186] employs a similar strategy by training a diffusion model on the CLIP image embedding space, followed by training a separate decoder to create images based on the CLIP image embeddings.

6 CONNECTIONS WITH OTHER GENERATIVE MODELS

In this section, we first introduce five other important classes of generative models and analyze their advantages and limitations. Then we introduce how diffusion models are connected with them, and illustrate how these generative models are promoted by incorporating diffusion models. The algorithms that integrate diffusion models with other generative models are summarized in Table 2, and we also provide a schematic illustration in Fig. 3.

Table 2. Diffusion models are incorporated into different generative models.

Model	Article	Year
VAE	Luo et al. [149]	2022
	Hunag et al. [98]	2021
	Vadhat et al. [234]	2021
GAN	Wang et al. [241]	2022
	Xiao et al. [253]	2021
Normalizing Flow	Zhang et al.[276]	2021
	Gong et al. [72]	2021
Autoregressive Model	Hoogeboom et al.[95]	2021
	Rasul et al. [190]	2021
Energy-based Model	Gao et al. [69]	2021
	Yu et al. [272]	2022

6.1 Variational Autoencoders and Connections with Diffusion Models

Variational Autoencoders [56, 124, 195] aim to learn both an encoder and a decoder to map input data to values in a continuous latent space. In these models, the embedding can be interpreted as a latent variable in a probabilistic generative model, and a probabilistic decoder can be formulated by a parameterized likelihood function. In addition, the data \mathbf{x} is assumed to be generated by some unobserved latent variable \mathbf{z} using conditional distribution $p_\theta(\mathbf{x} | \mathbf{z})$.

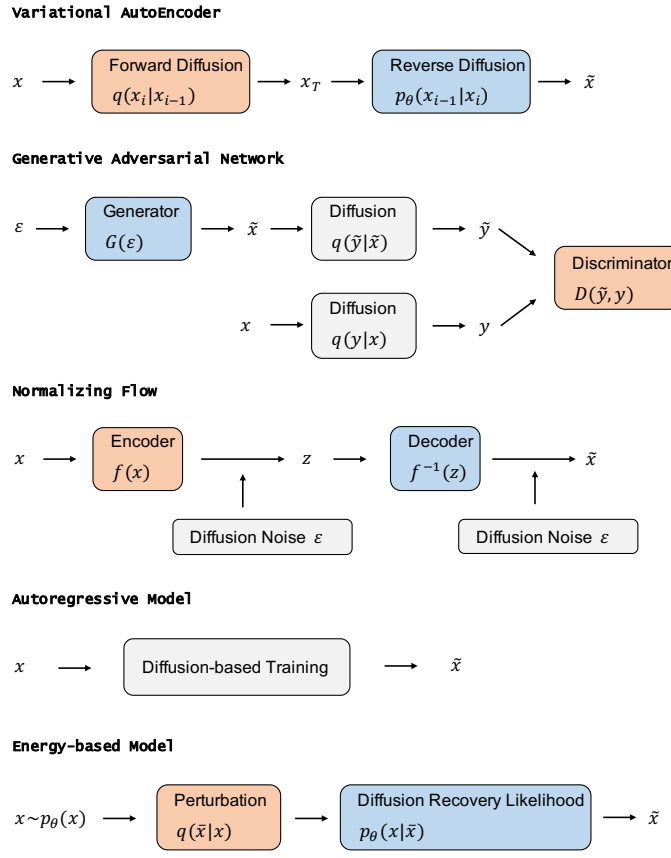


Fig. 3. Illustrations about works incorporating diffusion models with other generative models.

and $q_\phi(z | x)$ is used to approximately inference z . To guarantee an effective inference, a variational Bayes approach is used to maximize the evidence lower bound:

$$\mathcal{L}(\phi, \theta; x) = \mathbb{E}_{q(z|x)} [\log p_\theta(x, z) - \log q_\phi(z | x)] \quad (40)$$

with $\mathcal{L}(\phi, \theta; x) \leq \log p_\theta(x)$. Provided that the parameterized likelihood function $p_\theta(x | z)$ and the parameterized posterior approximation $q_\phi(z | x)$ can be computed in a point-wise way and are differentiable with their parameters, the ELBO can be maximized with gradient descent. This formulation allows flexible choices of encoder and decoder models. Typically, these models are represented by exponential family distributions whose parameters are generated by multi-layer neural networks.

