

REFINING DEEP GENERATIVE MODELS VIA DISCRIMINATOR GRADIENT FLOW

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ABSTRACT

Deep generative modeling has seen impressive advances in recent years, to the point where it is now commonplace to see simulated samples (e.g., images) that closely resemble real-world data. However, generation quality is generally inconsistent for any given model and can vary dramatically between samples. We introduce Discriminator Gradient flow (DGflow), a new technique that improves generated samples via the gradient flow of entropy-regularized f -divergences between the real and the generated data distributions. The gradient flow takes the form of a *non-linear* Fokker-Plank equation, which can be easily simulated by sampling from the equivalent McKean-Vlasov process. By refining inferior samples, our technique avoids wasteful sample rejection used by previous methods (DRS & MH-GAN). Compared to existing works that focus on specific GAN variants, we show our refinement approach can be applied to GANs with vector-valued critics and even other deep generative models such as VAEs and Normalizing Flows. Empirical results on multiple synthetic, image, and text datasets demonstrate that DGflow leads to significant improvement in the quality of generated samples for a variety of generative models, outperforming the state-of-the-art Discriminator Optimal Transport (DOT) and Discriminator Driven Latent Sampling (DDLS) methods.

1 INTRODUCTION

Deep generative models (DGMs) have excelled at numerous tasks, from generating realistic images (Brock et al., 2019) to learning policies in reinforcement learning (Ho & Ermon, 2016). Among the variety of proposed DGMs, Generative Adversarial Networks (GANs) (Goodfellow et al., 2014) have received widespread popularity for their ability to generate high quality samples that resemble real data. Unlike Variational Autoencoders (VAEs) (Kingma & Welling, 2014) and Normalizing Flows (Rezende & Mohamed, 2015; Kingma & Dhariwal, 2018), GANs are likelihood-free methods; training is formulated as a minimax optimization problem involving a generator and a discriminator. The generator seeks to generate samples that are similar to the real data by minimizing a measure of discrepancy (between the generated samples and real samples) furnished by the discriminator. The discriminator is trained to distinguish the generated samples from the real samples. Once trained, the generator is used to simulate samples and the discriminator has traditionally been discarded.

However, recent work has shown that discarding the discriminator is wasteful — it actually contains useful information about the underlying data distribution. This insight has led to *sample improvement* techniques that use this information to improve the quality of generated samples (Azadi et al., 2019; Turner et al., 2019; Tanaka, 2019; Che et al., 2020). Unfortunately, current methods either rely on wasteful rejection operations in the data space (Azadi et al., 2019; Turner et al., 2019), or require a sensitive diffusion term to ensure sample diversity (Che et al., 2020). Prior work has also focused on improving GANs with scalar-valued discriminators, which excludes a large family of GANs with vector-valued critics, e.g., MMDGAN (Li et al., 2017; Bińkowski et al., 2018) and OCFGAN (Ansari et al., 2020), and likelihood-based generative models.

In this work, we propose Discriminator Gradient flow (DGflow) which formulates sample improvement as refining inferior samples using the *gradient flow* of f -divergences between the generator and the real data distributions (Fig. 1). DGflow avoids wasteful rejection operations and can be used in a deterministic setting without a diffusion term. Existing state-of-the-art methods — specifically, Discriminator Optimal Transport (DOT) (Tanaka, 2019) and Discriminator Driven Latent Sampling (DDLS) (Che et al., 2020) — can be viewed as special cases of DGflow. Similar to DDLS, DGflow recovers the real data distribution when the gradient flow is simulated exactly.

We further present a generalized framework that employs existing pre-trained discriminators to refine samples from a *variety* of deep generative models: we demonstrate our method can be applied to GANs with vector-valued critics, and even likelihood-based models such as VAEs and Normalizing Flows. Empirical results on synthetic datasets, and benchmark image (CIFAR10, STL10) and text (Billion Words) datasets demonstrate that our gradient flow-based approach outperforms DOT and DDLS on multiple quantitative evaluation metrics.

In summary, this paper’s key contributions are:

- DGflow, a method to refine deep generative models using the gradient flow of f -divergences;
- a framework that extends DGflow to GANs with vector-valued critics, VAEs, and Normalizing Flows;
- experiments on a variety of generative models trained on synthetic, image (CIFAR10 & STL10), and text (Billion Words) datasets demonstrating that DGflow is effective in improving samples from generative models.

2 BACKGROUND: GRADIENT FLOWS

The following gives a brief introduction to gradient flows; we refer readers to the excellent overview by Santambrogio (2017) for a more thorough introduction.

Let $(\mathcal{X}, \|\cdot\|_2)$ be a Euclidean space and $F : \mathcal{X} \rightarrow \mathbb{R}$ be a smooth energy function. The gradient flow of F is the smooth curve $\{\mathbf{x}_t\}_{t \in \mathbb{R}_+}$ that follows the direction of steepest descent, i.e.,

$$\mathbf{x}'(t) = -\nabla F(\mathbf{x}(t)). \quad (1)$$

The value of the energy F is minimized along this curve. This idea of steepest descent curves can be characterized in arbitrary metric spaces via the *minimizing movement scheme* (Jordan et al., 1998). Of particular interest is the metric space of probability measures that is endowed with the Wasserstein distance (\mathcal{W}_p); the Wasserstein distance is a metric and the \mathcal{W}_p topology satisfies weak convergence of probability measures (Villani, 2008, Theorem 6.9). Gradient flows in the 2-Wasserstein space ($\mathcal{P}_2(\Omega)$, \mathcal{W}_2) — i.e., the space of probability measures with finite second moments and the 2-Wasserstein metric — have been studied extensively. Let $\{\rho_t\}_{t \in \mathbb{R}_+}$ be the gradient flow of a functional \mathcal{F} in the 2-Wasserstein space, where ρ_t is absolutely continuous with respect to the Lebesgue measure. The curve $\{\rho_t\}_{t \in \mathbb{R}_+}$ satisfies the continuity equation (Ambrosio et al., 2008, Theorem 8.3.1),

$$\partial_t \rho_t + \nabla \cdot (\rho_t \mathbf{v}_t) = 0. \quad (2)$$

The velocity field \mathbf{v}_t in Eq. (2) is given by

$$\mathbf{v}_t(\mathbf{x}) = -\nabla_{\mathbf{x}} \frac{\delta \mathcal{F}}{\delta \rho}(\rho), \quad (3)$$

where $\frac{\delta \mathcal{F}}{\delta \rho}$ denotes the first variation of the functional \mathcal{F} .

Since the seminal work of Jordan et al. (1998) that showed that the Fokker-Plank equation is the gradient flow of a particular functional in the Wasserstein space, gradient flows in the Wasserstein metric have been a popular tool in the analysis of partial differential equations (PDEs). For example, they have been applied to the study of the porous-medium equation (Otto, 2001), crowd modeling (Maury et al., 2010; 2011), and mean-field games (Almulla et al., 2017). More recently, gradient flows of various distances used in deep generative modeling literature have been proposed, notably that of the sliced Wasserstein distance (Liutkus et al., 2019), the maximum mean discrepancy (Arbel et al., 2019), the Stein discrepancy (Liu, 2017), and the Sobolev discrepancy (Mroueh et al., 2019). Gradient flows have also been used for learning non-parametric and parametric implicit generative models (Liutkus et al., 2019; Gao et al., 2019; 2020). As an example of the latter, Variational Gradient Flow (Gao et al., 2019) learns a mapping between latent vectors and samples evolved using the gradient flow of f -divergences. In this work, we present a method using gradient flows of entropy-regularized f -divergences for refining samples from deep generative models employing existing discriminators as density-ratio estimators.