The DDPM can be conceptualized as a hierarchical Markovian VAE with a fixed encoder. Specifically, DDPM's forward process functions as the encoder, and this process is structured as a linear Gaussian model (as described by Eq. (2)). The DDPM's reverse process, on the other hand, corresponds to the decoder, which is shared across multiple decoding steps. The latent variables within the decoder are all the same size as the sample data.

In a continuous-time setting, Song et al. (2021) [225], Huang et al. (2021) [98], and Kingma et al. (2021) [121] demonstrate that the score matching objective may be approximated by the Evidence Lower Bound (ELBO) of a deep

hierarchical VAE. Consequently, optimizing a diffusion model can be seen as training an infinitely deep hierarchical VAE—a finding that supports the common belief that Score SDE diffusion models can be interpreted as the continuous limit of hierarchical VAEs.

The Latent Score-Based Generative Model (LSGM) [234] furthers this line of research by illustrating that the ELBO can be considered a specialized score matching objective in the context of latent space diffusion. Though the cross-entropy term in the ELBO is intractable, it can be transformed into a tractable score matching objective by viewing the score-based generative model as an infinitely deep VAE.

6.2 Generative Adversarial Networks and Connections with Diffusion Models

Generative Adversarial Networks (GANs) [40, 73, 84] mainly consist of two models: a generator G and a discriminator D . These two models are typically constructed by neural networks but could be implemented in any form of a differentiable system that maps input data from one space to another. The optimization of GANs can be viewed as a mini-max optimization problem with value function $V(G, D)$:

$$\min_G \max_D \mathbb{E}_{\mathbf{x} \sim p_{\text{data}}(\mathbf{x})} [\log D(\mathbf{x})] + \mathbb{E}_{\mathbf{z} \sim p_{\mathbf{z}}(\mathbf{z})} [\log(1 - D(G(\mathbf{z})))]. \quad (41)$$

The generator G aims to generate new examples and implicitly model the data distribution. The discriminator D is usually a binary classifier that is used to identify generated examples from true examples with maximally possible accuracy. The optimization process ends at a saddle point that produces a minimum about the generator and a maximum about the discriminator. Namely, the goal of GAN optimization is to achieve Nash equilibrium [193]. At that point, the generator can be considered that it has captured the accurate distribution of real examples.

One of the issues of GAN is the instability in the training process, which is mainly caused by the non-overlapping between the distribution of input data and that of the generated data. One solution is to inject noise into the discriminator input for widening the support of both the generator and discriminator distributions. Taking advantage of the flexible diffusion model, Wang et al. (2022) [241] inject noise to the discriminator with an adaptive noise schedule determined by a diffusion model. On the other hand, GAN can facilitate sampling speed of diffusion models. Xiao et al. (2021) [253] show that slow sampling is caused by the Gaussian assumption in the denoising step, which is justified only for small step sizes. As such, each denoising step is modeled by a conditional GAN, allowing larger step size.

6.3 Normalizing Flows and Connections with Diffusion Models

Normalizing flows [51, 194] are generative models that generate tractable distributions to model high-dimensional data [53, 122]. Normalizing flows can transform simple probability distribution into an extremely complex probability distribution, which can be used in generative models, reinforcement learning, variational inference, and other fields. Existing normalizing flows are constructed based on the change of variable formula [51, 194]. The trajectory in normalizing flows is formulated by a differential equation. In the discrete-time setting, the mapping from data \mathbf{x} to latent \mathbf{z} in normalizing flows is a composition of a sequence of bijections, taking the form of $F = F_N \circ F_{N-1} \circ \dots \circ F_1$. The trajectory $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_N\}$ in normalizing flows satisfies :

$$\mathbf{x}_i = F_i(\mathbf{x}_{i-1}, \theta), \quad \mathbf{x}_{i-1} = F_i^{-1}(\mathbf{x}_i, \theta) \quad (42)$$

for all $i \leq N$.

Similar to the continuous setting, normalizing flows allow for the retrieval of the exact log-likelihood through a change of variable formula. However, the bijection requirement limits the modeling of complex data in both practical

and theoretical contexts [39, 246]. Several works attempt to relax this bijection requirement [53, 246]. For example, DiffFlow [276] introduces a generative modeling algorithm that combines the benefits of both flow-based and diffusion models. As a result, DiffFlow produces sharper boundaries than normalizing flow and learns more general distributions with fewer discretization steps compared to diffusion probabilistic models.