3 GENERATOR REFINEMENT VIA DISCRIMINATOR GRADIENT FLOW

This section describes our main contribution: Discriminator Gradient flow (DGflow). As an overview, we begin with the construction of the gradient flow of entropy-regularized f -divergences and describe its application to sample refinement. We then discuss how to simulate the gradient flow in the latent space of the generator — a procedure more suitable for high-dimensional datasets. Finally, we present a simple technique that extends our method to generative models that have not yet been studied in the context of refinement. Due to space constraints, we focus on conveying the key concepts and relegate details (e.g., proofs) to the appendix.

The entropy-regularized f -divergence functional is defined as

$$\mathcal{F}_\mu^f(\rho) \triangleq \mathcal{D}_f(\mu\|\rho) - \gamma\mathcal{H}(\rho), \quad (4)$$

where the f -divergence term $\mathcal{D}_f(\mu\|\rho)$ ensures that the “distance” between the probability density ρ and the target density μ decreases along the gradient flow. The differential entropy term $\mathcal{H}(\rho)$ improves diversity and expressiveness when the gradient flow is simulated for finite time-steps. We now construct the gradient flow of \mathcal{F}_μ^f .

Lemma 3.1. Define the functional $\mathcal{F}_\mu^f : \mathcal{P}_2(\Omega) \rightarrow \mathbb{R}$ as

$$\mathcal{F}_\mu^f(\rho) \triangleq \underbrace{\int f(\rho(\mathbf{x})/\mu(\mathbf{x})) \mu(\mathbf{x}) d\mathbf{x}}_{f\text{-divergence}} + \underbrace{\gamma \int \rho(\mathbf{x}) \log \rho(\mathbf{x}) d\mathbf{x}}_{\text{negative entropy}}, \quad (5)$$

where f is a twice-differentiable convex function with $f(1) = 0$. The gradient flow of the functional $\mathcal{F}_\mu^f(\rho)$ in the Wasserstein space ($\mathcal{P}_2(\Omega)$, \mathcal{W}_2) is given by the following PDE,

$$\partial_t \rho_t(\mathbf{x}) - \nabla_{\mathbf{x}} \cdot (\rho_t(\mathbf{x}) \nabla_{\mathbf{x}} f'(\rho_t(\mathbf{x})/\mu(\mathbf{x}))) - \gamma \Delta_{\mathbf{x}\mathbf{x}} \rho_t(\mathbf{x}) = 0, \quad (6)$$

where $\nabla_{\mathbf{x}} \cdot$ and $\Delta_{\mathbf{x}\mathbf{x}}$ denote the divergence and the Laplace operators respectively.

The proof is given in Appendix A.1. The PDE in Eq. (6) is a type of Fokker-Plank equation (FPE). FPEs have been studied extensively in the literature of stochastic processes and have a Stochastic Differential Equation (SDE) counterpart (Risken, 1996). In the case of Eq. (6), the equivalent SDE is given by

$$d\mathbf{x}_t = \underbrace{-\nabla_{\mathbf{x}} f'(\rho_t/\mu)(\mathbf{x}_t)}_{\text{drift}} dt + \underbrace{\sqrt{2\gamma} d\mathbf{w}_t}_{\text{diffusion}}, \quad (7)$$

where $d\mathbf{w}_t$ denotes the standard Wiener process. Eq. (7) defines the evolution of a particle \mathbf{x}_t under the influence of drift and diffusion. Specifically, it is a McKean-Vlasov process (Braun & Hepp, 1977) which is a type of *non-linear* stochastic process as the drift term at any time t depends on the distribution ρ_t of the particle \mathbf{x}_t . Eqs. (6) and (7) are equivalent in the sense that the distribution of the particle \mathbf{x}_t in Eq. (7) solves the PDE in Eq. (6). Consequently, samples from the density ρ_t along the gradient flow can be obtained by first drawing samples $\mathbf{x}_0 \sim \rho_0$ and then simulating the

SDE in Eq. (7). The SDE can be approximately simulated via the stochastic Euler scheme (also known as the Euler-Maruyama method) (Beyn & Kruse, 2011) given by

$$\mathbf{x}_{\tau_{n+1}} = \mathbf{x}_{\tau_n} - \eta \nabla_{\mathbf{x}} f'(\rho_{\tau_n}/\mu)(\mathbf{x}_{\tau_n}) + \sqrt{2\gamma\eta}\xi_{\tau_n}, \quad (8)$$

where $\xi_{\tau_n} \sim \mathcal{N}(\mathbf{0}, \mathbf{I})$, the time interval $[0, T]$ is partitioned into equal intervals of size η and $\tau_0 < \tau_1 < \dots < \tau_N$ denote the discretized time-steps.

Eq. (8) provides a non-parametric procedure to refine samples from a generator g_θ where we let μ be the density of real samples and ρ_{τ_0} the density of samples generated from g_θ obtained by first sampling from the prior latent distribution $\mathbf{z} \sim p_Z(\mathbf{z})$ and then feeding \mathbf{z} into g_θ . We first generate particles $\mathbf{x}_0 \sim \rho_{\tau_0}$ and then update the particles using Eq. (8) for N time steps.

Given a binary classifier (discriminator) D that has been trained to distinguish between samples from μ and ρ_{τ_0} , the density-ratio $\rho_{\tau_0}(\mathbf{x})/\mu(\mathbf{x})$ can be estimated via the well-known *density-ratio trick* (Sugiyama et al., 2012),

$$\rho_{\tau_0}(\mathbf{x})/\mu(\mathbf{x}) = \frac{1 - D(y = 1|\mathbf{x})}{D(y = 1|\mathbf{x})} = \exp(-d(\mathbf{x})), \quad (9)$$

where $D(y = 1|\mathbf{x})$ denotes the conditional probability of the sample \mathbf{x} being from μ and $d(\mathbf{x})$ denotes the logit output of the classifier D . We term this procedure where samples are refined via gradient flow of f -divergences as Discriminator Gradient flow (DGflow).

3.1 REFINEMENT IN THE LATENT SPACE

Eq. (8) requires a running estimate of the density-ratio $\rho_{\tau_n}(\mathbf{x})/\mu(\mathbf{x})$, which can be approximated using the *stale estimate* $\rho_{\tau_n}(\mathbf{x})/\mu(\mathbf{x}) \approx \rho_{\tau_0}(\mathbf{x})/\mu(\mathbf{x})$ for $\eta \rightarrow 0$ and small N , where the density ρ_{τ_n} will be close to ρ_{τ_0} . However, our initial image experiments showed that refining directly in high-dimensional data-spaces with the stale estimate is problematic; error is accumulated at each time-step leading to a visible degradation in the quality of data samples (e.g., appearance of artifacts in images).

To tackle this problem, we propose refining the latent vectors before mapping them to samples in data-space using g_θ . We describe a procedure analogous to Eq. (8) but in the latent space for generators g_θ that take a latent vector $\mathbf{z} \in \mathcal{Z}$ as input and generate a sample $\mathbf{x} \in \mathcal{X}$. We first show in Lemma 3.2 that the density-ratio in the latent space between two distributions can be estimated via the density-ratio of corresponding distributions in the data space.

Lemma 3.2. *Let $g : \mathcal{Z} \rightarrow \mathcal{X}$ be a sufficiently well-behaved injective function where $\mathcal{Z} \subseteq \mathbb{R}^n$ and $\mathcal{X} \subseteq \mathbb{R}^m$ with $m > n$. Let $p_Z(\mathbf{z})$, $p_{\hat{\mathcal{Z}}}(\hat{\mathbf{z}})$ be probability densities on \mathcal{Z} and $q_X(\mathbf{x})$, $q_{\hat{\mathcal{X}}}(\hat{\mathbf{x}})$ be the densities of the pushforward measures $g\#\mathcal{Z}$, $g\#\hat{\mathcal{Z}}$ respectively. Assume that $p_Z(\mathbf{z})$ and $p_{\hat{\mathcal{Z}}}(\hat{\mathbf{z}})$ have same support, and the Jacobian matrix \mathbf{J}_g has full column rank. Then, the density-ratio $p_{\hat{\mathcal{Z}}}(\mathbf{u})/p_Z(\mathbf{u})$ at the point $\mathbf{u} \in \mathcal{Z}$ is given by*

$$\frac{p_{\hat{\mathcal{Z}}}(\mathbf{u})}{p_Z(\mathbf{u})} = \frac{q_{\hat{\mathcal{X}}}(g(\mathbf{u}))}{q_X(g(\mathbf{u}))}. \quad (10)$$

The proof is in Appendix A.2. We let $p_{\hat{\mathcal{Z}}}(\hat{\mathbf{z}})$ be the density of the “correct” latent space distribution induced by a generator g_θ , i.e., $p_{\hat{\mathcal{Z}}}(\hat{\mathbf{z}})$ is the density of a probability measure whose pushforward under g_θ approximately equals the target data density μ . The density-ratio of the prior latent distribution $p_Z(\mathbf{z})$ and $p_{\hat{\mathcal{Z}}}(\hat{\mathbf{z}})$ can now be computed by combining Lemma 3.2 with Eq. (9),

$$\frac{p_Z(\mathbf{u})}{p_{\hat{\mathcal{Z}}}(\mathbf{u})} = \frac{\rho_{\tau_0}(g_\theta(\mathbf{u}))}{\mu(g_\theta(\mathbf{u}))} = \exp(-d(g_\theta(\mathbf{u}))). \quad (11)$$

Algorithm 1 Refinement in the Latent Space using DGflow.

Require: First derivative of f (f'), generator (g_θ), discriminator (d_ϕ), number of update steps (N), step-size (η), noise factor (γ).

- 1: $\mathbf{z}_0 \sim p_Z(\mathbf{z})$ ▷ Sample from the prior.
 - 2: **for** $i \leftarrow 0, N$ **do**
 - 3: $\xi_i \sim \mathcal{N}(0, I)$
 - 4: $\mathbf{z}_{i+1} = \mathbf{z}_i - \eta \nabla_{\mathbf{z}_i} f'(e^{-d_\phi(g_\theta(\mathbf{z}_i))}) + \sqrt{2\eta\gamma}\xi_i$
 - 5: **end for**
 - 6: **return** $g_\theta(\mathbf{z}_n)$ ▷ The refined sample.
-

Although a generator g_θ parameterized by a neural network may not satisfy the conditions of injectivity and full column rank Jacobian matrix \mathbf{J}_{g_θ} , Eq. (11) provides an approximation that works well in practice as shown by our experiments. Combining Eq. (11) with Eq. (8) provides us with an update rule for refining samples in the latent space,

$$\mathbf{u}_{\tau_{n+1}} = \mathbf{u}_{\tau_n} - \eta \nabla_{\mathbf{u}} f' (p_{\mathbf{u}_{\tau_n}} / p_{\hat{Z}}) (\mathbf{u}_{\tau_n}) + \sqrt{2\gamma\eta} \xi_{\tau_n}, \quad (12)$$

where $\mathbf{u}_{\tau_0} \sim p_Z(\mathbf{z})$ and the density-ratio $p_{\mathbf{u}_{\tau_n}} / p_{\hat{Z}}$ is approximated using the stale estimate $p_{\mathbf{u}_{\tau_0}} / p_{\hat{Z}} = \exp(-d(g_\theta(\mathbf{u})))$. We summarize the complete algorithm in Algorithm 1.

3.2 REFINEMENT FOR ALL

Thus far, prior work (Azadi et al., 2019; Turner et al., 2019; Tanaka, 2019; Che et al., 2020) has focused on improving samples for GANs with scalar-valued discriminators, which comprises the canonical GAN as well as recent variants, e.g., WGAN (Gulrajani et al., 2017), and SNGAN (Miyato et al., 2018). Here, we propose a technique that extends our approach to refine samples from a larger class of DGMs including GANs with vector-valued critics, VAEs, and Normalizing Flows.

Let p_θ be the density of the samples generated by a generator g_θ and μ be the density of the real data distribution. We are interested in refining samples from g_θ ; however, a corresponding density-ratio estimator for p_θ / μ is unavailable, as is the case with the aforementioned generative models.

Let D_ϕ be a discriminator that has been trained on the same dataset but for a different generative model g_ϕ (e.g., let g_ϕ and D_ϕ be the generator and discriminator of SNGAN respectively). D_ϕ can be used to compute the density ratio p_ϕ / μ . A straightforward technique would be to use the crude approximation $p_\theta / \mu \approx p_\phi / \mu$, which could work provided p_θ and p_ϕ are not too far from each other. Our experiments show that this simple approximation works to a limited extent (see appendix E).

To improve upon the crude approximation above, we propose to correct the density-ratio estimate. Specifically, a discriminator D_λ is initialized with the weights from D_ϕ and is fine-tuned on samples from g_ϕ and g_θ . D_ϕ and D_λ are then used to approximate the density-ratio p_θ / μ ,

$$\frac{p_\theta(\mathbf{x})}{\mu(\mathbf{x})} = \frac{p_\phi(\mathbf{x})}{\mu(\mathbf{x})} \frac{p_\theta(\mathbf{x})}{p_\phi(\mathbf{x})} = \exp(-d_\phi(\mathbf{x})) \cdot \exp(-d_\lambda(\mathbf{x})), \quad (13)$$

where d_ϕ and d_λ are logits output from D_ϕ and D_λ , respectively. We term the network D_λ the *density ratio corrector*, which experiments show produces higher quality samples than using $p_\theta / \mu \approx p_\phi / \mu$. The estimate in Eq. (13) is similar to telescoping density-ratio estimation (TRE), a technique proposed in very recent independent work (Rhodes et al., 2020). In brief, Rhodes et al. (2020) show that classifier-based density ratio estimators perform poorly when distributions are “too far apart”; the classifier can easily distinguish between the distributions, even with a poor estimate of the density ratio. TRE expands the standard density ratio into a telescoping product of more difficult-to-distinguish intermediate density ratios. Likewise, in Eq. (13), we treat p_ϕ as an intermediate distribution and estimate the final density-ratio as a product of two density-ratios.

4 RELATED WORK

Azadi et al. (2019) first proposed the idea of improving samples from a GAN’s generator by discriminator rejection sampling (DRS), making use of the density-ratio provided by the discriminator to estimate the acceptance probability. Metropolis-Hastings GAN (MH-GAN) (Turner et al., 2019) improved upon the costly rejection sampling procedure via the Metropolis-Hastings algorithm. Unlike DG flow, both of these methods reject inferior samples instead of refining them.

Recent work has also sought to improve generative models via sample evolution in the training/generation process. In energy-based generative models (Arbel et al., 2021; Deng et al., 2020), the energy functions can be viewed as a component that improves some base generator. For example, the Generalized Energy-Based Model (GEBM) (Arbel et al., 2021) jointly trains an implicit generative model with an energy function and uses Langevin dynamics to sample from the combination of the two. The Noise Conditional Score Network (NCSN) (Song & Ermon, 2019; 2020) — a score-based generative model — can be seen as a gradient flow that refines a sample right from noise to data. Latent Optimization GAN (LOGAN) (Wu et al., 2020) optimizes a latent vector via natural gradient descent as part of the GAN training process. While related, these techniques re-

sult in “self-contained” generative models; unlike DGflow, they are not general sample refinement techniques that can be applied across generative models¹.

Our method is closely related to recent state-of-the-art techniques, specifically Discriminator-Driven Latent Sampling (DDLS) (Che et al., 2020) and Discriminator Optimal Transport (DOT) (Tanaka, 2019). In fact, both these methods can be seen as special cases of DGflow.