6.4 Autoregressive Models and Connections with Diffusion Models

Autoregressive Models (ARMs) work by decomposing the joint distribution of data into a product of conditional distributions using the probability chain rule:

$$\log p(\mathbf{x}_{1:T}) = \sum_{t=1}^T \log p(x_t | \mathbf{x}_{<t}) \quad (43)$$

where $\mathbf{x}_{<t}$ is a shorthand for x_1, x_2, \dots, x_{t-1} [11, 130]. Recent advances in deep learning have facilitated significant progress for various data modalities [25, 162, 207], such as images [34, 237], audio [112, 236], and text [12, 18, 80, 160, 163]. Autoregressive models (ARMs) offer generative capabilities through the use of a single neural network. Sampling from these models requires the same number of network calls as the data's dimensionality. While ARMs are effective density estimators, sampling is a continuous, time-consuming process—particularly for high-dimensional data.

The Autoregressive Diffusion Model (ARDM) [95], on the other hand, is capable of generating arbitrary-order data, including order-agnostic autoregressive models and discrete diffusion models as special cases [6, 96, 216]. Instead of using causal masking on representations like ARMs, the ARDM is trained with an effective objective that mirrors that of diffusion probabilistic models. At the testing stage, the ARDM is able to generate data in parallel—enabling its application to a range of arbitrary-generation tasks.

6.5 Energy-based Models and Connections with Diffusion Models

Energy-based Models (EBMs) [26, 48, 58, 64, 67, 68, 75, 78, 79, 120, 129, 132, 165, 170, 182, 196, 254, 281] can be viewed as one generative version of discriminators [79, 104, 131, 134], while can be learned from unlabeled input data. Let $\mathbf{x} \sim p_{\text{data}}(\mathbf{x})$ denote a training example, and $p_\theta(\mathbf{x})$ denote a probability density function that aims to approximates $p_{\text{data}}(\mathbf{x})$. An energy-based model is defined as:

$$p_\theta(\mathbf{x}) = \frac{1}{Z_\theta} \exp(f_\theta(\mathbf{x})), \quad (44)$$

where $Z_\theta = \int \exp(f_\theta(\mathbf{x})) d\mathbf{x}$ is the partition function, which is analytically intractable for high-dimensional \mathbf{x} . For images, $f_\theta(\mathbf{x})$ is parameterized by a convolutional neural network with a scalar output. Salimans et al. (2021) [204] compare both constrained score models and energy-based models for modeling the score of the data distribution, finding that constrained score models, i.e., energy based models, can perform just as well as unconstrained models when using a comparable model structure.

Although EBMs have a number of desirable properties, two challenges remain for modeling high-dimensional data. First, learning EBMs by maximizing the likelihood requires MCMC method to generate samples from the model, which can be very computationally expensive. Second, as demonstrated in [169], the energy potentials learned with non-convergent MCMC are not stable, in the sense that samples from long-run Markov chains can be significantly different from the observed samples, and thus it is difficult to evaluate the learned energy potentials. In a recent study, Gao et al. (2021) [69] present a diffusion recovery likelihood method to tractably learn samples from a sequence of EBMs in the reverse process of the diffusion model. Each EBM is trained with recovery likelihood, which aims to

maximize the conditional probability of the data at a certain noise level, given their noisy versions at a higher noise level. EBMs maximize the recovery likelihood because it is more tractable than marginal likelihood, as sampling from the conditional distributions is much easier than sampling from the marginal distributions. This model can generate high-quality samples, and long-run MCMC samples from the conditional distributions still resemble realistic images.

7 APPLICATIONS OF DIFFUSION MODELS

Diffusion models have recently been employed to address a variety of challenging real-world tasks due to their flexibility and strength. We have grouped these applications into six different categories based on the task: computer vision, natural language processing, temporal data modeling, multi-modal learning, robust learning, and interdisciplinary applications. For each category, we provide a brief introduction to the task, followed by a detailed explanation of how diffusion models have been applied to improve performance. Table 3 summarizes the various applications that have made use of diffusion models.

Table 3. Summary of all the applications utilizing the diffusion models.