DDLS treats a GAN as an energy-based model and uses Langevin dynamics to sample from the energy-based latent distribution $p_t(\mathbf{z}) \propto p_Z(\mathbf{z}) \exp(d(g_\theta(\mathbf{z})))$ induced by performing rejection sampling in the latent space. This distribution is the same as $p_{\hat{Z}}(\hat{\mathbf{z}})$, which can be seen by rearranging terms in Eq. (11). If we use the KL-divergence by setting $f = r \log r$, DGflow is equivalent to DDLS. However, there are practical differences that make DGflow more appealing. DDLS requires estimation of the score function $\nabla_{\mathbf{z}} \{\log p_Z(\mathbf{z}) + d(g_\theta(\mathbf{z}))\}$ to perform the update which becomes undefined if \mathbf{z} escapes the support of $p_Z(\mathbf{z})$, e.g., in the case of the uniform prior distribution commonly used in GANs; handling such cases would require techniques such as projected gradient descent. This problem does not arise in the case of DGflow since it only uses the density-ratio that is implicitly defined by the discriminator. Moreover, DDLS uses Langevin dynamics which *requires* the sensitive diffusion term to ensure diversity and to prevent points from collapsing to the maximum-likelihood point. In DGflow, the sample diversity is ensured by the density-ratio term and the diffusion term serves as an enhancement. Note that DGflow performs well even without the diffusion term (i.e., with $\gamma = 0$, see Tables 13 & 14 in the appendix). This deterministic variant of DGflow is a practical alternative with one less hyperparameter to tune.

DOT refines samples by constructing an Optimal Transport (OT) map induced by the WGAN discriminator. The OT map is realized by means of a deterministic optimization problem in the vicinity of the generated samples. If we further analyze the case of DGflow with $\gamma = 0$ and solve the resulting ordinary differential equation (ODE) using the backward Euler method,

$$\mathbf{u}_{\tau_{n+1}} = \arg \min_{\mathbf{u} \in \mathbb{R}^n} \left\{ f' \left(p_{\mathbf{u}_{\tau_n}} / p_{\hat{Z}} \right) (\mathbf{u}) + \frac{1}{2\lambda} \|\mathbf{u} - \mathbf{u}_{\tau_n}\|^2 \right\}, \quad (14)$$

DOT emerges as a special case when we consider a single update step of Eq. (14) using gradient descent and set $f'(t) = \log(t)^2$ with $\lambda = \frac{1}{2}$. This connection of DGflow to DOT, an optimal transport technique, is perhaps unsurprising given the relationship between gradient flows and the dynamical Benamou-Brenier formulation of optimal transport (Santambrogio, 2017).

5 EXPERIMENTS

In this section, we present empirical results on various deep generative models trained on multiple synthetic and real world datasets. Our primary goals were to determine if (a) DGflow is effective in improving the quality of samples from generative models, (b) the proposed extension to other generative models improves their sample quality, and (c) DGflow is generalizable to different types of data and metrics. Note that we did not seek to achieve state-of-the-art results for the datasets studied but to demonstrate that DGflow is able to significantly improve samples from the bare generators for different models.

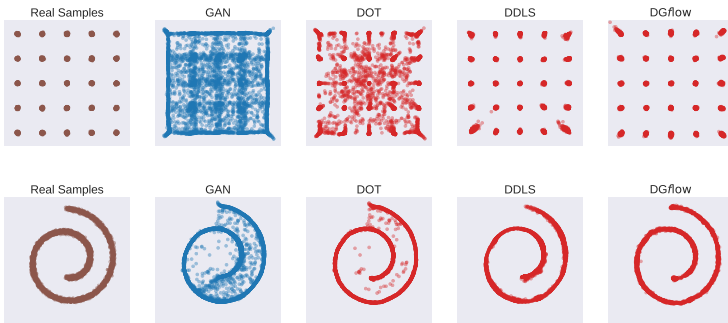
We experimented with three f -divergences, namely the Kullback-Leibler (KL) divergence, the Jensen-Shannon (JS) divergence, and the log D divergence (Gao et al., 2019). The specific forms of the functions f and corresponding derivatives are tabulated in Table 7 (appendix). We compare DGflow with two state-of-the-art competing methods: DOT and DDLS. In this section we discuss the main results and relegate details to the appendix. Our code is available online at <https://github.com/clear-nus/DGflow>.

5.1 2D DATASETS

We first tested DGflow on two synthetic datasets, 25Gaussians and 2DSwissroll, to visually inspect the improvement in the quality of generated samples. We generated 5000 samples from a trained WGAN-GP generator and refined them using DOT, DDLS, and DGflow. We performed refinement in the latent space for DDLS and directly in the data-space for DOT and DGflow. Fig. 2 shows

¹For further discussion about these techniques, please refer to appendix C.

²This implies that $f(t) = t \log t - t + 1$, which is a twice-differentiable convex function with $f(0) = 1$.

Figure 2: Qualitative comparison of $DGflow_{(KL)}$ with DOT and DDLS on synthetic 2D datasets.

the samples generated from the WGAN-GP generator (blue) and the refined samples using different techniques (red). Although the WGAN-GP generator learned the overall structure of the dataset, it also learned a number of spurious modes. DOT is able to refine the spurious samples but to a limited degree. In contrast, DDLS and $DGflow$ are able to correct almost all spurious samples and are able to recover the correct structure of the data. Visualizations for $DGflow$ with different f -divergences can be found in the appendix (Fig. 4).

We also compared the different methods quantitatively on two metrics: % high quality samples and kernel density estimate (KDE) score. A sample is classified as a high quality sample if it lies within 4 standard deviations of its nearest Gaussian. The KDE score is computed by first estimating the KDE using generated samples and then computing the log-likelihood of the training samples under the KDE estimate. We computed both the metrics 10 times using 5000 samples and report the mean in Table 1. The quantitative metrics reinforce the qualitative analysis and show that DDLS and $DGflow$ significantly improve the samples from the generator, with $DGflow$ performing slightly better than DDLS in terms of the KDE score.

5.2 IMAGE EXPERIMENTS

We conducted experiments on the CIFAR10 and STL10 datasets to demonstrate the efficacy of $DGflow$ in the real-world setting. We followed the setup of Tanaka (2019) for our image experiments. We used the Fréchet Inception Distance (FID) (Heusel et al., 2017) and Inception Score (IS) (Salimans et al., 2016) metrics to evaluate the quality of generated samples before and after refinement. A high value of IS and a low value of FID corresponds to high quality samples, respectively.

We first applied $DGflow$ to GANs with scalar-valued discriminators (e.g., WGAN-GP, SNGAN) trained on the CIFAR10 and the STL10 datasets. Table 2 shows that $DGflow$ significantly improves the quality of

Table 1: Quantitative comparison on the 25Gaussians dataset. Higher scores are better.

	% High Quality	KDE Score
GAN	$26.6 \pm .8$	-7037 ± 64
DOT	$69.8 \pm .7$	-4149 ± 39
DDLS	$89.3 \pm .6$	-2997 ± 17
$DGflow_{(KL)}$	$89.5 \pm .4$	-2893 ± 07
$DGflow_{(JS)}$	$82.6 \pm .4$	-3118 ± 19
$DGflow_{(\log D)}$	$84.5 \pm .3$	-3036 ± 14

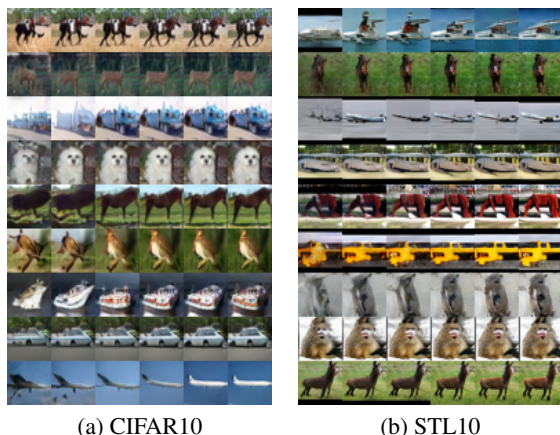
Figure 3: Improvement in the quality of samples generated from the base model (leftmost columns) over the steps of $DGflow$ for SN-ResNet-GAN and SN-DCGAN on the CIFAR10 and STL10 datasets respectively.

Table 2: Comparison of different variants of DG_{flow} with DOT on the CIFAR10 and STL10 datasets. For SN-DCGAN, (hi) denotes the hinge loss and (ns) denotes the non-saturating loss. Lower scores are better. DG_{flow}'s results have been averaged over 5 random runs with the standard deviation in parentheses.

		Fréchet Inception Distance				
Model		Base Model	DOT	DG _{flow} (KL)	DG _{flow} (JS)	DG _{flow} (log D)
CIFAR10	WGAN-GP	28.37 (.08)	24.14	24.68 (.09)	23.15 (.07)	24.53 (.11)
	SN-DCGAN (hi)	20.70 (.05)	17.12	15.68 (.07)	16.45 (.06)	17.36 (.05)
	SN-DCGAN (ns)	20.90 (.11)	15.78	15.30 (.08)	15.90 (.11)	16.42 (.05)
	SN-ResNet-GAN	14.10 (.06)	—	9.62 (.03)	9.79 (.02)	9.73 (.05)
STL10	WGAN-GP	51.50 (.15)	44.45	39.07 (.07)	50.83 (.06)	39.71 (.29)
	SN-DCGAN (hi)	40.54 (.17)	34.85	34.95 (.06)	36.37 (.12)	36.56 (.08)
	SN-DCGAN (ns)	41.86 (.12)	34.84	34.60 (.11)	35.37 (.12)	37.07 (.14)

Table 3: Inception scores of different generative models, DRS, MH-GAN, DDLS, and DG_{flow} on the CIFAR10 dataset. Higher scores are better.

Model	Inception Score
WGAN-GP (Gulrajani et al., 2017)	7.86 (.07)
ProgressiveGAN (Karras et al., 2017)	8.80 (.05)
SN-ResNet-GAN (Miyato et al., 2018)	8.22 (.05)
NCSN (Song & Ermon, 2019)	8.87 (.12)
DCGAN	2.88
DCGAN + DRS (cal) (Azadi et al., 2019)	3.07
DCGAN + MH (cal) (Turner et al., 2019)	3.38
SN-ResNet-GAN (our evaluation)	8.38 (.03)
SN-ResNet-GAN + DDLS (cal) (Che et al., 2020)	9.09 (.10)
SN-ResNet-GAN + DG _{flow} (KL)	9.35 (.03)
BigGAN	9.22

the samples in terms of the FID score and outperforms DOT on multiple models. The corresponding values of the Inception score can be found in the Appendix (Table 11), which shows DG_{flow} outperforms DOT on all models. In Table 3, we reproduce previously reported IS results for generative models and other sample improvement methods (DRS, MH-GAN, and DDLS) for completeness. DG_{flow} performs the best in terms of relative improvement from the base score and even outperforms the state-of-the-art BigGAN (Brock et al., 2019), a conditional generative model, without the need for additional labels. Qualitatively, DG_{flow} improves the vibrance of the samples and corrects deformations in the foreground object. Fig. 3 shows the change in the quality of samples when using DG_{flow} where the leftmost columns show the image generated from the base models and the successive columns show the refined sample using DG_{flow} over increments of 5 update steps.

We then evaluated the ability of DG_{flow} to refine samples from generative models *without corresponding discriminators*, namely MMDGAN, OCFGAN-GP, VAEs, and Normalizing Flows (Glow). We used the SN-DCGAN (ns) as the surrogate discriminator D_ϕ for these models and fine-tuned density ratio correctors D_λ for each model as described in section 3.2. Table 4 shows the FID scores achieved by these models without and with refinement using DG_{flow}. We obtain a clear improvement in quality of samples when these generative models are combined with DG_{flow}.

5.3 CHARACTER-LEVEL LANGUAGE MODELING

Finally, we conducted an experiment on the character-level language modeling task proposed by Gulrajani et al. (2017) to show that DG_{flow} works on different types of data. We trained a character-level GAN language model on the Billion Words Dataset (Chelba et al., 2013), which was pre-processed into 32-character long strings. We evaluated the generated samples using the JS-4 and JS-6 scores which compute the Jensen-Shannon divergence between the 4-gram and 6-gram probabilities of the data generated by the model and the real data. Table 5 (a) shows that DG_{flow} leads to

Table 4: Comparison of different variants of DGflow applied to MMDGAN, OCFGAN-GP, VAE, and Glow models. Lower scores are better. Results have been averaged over 5 random runs with the standard deviation in parentheses.

Model	Fréchet Inception Distance				
	Base Model	DGflow _(KL)	DGflow _(JS)	DGflow _(log D)	
CIFAR10	MMDGAN	41.98 (.12)	36.75 (.09)	38.06 (.14)	37.75 (.10)
	OCFGAN-GP	31.98 (.12)	26.89 (.06)	28.20 (.06)	27.82 (.09)
	VAE	129.5 (.13)	116.0 (.21)	128.9 (.13)	115.2 (.06)
	Glow	100.5 (.52)	79.02 (.23)	94.61 (.34)	81.12 (.35)
STL10	MMDGAN	47.20 (.07)	43.21 (.06)	46.74 (.05)	43.06 (.05)
	OCFGAN-GP	36.55 (.08)	31.12 (.13)	36.05 (.11)	30.61 (.14)
	VAE	150.5 (.09)	130.1 (.18)	149.9 (.08)	132.5 (.28)

Table 5: Results of DGflow on a character-level GAN language model.

(a) JS-4 and JS-6 scores. Lower scores are better.

Model	JS-4	JS-6
WGAN-GP	0.224 (.0009)	0.574 (.0015)
DGflow _(KL)	0.212 (.0008)	0.512 (.0012)
DGflow _(JS)	0.186 (.0007)	0.508 (.0011)
DGflow _(log D)	0.209 (.0005)	0.506 (.0008)

(b) Examples of text samples refined by DGflow.

Generated by WGAN-GP	Refined by DGflow
In Ruoduce that fhance would pol	In product that chance could rol
I said thowe toot lind talker .	I said this stood line talked 10
Now their rarning injurer hows	Now their warning injurer shows
Police report in B0sbu does off	Police report inturner will befe
We gine jaid 121 , one bub like	We gave wall said left out like
In years in 19mbisuch said he h	In years in 1900b such said he h

an improvement in the JS-4 and JS-6 scores. Table 5 (b) shows example sentences where DGflow visibly improves the quality of generated text.

6 CONCLUSION

In this paper, we proposed a technique to improve samples from deep generative models by refining them using gradient flow of f -divergences between the real and the generator data distributions. We also presented a simple framework that extends the proposed technique to commonly used deep generative models: GANs, VAEs, and Normalizing Flows. Experimental results indicate that gradient flows provide an excellent alternative methodology to refine generative models. Moving forward, we are considering several technical enhancements to improve DGflow’s performance. At present, DGflow uses a stale estimate of the density-ratio, which could adversely affect sample evolution when the gradient flow is simulated for larger number of steps; how we can efficiently update this estimate is an open question. Another related question is when the evolution of the samples should be stopped; running chains for too long may modify characteristics of the original sample (e.g., orientation and color) which may be undesirable. This issue does not just affect DGflow; a method for automatically stopping sample evolution could improve results across refinement techniques.

ACKNOWLEDGEMENTS

This research is supported by the National Research Foundation Singapore under its AI Singapore Programme (Award Number: AISG-RP-2019-011) to H. Soh. Thank you to J. Scarlett for his comments regarding the proofs.