Primary	Secondary	Article
Computer Vision	Super Resolution, Inpainting, and Translation	[137], [202], [198], [147], [200], [181], [91], [10], [174], [38], [224], [36], [161]
	Semantic Segmentation	[9], [17], [76]
	Video Generation	[89], [93], [268], [275]
	Point Cloud Completion and Generation	[286], [150], [155], [143], [274]
Natural Language Processing	Anomaly Detection	[252], [70]
	Natural Language Processing	[6], [141], [32], [88]
Temporal Data Modeling	Time Series Imputation	[230], [1], [177]
	Time Series Forecasting	[191], [1]
	Waveform Signal Processing	[29], [126]
Multi-Modal Learning	Text-to-Image Generation	[7], [186], [201], [167], [83], [62], [199], [116], [235]
	Text-to-Audio Generation	[180], [261], [249], [136], [228], [99], [119]
Robust Learning	Robust Learning	[168], [270], [16], [240], [248], [227]
	Molecular Graph Modeling	[107], [94], [3], [259], [233], [247], [210], [152]
Interdisciplinary Applications	Material Design	[256], [153]
	Medical Image Reconstruction	[224], [36], [37], [38], [178], [257]

7.1 Computer Vision

7.1.1 Super Resolution, Inpainting, and Translation. Generative models have been used to tackle a variety of image restoration tasks including super-resolution, inpainting, and translation [10, 47, 61, 103, 137, 174, 187, 282]. Image super-resolution aims to restore high-resolution images from low-resolution inputs, while image inpainting revolves around reconstructing missing or damaged regions in an image.

Several methods make use of diffusion models for these tasks. For example, Super-Resolution via Repeated Refinement (SR3) [202] uses DDPM to enable conditional image generation. SR3 conducts super-resolution through a stochastic, iterative denoising process. The Cascaded Diffusion Model (CDM) [91] consists of multiple diffusion models in sequence, each generating images of increasing resolution. Both the SR3 and CDM directly apply the diffusion process to input images, which leads to larger evaluation steps.

In order to allow for the training of diffusion models with limited computational resources, some methods [198, 234] have shifted the diffusion process to the latent space using pre-trained autoencoders. The Latent Diffusion Model (LDM) [198] streamlines the training and sampling processes for denoising diffusion models without sacrificing quality.

For inpainting tasks, RePaint [147] features an enhanced denoising strategy that uses resampling iterations to better condition the image (see Figure Fig. 5). Meanwhile, Palette [200] employs conditional diffusion models to create a unified framework for four image generation tasks: colorization, inpainting, uncropping, and JPEG restoration.

Image translation focuses on synthesizing images with specific desired styles [103]. SDEdit [161] uses a Stochastic Differential Equation (SDE) prior to improve fidelity. Specifically, it begins by adding noise to the input image, then denoises the image through the SDE.



Fig. 4. **Image super resolution results produced by LDM [198].**

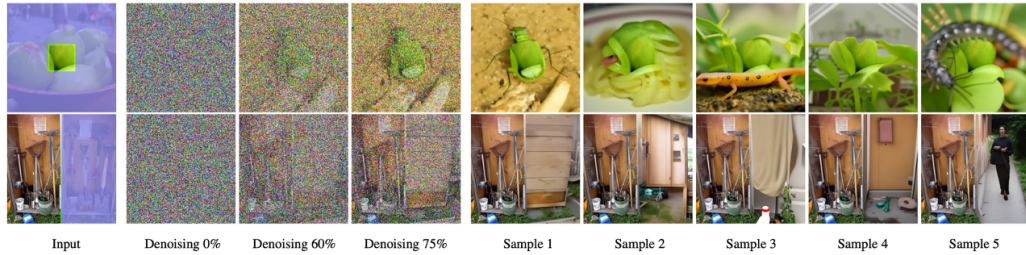


Fig. 5. **Image inpainting results produced by RePaint [147].**

7.1.2 Semantic Segmentation. Semantic segmentation aims to label each image pixel according to established object categories. Generative pre-training can enhance the label utilization of semantic segmentation models, and recent work has shown that representations learned through DDPM contain high-level semantic information that is useful for segmentation tasks [9, 76]. The few-shot method that leverages these learned representations has outperformed alternatives such as VDVAE [33] and ALAE [179]. Similarly, Decoder Denoising Pretraining (DDeP) [17] integrates diffusion models with denoising autoencoders [239] and delivers promising results on label-efficient semantic segmentation.