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A PROOFS

A.1 LEMMA 3.1

Proof. Gradient flows in the Wasserstein space are of the form of the continuity equation (see Ambrosio et al. (2008), page 281), i.e,

$$\partial_t \rho_t + \nabla \cdot (\rho_t \mathbf{v}) = 0. \quad (15)$$

The velocity field \mathbf{v} in Eq. (15) is given by

$$\mathbf{v}(\mathbf{x}) = -\nabla_{\mathbf{x}} \frac{\delta \mathcal{F}}{\delta \rho}(\rho), \quad (16)$$

where $\frac{\delta \mathcal{F}}{\delta \rho}(\rho)$ denotes the first variation of the functional \mathcal{F} . The first variation is defined as

$$\frac{d}{d\varepsilon} \mathcal{F}(\rho + \varepsilon \chi) \Big|_{\varepsilon=0} = \int \frac{\delta \mathcal{F}}{\delta \rho}(\rho) \chi, \quad (17)$$

where $\chi = \nu - \rho$ for some $\nu \in \mathcal{P}_2(\Omega)$.

Let's derive an expression for the the first variation of \mathcal{F} . In the following, we drop the notation for dependence on \mathbf{x} for clarity,

$$\frac{d}{d\varepsilon} \mathcal{F}(\rho + \varepsilon \chi) \Big|_{\varepsilon=0} = \frac{d}{d\varepsilon} \int f \left(\frac{\rho + \varepsilon \chi}{\mu} \right) \mu + \gamma \int (\rho + \varepsilon \chi) \log(\rho + \varepsilon \chi) \Big|_{\varepsilon=0} \quad (18)$$

$$= \int f' \left(\frac{\rho + \varepsilon \chi}{\mu} \right) \chi + \gamma \int (\log(\rho + \varepsilon \chi) + 1) \chi \Big|_{\varepsilon=0} \quad (19)$$

$$= \int \left[f' \left(\frac{\rho}{\mu} \right) + \gamma \log(\rho) + \gamma \right] \chi. \quad (20)$$

Substituting $\frac{\delta \mathcal{F}}{\delta \rho}(\rho)$ in Eq. (16) we get,

$$\mathbf{v}(\mathbf{x}) = -\nabla_{\mathbf{x}} \left[f' \left(\frac{\rho}{\mu} \right) + \gamma \log(\rho) + \gamma \right] \quad (21)$$

$$= -\nabla_{\mathbf{x}} f' \left(\frac{\rho}{\mu} \right) - \frac{\gamma}{\rho} \nabla_{\mathbf{x}} \rho. \quad (22)$$

Substituting \mathbf{v} in Eq. (15) we get the gradient flow,

$$\partial_t \rho_t - \nabla_{\mathbf{x}} \cdot \left(\rho_t \nabla_{\mathbf{x}} f' \left(\frac{\rho}{\mu} \right) + \rho_t \frac{\gamma}{\rho} \nabla_{\mathbf{x}} \rho \right) = 0 \quad (23)$$

$$\partial_t \rho_t(\mathbf{x}) - \nabla_{\mathbf{x}} \cdot \left(\rho_t \nabla_{\mathbf{x}} f' \left(\frac{\rho(\mathbf{x})}{\mu(\mathbf{x})} \right) \right) - \gamma \Delta_{\mathbf{x}\mathbf{x}} \rho_t(\mathbf{x}) = 0, \quad (24)$$

where $\Delta_{\mathbf{x}\mathbf{x}}$ and $\nabla_{\mathbf{x}} \cdot$ denote the Laplace and the divergence operators respectively. \square

A.2 LEMMA 3.2

Proof. Let f be an integrable function on \mathcal{X} . If \mathbf{J}_g has full column rank and g is an injective function, then we have the following change-of-variables equation (Ben-Israel, 1999; Gemici et al., 2016),

$$\int_{\mathcal{X}} f(\mathbf{x}) d\mathbf{x} = \int_{\mathcal{Z}} (f \circ g)(\mathbf{z}) \sqrt{\det \mathbf{J}_g^\top \mathbf{J}_g(\mathbf{z})} d\mathbf{z}. \quad (25)$$

This implies that the infinitesimal volumes $d\mathbf{x}$ and $d\mathbf{z}$ are related as $d\mathbf{x} = \sqrt{\det \mathbf{J}_g^\top \mathbf{J}_g(\mathbf{z})} d\mathbf{z}$ and the densities $p_Z(\mathbf{z})$ and $q_X(\mathbf{x})$ are related as $p_Z(\mathbf{z}) = q_X(g(\mathbf{z})) \sqrt{\det \mathbf{J}_g^\top \mathbf{J}_g(\mathbf{z})}$. Similarly, $p_{\hat{Z}}(\hat{\mathbf{z}}) = q_{\hat{X}}(g(\hat{\mathbf{z}})) \sqrt{\det \mathbf{J}_g^\top \mathbf{J}_g(\hat{\mathbf{z}})}$. Finally, the density-ratio $p_{\hat{Z}}(\mathbf{u})/p_Z(\mathbf{u})$ at the point $\mathbf{u} \in \mathcal{Z}$ is given by

$$\frac{p_{\hat{Z}}(\mathbf{u})}{p_Z(\mathbf{u})} = \frac{q_{\hat{X}}(g(\mathbf{u})) \sqrt{\det \mathbf{J}_g^\top \mathbf{J}_g(\mathbf{u})}}{q_X(g(\mathbf{u})) \sqrt{\det \mathbf{J}_g^\top \mathbf{J}_g(\mathbf{u})}} = \frac{q_{\hat{X}}(g(\mathbf{u}))}{q_X(g(\mathbf{u}))}. \quad (26)$$

□

B A DISCUSSION ON DGflow FOR WGAN

We apply DGflow to WGAN models by treating the output from their critics as the logit for the estimation of density-ratio. However, it is well-known that WGAN critics are not density-ratio estimators as they are trained to maximize the 1-Wasserstein distance with an unconstrained output. In this section, we provide theoretical justification for the good performance of DGflow on WGAN models. We show that DGflow is related to the gradient flow of the entropy-regularized 1-Wasserstein functional $\mathcal{F}_\mu^W : \mathcal{P}_2(\Omega) \rightarrow \mathbb{R}$,

$$\mathcal{F}_\mu^W(\rho) \triangleq \underbrace{\sup_{\|d\|_{\text{Lip}} \leq 1} \int d(\mathbf{x}) \mu(\mathbf{x}) d\mathbf{x} - \int d(\mathbf{x}) \rho(\mathbf{x}) d\mathbf{x}}_{\text{1-Wasserstein distance}} + \underbrace{\gamma \int \rho(\mathbf{x}) \log \rho(\mathbf{x}) d\mathbf{x}}_{\text{negative entropy}}, \quad (27)$$

where μ denotes the target density, $\|d\|_{\text{Lip}}$ denotes the Lipschitz constant of the function d .

Let d^* be the function that achieves the supremum in Eq. (27). This results in the functional,

$$\mathcal{F}_\mu^W(\rho) = \int d^*(\mathbf{x}) \mu(\mathbf{x}) d\mathbf{x} - \int d^*(\mathbf{x}) \rho(\mathbf{x}) d\mathbf{x} + \gamma \int \rho(\mathbf{x}) \log \rho(\mathbf{x}) d\mathbf{x}. \quad (28)$$

Following a similar derivation as in Appendix A.1, the gradient flow of $\mathcal{F}_\mu^W(\rho)$ is given by the following PDE,

$$\partial_t \rho_t(\mathbf{x}) + \nabla_{\mathbf{x}} \cdot (\rho_t \nabla_{\mathbf{x}} d^*(\mathbf{x})) - \gamma \Delta_{\mathbf{xx}} \rho_t(\mathbf{x}) = 0. \quad (29)$$

If d^* is approximated using the critic (d_ϕ) of WGAN, we get the following gradient flow,

$$\partial_t \rho_t(\mathbf{x}) + \nabla_{\mathbf{x}} \cdot (\rho_t \nabla_{\mathbf{x}} d_\phi(\mathbf{x})) - \gamma \Delta_{\mathbf{xx}} \rho_t(\mathbf{x}) = 0, \quad (30)$$

which is same as the gradient flow of entropy-regularized f -divergence with $f = r \log r$ (i.e., the KL divergence) when d_ϕ is treated as a density-ratio estimator. The gradient flow of entropy-regularized f -divergence with $f = r \log r$ is simplified below,

$$\partial_t \rho_t(\mathbf{x}) - \nabla_{\mathbf{x}} \cdot (\rho_t \nabla_{\mathbf{x}} f'(\exp(-d_\phi(\mathbf{x})))) - \gamma \Delta_{\mathbf{xx}} \rho_t(\mathbf{x}) = 0 \quad (31)$$

$$\partial_t \rho_t(\mathbf{x}) - \nabla_{\mathbf{x}} \cdot (\rho_t \nabla_{\mathbf{x}} (\log(\exp(-d_\phi(\mathbf{x}))) + 1)) - \gamma \Delta_{\mathbf{xx}} \rho_t(\mathbf{x}) = 0 \quad (32)$$

$$\partial_t \rho_t(\mathbf{x}) + \nabla_{\mathbf{x}} \cdot (\rho_t \nabla_{\mathbf{x}} d_\phi(\mathbf{x})) - \gamma \Delta_{\mathbf{xx}} \rho_t(\mathbf{x}) = 0. \quad (33)$$

The equality of Eq. (30) and Eq. (33) implies that DGflow approximates the gradient flow of the 1-Wasserstein distance when the critic of WGAN is used for density-ratio estimation.

C FURTHER DISCUSSION ON RELATED WORK

Energy-based & Score-based Generative Models DGflow is related to recently proposed energy-based generative models (Arbel et al., 2021; Deng et al., 2020) — one can view the energy functions used in these methods as a component that improves some base model. For example, the Generalized Energy-Based Model (GEBM) (Arbel et al., 2021) jointly trains an implicit generative model with an energy function and uses Langevin dynamics to sample from the combination

of the two. Similarly, in Deng et al. (2020), a discriminator that estimates the energy function is combined with a language model to train an energy-based text-generation model.

Score-based generative modeling (SBGM) (Song & Ermon, 2019; 2020) is another active area of research closely-related to energy-based models. Noise Conditional Score Network (NCSN) (Song & Ermon, 2019; 2020), a SBGM, trains a neural network to estimate the score function of a probability density at various noise levels. Once trained, this score network is used to evolve samples from noise to the data distribution using Langevin dynamics. NCSN can be viewed as a gradient flow that refines a sample right from noise to data; however, unlike DGflow, NCSN is a complete generative models in itself and not a sample refinement technique that can be applied to other generative models.

Other Related Work Monte Carlo techniques have been used for improving various components in generative models, e.g., Grover et al. (2018) proposed Variational Rejection Sampling which performs rejection sampling in the latent space of VAEs to improve the variational posterior and Grover et al. (2019) used likelihood-free importance sampling for bias correction in generative models . Wu et al. (2020) proposed Latent Optimization GAN (LOGAN) which optimizes the latent vector as part of the training process unlike DGflow that refines the latent vector post training.

D IMPLEMENTATION DETAILS

D.1 2D DATASETS

Datasets The 25 Gaussians dataset was constructed by generating 100000 samples from a mixture of 25 equally likely 2D isotropic Gaussians with means $\{-4, -2, 0, 2, 4\} \times \{-4, -2, 0, 2, 4\} \subset \mathbb{R}^2$ and standard deviation 0.05. Once generated, the data-points were normalized by $2\sqrt{2}$ following Tanaka (2019). The 2DSwissroll dataset was constructed by first generating 100000 samples of the 3D swissroll dataset using `make_swiss_roll` from `scikit-learn` with `noise=0.25` and then only keeping dimensions $\{0, 2\}$. The generated samples were normalized by 7.5.

Base Models We trained a WGAN-GP model for both the datasets. The generator was a fully-connected network with ReLU non-linearities that mapped $z \sim \mathcal{N}(0, I_{2 \times 2})$ to $x \in \mathbb{R}^2$. Similarly, the discriminator was a fully-connected network with ReLU non-linearities that mapped $x \in \mathbb{R}^2$ to \mathbb{R} . We refer the reader to Gulrajani et al. (2017) for the exact network structures. The gradient penalty factor was set to 10. The models were trained for 10K generator iterations with a batch size of 256 using the Adam optimizer with a learning rate of 10^{-4} , $\beta_1 = 0.5$, and $\beta_2 = 0.9$. We updated the discriminator 5 times for each generator iteration.

Hyperparameters We ran DOT for 100 steps and performed gradient descent using the Adam optimizer with a learning rate of 0.01 and $\beta = (0., 0.9)$ as suggested by Tanaka (2019). DDLS was run for 50 iterations with a step-size of 0.01 and the Gaussian noise was scaled by a factor of 0.1 as suggested by Che et al. (2020). For DGflow, we set the step-size $\eta = 0.01$, the number of steps $n = 100$, and the noise regularizer $\gamma = 0.01$. We used the output from the WGAN-GP discriminator directly as a logit for estimating the density ratio for DDLS and DGflow.

Metrics We compared the different methods quantitatively on two metrics: % high quality samples and kernel density estimate (KDE) score. A sample is classified as a high quality sample if it lies within 4 standard deviations of its nearest Gaussian. The KDE score is computed by first estimating the KDE using generated samples and then computing the log-likelihood of the training samples under the KDE estimate. KDE was performed using `sklearn.neighbors.KernelDensity` with a Gaussian kernel and a kernel bandwidth of 0.1. The quantitative metrics were averaged over 10 runs with 5000 samples from each method.

D.2 IMAGE EXPERIMENTS

Datasets CIFAR10 (Krizhevsky et al., 2009) is a dataset of 60K natural RGB images of size 32×32 from 10 classes. STL10 is a dataset of 100K natural RGB images of size 96×96 from 10

Table 6: Network architectures used for MMDGAN and VAE models.

(a) Generator or Decoder	(b) Discriminator or Encoder
Input Shape: (b, d, 1, 1)	Input Shape: (b, 3, 32, 32)
Upconv(256)	Conv(64)
BatchNorm	LeakyReLU(0.2)
ReLU	Conv(128)
Upconv(128)	BatchNorm
BatchNorm	LeakyReLU(0.2)
ReLU	Conv(256)
Upconv(64)	BatchNorm
BatchNorm	LeakyReLU(0.2)
ReLU	Conv(m)
Upconv(3)	Output Shape: (b, m, 1, 1)
Tanh	
Output Shape: (b, 3, 32, 32)	

classes. We resized the STL10 (Coates et al., 2011) dataset to 48×48 for SNGAN and WGAN-GP, and to 32×32 for MMDGAN, OCFGAN-GP, and VAE since the respective base models were trained on these sizes.

Base Models for CIFAR10 We used the publicly available pre-trained models for WGAN-GP, SN-DCGAN (hi), and SN-DCGAN (ns). We refer the reader to Tanaka (2019) for exact details about these models. For SN-ResNet-GAN and OCFGAN-GP we used the pre-trained models from Miyato et al. (2018) and Ansari et al. (2020) respectively. We used the respective discriminators of SN-DCGAN (ns), SN-DCGAN (hi), and WGAN-GP for density-ratio estimation when refining their generators. For the SN-ResNet-GAN (hi) generator, we used SN-DCGAN (ns) discriminator as the non-saturating loss provides a better density-ratio estimation than a discriminator trained using the hinge loss.

We trained our own models for MMDGAN, VAE, and Glow. We used the generator and discriminator architectures shown in Table 6 for MMDGAN with $d = 32$. VAE used the same architecture with $d = 64$. Our Glow model was trained using the code available at <https://github.com/y0ast/Glow-PyTorch> with a batch size of 56 for 150 epochs. The density ratio correctors, D_λ (see section 3.2), were initialized with the weights from the SN-DCGAN (ns) released by Tanaka (2019). D_λ was then fine-tuned on images from SN-DCGAN (ns)'s generator and the generator being improved (e.g., MMDGAN and OCFGAN-GP) using SGD with a learning rate of 10^{-4} and momentum of 0.9. We fine-tuned D_λ for 10000 iterations with a batch size of 64.