7.1.3 Video Generation. Generating high-quality videos remains a challenge in the deep learning era due to the complexity and spatio-temporal continuity of video frames [265, 273]. Recent research has turned to diffusion models to improve the quality of generated videos [93]. For example, the Flexible Diffusion Model (FDM) [89] uses a generative model to allow for the sampling of any arbitrary subset of video frames, given any other subset. The FDM also includes a

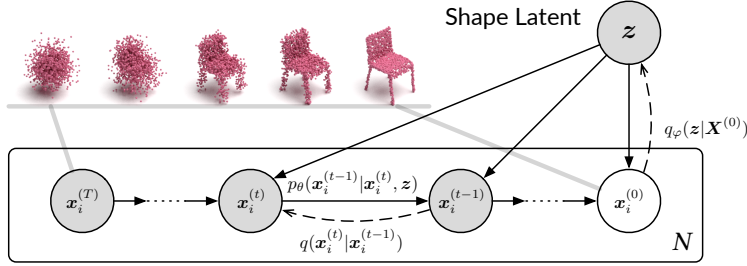


Fig. 6. **The directed graphical model of the diffusion process for point clouds [150].**

specialized architecture designed for this purpose. Additionally, the Residual Video Diffusion (RVD) model [268] utilizes an autoregressive, end-to-end optimized video diffusion model. It generates future frames by amending a deterministic next-frame prediction, using a stochastic residual produced through an inverse diffusion process.

7.1.4 Point Cloud Completion and Generation. Point clouds are a critical form of 3D representation for capturing real-world objects. However, scans often generate incomplete point clouds due to partial observation or self-occlusion. Recent studies have applied diffusion models to address this challenge, using them to infer missing parts in order to reconstruct complete shapes. This work has implications for many downstream tasks such as 3D reconstruction, augmented reality, and scene understanding [151, 155, 274].

Luo et al. 2021 [150] has taken the approach of treating point clouds as particles in a thermodynamic system, using a heat bath to facilitate diffusion from the original distribution to a noise distribution. Meanwhile, the Point-Voxel Diffusion (PVD) model [286] joins denoising diffusion models with the pointvoxel representation of 3D shapes. The Point Diffusion-Refinement (PDR) model [155] uses a conditional DDPM to generate a coarse completion from partial observations; it also establishes a point-wise mapping between the generated point cloud and the ground truth.

7.1.5 Anomaly Detection. Anomaly detection is a critical and challenging problem in machine learning [209, 283] and computer vision [262]. Generative models have been shown to own a powerful mechanism for anomaly detection [70, 87, 252], modeling normal or healthy reference data. AnoDDPM [252] utilizes DDPM to corrupt the input image and reconstruct a healthy approximation of the image. These approaches may perform better than alternatives based on adversarial training as they can better model smaller datasets with effective sampling and stable training schemes. DDPM-CD [70] incorporates large numbers of unsupervised remote sensing images into the training process through DDPM. Changes of remote sensed images is detected by utilizing a pre-trained DDPM and applying the multi-scale representations from the diffusion model decoder.

7.2 Natural Language Processing

Natural language processing aims to understand, model, and manage human languages from different sources such as text or audio. Text generation has become one of the most critical and challenging tasks in natural language processing [102, 138, 139]. It aims to compose plausible and readable text in the human language given input data (e.g., a sequence and keywords) or random noise. Numerous approaches based on diffusion models have been developed for text generation. Discrete Denoising Diffusion Probabilistic Models (D3PM) [6] introduces diffusion-like generative models for character-level text generation [28]. It generalizes the multinomial diffusion model [96] through going beyond corruption processes with uniform transition probabilities. Large autoregressive language models (LMs) is

able to generate high-quality text [18, 35, 185, 279]. To reliably deploy these LMs in real-world applications, the text generation process is usually expected to be controllable. It means we need to generate text that can satisfy desired requirements (e.g., topic, syntactic structure). Controlling the behavior of language models without re-training is a major and important problem in text generation [43, 117]. Although recent methods have achieved significant successes on controlling simple sentence attributes (e.g., sentiment) [127, 263], there is little progress on complex, fine-grained controls (e.g., syntactic structure). In order to tackle more complex controls, Diffusion-LM [141] proposes a new language model based on continuous diffusion. Diffusion-LM starts with a sequence of Gaussian noise vectors and incrementally denoises them into vectors corresponding to words. The gradual denoising steps help produce hierarchical continuous latent representations. This hierarchical and continuous latent variable can make it possible for simple, gradient-based methods to accomplish complex control. Analog Bits [32] generates the analog bits to represent the discrete variables and further improves the sample quality with self-conditioning and asymmetric time intervals. DiffuSeq [88] proposes a new conditional diffusion model to accomplish more challenging text generation tasks.

7.3 Temporal Data Modeling

7.3.1 Time Series Imputation. Time series data are widely used with many important real-world applications [60, 173, 265, 280]. Nevertheless, time series usually contain missing values for multiple reasons, caused by mechanical or artificial errors [213, 229, 269]. Recent years, imputation methods have been greatly for both deterministic imputation [23, 27, 154] and probabilistic imputation [65], including diffusion-based approaches. Conditional Score-based Diffusion models for Imputation (CSDI) [230] presents a novel time series imputation method that leverages score-based diffusion models. Specifically, for the purpose of exploiting correlations within temporal data, it adopts the form of self-supervised training to optimize diffusion models. Its application in some real-world datasets reveals its superiority over previous methods. Controlled Stochastic Differential Equation (CSDE) [177] proposes a novel probabilistic framework for modeling stochastic dynamics with a neural-controlled stochastic differential equation. Structured State Space Diffusion (SSSD) [1] integrates conditional diffusion models and structured state-space models [82] to particularly capture long-term dependencies in time series. It performs well in both time series imputation and forecasting tasks.

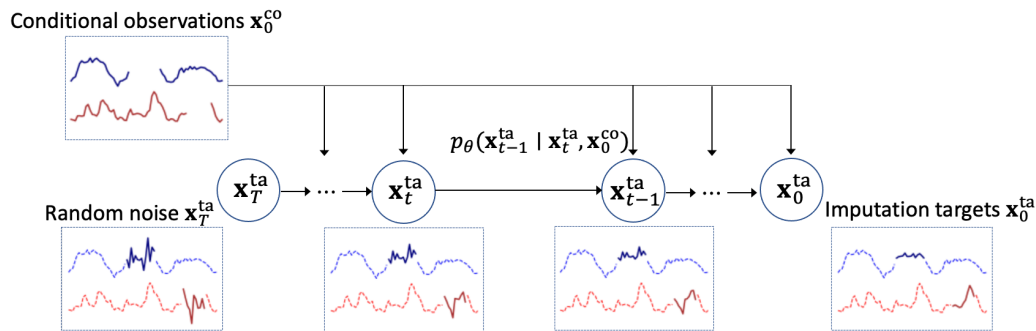


Fig. 7. The procedure of time series imputation with CSDI [230].

7.3.2 Time Series Forecasting. Time series forecasting is the task of forecasting or predicting the future value over a period of time. Neural methods have recently become widely-used for solving the prediction problem with univariate point forecasting methods [172] or univariate probabilistic methods [205]. In the multivariate setting, we also have

point forecasting methods [140] as well as probabilistic methods, which explicitly model the data distribution using Gaussian copulas [206], GANs [271], or normalizing flows [192]. TimeGrad [191] presents an autoregressive model for forecasting multivariate probabilistic time series, which samples from the data distribution at each time step through estimating its gradient. It utilizes diffusion probabilistic models, which are closely connected with score matching and energy-based methods. Specifically, it learns gradients by optimizing a variational bound on the data likelihood and transforms white noise into a sample of the distribution of interest through a Markov chain using Langevin sampling [220] during inference time.

7.3.3 Waveform Signal Processing. In electronics, acoustics, and some related fields, the waveform of a signal is denoted by the shape of its graph as a function of time, independent of its time and magnitude scales. WaveGrad [29] introduces a conditional model for waveform generation that estimates gradients of the data density. It receives a Gaussian white noise signal as input and iteratively refines the signal with a gradient-based sampler. WaveGrad naturally trades inference speed for sample quality by adjusting the number of refinement steps, and make a connection between non-autoregressive and autoregressive models with respect to audio quality. DiffWave [126] presents a versatile and effective diffusion probabilistic model for conditional or unconditional waveform generation. The model is non-autoregressive and is efficiently trained by optimizing a variant of variational bound on the data likelihood. Moreover, it produces high-fidelity audio in different waveform generation tasks, such as class-conditional generation and unconditional generation.

7.4 Multi-Modal Learning

7.4.1 Text-to-Image Generation. Vision-language models have attracted a lot of attention recently due to the number of potential applications [184]. Text-to-Image generation is the task of generating a corresponding image from a descriptive text [57, 116, 235]. An example is shown in Fig. 8. Blended diffusion [7] utilizes both pre-trained DDPM [49] and CLIP [184] models, and it proposes a solution for region-based image editing for general purposes, which uses natural language guidance and is applicable to real and diverse images. On the other hand, unCLIP (DALLE-2) [186] proposes a two-stage approach, a prior model that can generate a CLIP-based image embedding conditioned on a text caption, and a diffusion-based decoder that can generate an image conditioned on the image embedding. Recently, Imagen [201] proposes a text-to-image diffusion model and a comprehensive benchmark for performance evaluation. It shows that Imagen performs well against the state-of-the-art approaches including VQ-GAN+CLIP [41], Latent Diffusion Models [146], and DALL-E 2 [186]. Inspired by the ability of guided diffusion models [49, 92] to generate photorealistic samples and the ability of text-to-image models to handle free-form prompts, GLIDE [167] applies guided diffusion to the application of text-conditioned image synthesis. VQ-Diffusion [83] proposes a vector-quantized diffusion model for text-to-image generation, and it eliminates the unidirectional bias and avoids accumulative prediction errors.

7.4.2 Text-to-Audio Generation. Text-to-audio generation is the task to transform normal language texts to voice outputs [136, 249]. Grad-TTS [180] presents a novel text-to-speech model with a score-based decoder and diffusion models. It gradually transforms noise predicted by the encoder and is further aligned with text input by the method of Monotonic Alignment Search [183]. Grad-TTS2 [119] improves Grad-TTS in an adaptive way. Diffsound [261] presents a non-autoregressive decoder based on the discrete diffusion model [6, 215], which predicts all the mel-spectrogram tokens in every single step, and then refines the predicted tokens in the following steps. EdiTTS [228] leverages the score-based text-to-speech model to refine a mel-spectrogram prior that is coarsely modified. Instead of estimating the gradient of data density, ProDiff [99] parameterizes the denoising diffusion model by directly predicting the clean data.

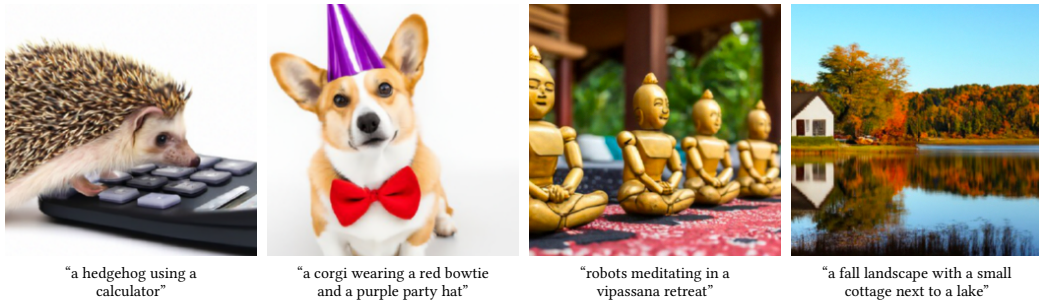


Fig. 8. Text-to-image result generated by GLIDE [167].

7.5 Robust Learning

Robust learning is a class of defense methods that help learning networks that are robust to adversarial perturbations or noises [16, 168, 179, 240, 248, 270]. While adversarial training [157] is viewed as a standard defense method against adversarial attacks for image classifiers, adversarial purification has shown significant performances as an alternative defense method [270], which purifies attacked images into clean images with a standalone purification model. Given an adversarial example, DiffPure [168] diffuses it with a small amount of noise following a forward diffusion process and then restores the clean image with a reverse generative process. Adaptive Denoising Purification (ADP) [270] demonstrates that an EBM trained with denoising score matching [238] can effectively purify attacked images within just a few steps. It further proposes an effective randomized purification scheme, injecting random noises into images before purification. Projected Gradient Descent (PGD) [16] presents a novel stochastic diffusion-based pre-processing robustification, which aims to be a model-agnostic adversarial defense and yield a high-quality denoised outcome. In addition, some works propose to apply a guided diffusion process for advanced adversarial purification [240, 248].

7.6 Interdisciplinary Applications

7.6.1 Molecular Graph Modeling. Graph Neural Networks [85, 251, 266, 285] and corresponding representation learning [86] techniques have achieved great success [14, 231, 250, 258, 264, 288] in many areas, including modeling molecule graph in various tasks ranging from property prediction [59, 71] to molecule generation [105, 111, 152, 211], where a molecule is naturally represented by a node-edge graph. Despite their effectiveness in different applications, more intrinsic and informative properties begin to be combined with diffusion models for enhancing molecular graph modeling. Torsional diffusion [107] presents a new diffusion framework that makes operations on the space of torsion angles with a diffusion process on the hyperspace and an extrinsic-to-intrinsic scoring model. GeoDiff [259] demonstrates that Markov chains evolving with equivariant Markov kernels can produce an invariant distribution, and further design blocks for the Markov kernels to preserve the desirable equivariance property. There are also other works incorporate the equivariance property into 3D molecule generation [94] and protein generation [3, 13]. Motivated by the classical force field methods for simulating molecular dynamics, ConfGF [210] directly estimates the gradient fields of the log density of atomic coordinates in molecular conformation generation.

7.6.2 Material Design. Solid state materials are the critical foundation of numerous key technologies [19]. Crystal Diffusion Variational Autoencoder (CDVAE) [256] incorporates stability as an inductive bias by proposing a noise

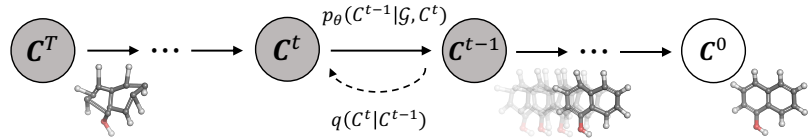


Fig. 9. Molecule-to-conformation diffusion process in GeoDiff [259].

conditional score network, which simultaneously utilizes permutation, translation, rotation, and periodic invariance properties. Luo et al. (2022) [153] model sequences and structures of complementarity-determining regions with equivariant diffusion, and explicitly target specific antigen structures to generate antibodies at atomic resolution.

7.6.3 Medical Image Reconstruction. An inverse problem is to recover an unknown signal from observed measurements, and it is an important problem in medical image reconstruction of Computed Tomography (CT) and Magnetic Resonance Imaging (MRI) [36, 37, 178, 224, 257]. Song et al. (2021) [224] utilize a score-based generative model to reconstruct an image consistent with both the prior and the observed measurements. Chung et al. (2022) [38] train a continuous time-dependent score function with denoising score matching, and iterate between the numerical SDE solver and data consistency step for reconstruction at the evaluation stage. Peng et al. (2022) [178] perform MR reconstruction by gradually guiding the reverse-diffusion process given observed k-space signal, and propose a coarse-to-fine sampling algorithm for efficient sampling.

8 FUTURE DIRECTIONS

Research on diffusion models is in its early stages, with much potential for improvement in both theoretical and empirical aspects. As discussed in early sections, key research directions include efficient sampling and improved likelihood, as well as exploring how diffusion models can handle special data structures, interface with other types of generative models, and be tailored to a range of applications. In addition, we foresee that future research on diffusion models will likely expand to the following avenues.

Revisiting Assumptions. Numerous typical assumptions in diffusion models need to be revisited and analyzed. For example, the assumption that the forward process of diffusion models completely erases any information in data and renders it equivalent to a prior distribution may not always hold. In reality, complete removal of information is unachievable in finite time. It is of great interest to understand when to halt the forward noising process in order to strike a balance between sampling efficiency and sample quality [66]. Recent advances in Schrödinger bridges and optimal transport [31, 44, 46, 212, 218] provide promising alternative solutions, suggesting new formulations for diffusion models that are capable of converging to a specified prior distribution in finite time.

Theoretical Understanding. Diffusion models have emerged as a powerful framework, notably as the only one that can rival generative adversarial networks (GANs) in most applications without resorting to adversarial training. Key to harnessing this potential is an understanding of why and when diffusion models are effective over alternatives for specific tasks. It is important to identify which fundamental characteristics differentiate diffusion models from other types of generative models, such as variational autoencoders, energy-based models, or autoregressive models. Understanding these distinctions will help elucidate why diffusion models are capable of generating samples of excellent quality while achieving top likelihood. Equally important is the need to develop theoretical guidance for selecting and determining various hyperparameters of diffusion models systematically.

Latent Representations. Unlike variational autoencoders or generative adversarial networks, diffusion models are less effective for providing good representations of data in their latent space. As a result, they cannot be easily used for tasks such as manipulating data based on semantic representations. Furthermore, since the latent space in diffusion models often possesses the same dimensionality as the data space, sampling efficiency is negatively affected and the models may not learn the representation schemes well [106].

9 CONCLUSION

We have provided a comprehensive look at diffusion models from various angles. We began with a self-contained introduction to three fundamental formulations: DDPMs, SGMs, and Score SDEs. We then discussed recent efforts to improve diffusion models, highlighting three major directions: sampling efficiency, likelihood maximization, and new techniques for data with special structures. We also explored connections between diffusion models and other generative models and outlined potential benefits of combining the two. A survey of applications across six domains illustrated the wide-ranging potential of diffusion models. Finally, we outlined possible avenues for future research.

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