Base Models for STL10 We used the publicly available pre-trained models (Tanaka, 2019; Ansari et al., 2020) for WGAN-GP, SN-DCGAN (hi), SN-DCGAN (ns), and OCFGAN-GP. We trained our own models for MMDGAN and VAE with the same architecture and training details as CIFAR10. We fine-tuned the density ratio correctors for STL10 for 5000 iterations with other details being the same as CIFAR10.

Hyperparameters We performed 25 updates of DGflow for CIFAR10 and STL10 with a step size of 0.1 for models that do not require density ratio corrections. For STL10 models that require a density ratio correction, we performed 15 updates with a step size of 0.05. The noise regularizer (γ), whenever used, was set to 0.01.

Metrics We used the Fréchet Inception Distance (FID) (Heusel et al., 2017) and Inception Score (IS) (Salimans et al., 2016) metrics to evaluate the quality of generated samples before and after refinement. The IS denotes the confidence in classification of the generated samples using a pretrained InceptionV3 network whereas the FID is the Fréchet distance between multivariate Gaussians fitted to the 2048 dimensional feature vectors extracted from the InceptionV3 network for real and generated data. Both the metrics were computed using 50K samples for all the models, except Glow

Table 7: f -divergences and their derivatives.

f -divergence	f	f'	f''
KL	$r \log r$	$\log r + 1$	$\frac{1}{r}$
JS	$r \log r - (r+1) \log \frac{r+1}{2}$	$\log \frac{2r}{r+1}$	$\frac{1}{r^2+r}$
log D	$(r+1) \log(r+1) - 2 \log 2$	$\log(r+1) + 1$	$\frac{1}{r+1}$

where we used 10K samples. Following Tanaka (2019), we used the entire training and test set (60K images) for CIFAR10 and the entire unlabeled set (100K images) for STL10 as the set of real images used to compute FID.

D.3 CHARACTER LEVEL LANGUAGE MODELING

Dataset We used the Billion Words dataset (Chelba et al., 2013) which was pre-processed into 32-character long strings.

Base Model Our generator was a 1D CNN which followed the architecture used by Gulrajani et al. (2017).

Hyperparameters We performed 50 updates of DGflow with a step size of 0.1 and noise factor $\gamma = 0$.

Metrics The JS-4 and JS-6 scores were computed using the code provided by Gulrajani et al. (2017) at https://github.com/igul222/improved_wgan_training. We used 10000 samples from the models to compute the JS-4 score.

E ADDITIONAL RESULTS

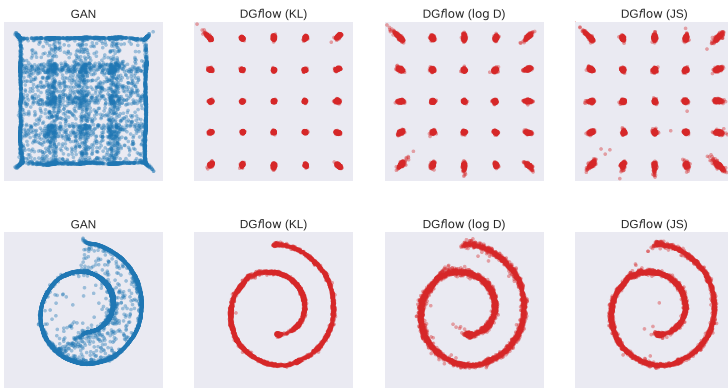


Figure 4: Qualitative comparison of DGflow with different f -divergences on the 25Gaussians and 2DSwissroll datasets.

Fig. 4 shows the samples generated by WGAN-GP (leftmost, blue) and refined samples generated using DGflow with different f -divergences (red). Fig. 5 shows the deterministic component, $-\nabla_{\mathbf{x}}f'(\rho_0/\mu)(\mathbf{x}_0)$, of the velocity for different f -divergences on the 2D datasets. Fig. 6 (right) shows the latent space distribution recovered by DGflow when applied in the latent space for the 2D datasets. This latent space is same as the one derived by Che et al. (2020), i.e., $p_t(\mathbf{z}) \propto p_Z(\mathbf{z}) \exp(d(g_\theta(\mathbf{z})))$ which is shown in Fig. 6 (left) for both datasets.

Table 11 shows the comparison of DGflow with DOT in terms of the inception score for the CIFAR10 and STL10 datasets. DGflow outperforms DOT significantly for all the base GAN models on both the datasets. Table 12 compares different variants of DGflow applied to MMDGAN, OCFGAN-GP, VAE, and Glow generators in terms of the inception score. DGflow leads to a

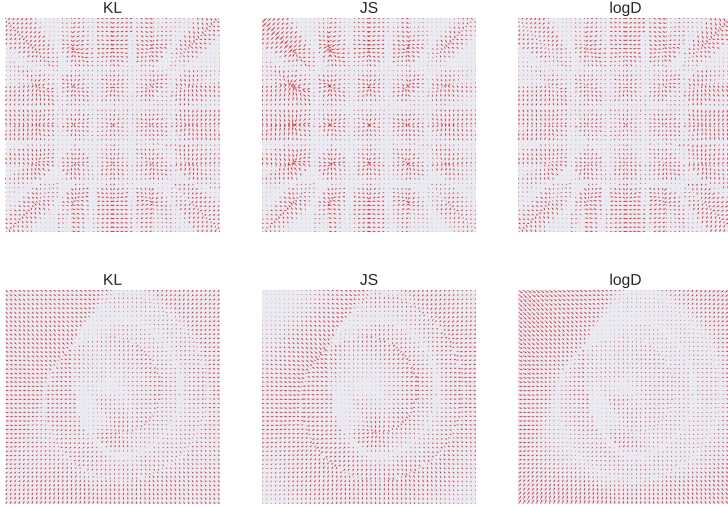


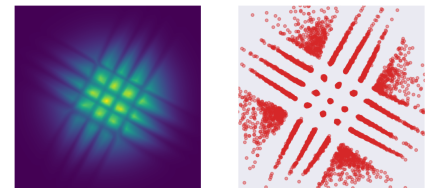
Figure 5: A vector plot showing the deterministic component of the velocity, i.e., the drift $-\nabla_{\mathbf{x}}f'(\rho_0/\mu)(\mathbf{x}_0)$, for different f -divergences on the 25Gaussians and 2DSwissroll dataset.

Table 8: Runtime comparison of DOT, DDLS, and DG $\text{flow}_{(\text{KL})}$ on the 25Gaussians dataset. The runtime is averaged over 100 runs with standard deviation reported in parentheses.

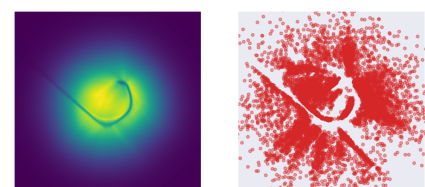
Method	Runtime (s) per 5K samples
DOT	2.24 (0.18)
DDLS	2.23 (0.14)
DG flow	2.22 (0.15)

significant improvement in the quality of samples for all the models. Tables 13 and 14 compare the deterministic variant of DG flow ($\gamma = 0$) against DOT and DDLS. These results show that the diffusion term only serves as an enhancement for DG flow , not a necessity, and it outperforms competing methods even without added noise. Table 15 shows the results of DG flow on MMDGAN, OCFGAN-GP, and VAE models when the SN-DCGAN (ns) discriminator is directly used as a density-ratio estimator without an additional density-ratio corrector. Figures 7, 8, 9, and 10 show the samples generated by the base model (left) and the refined samples (right) using DG flow for the CIFAR10 and STL10 datasets.

Runtime DG flow performs a backward pass through $d_\phi \circ g_\theta$ to compute the gradient of the density-ratio with respect to the latent vector. This results in the same runtime complexity as that of DOT and DDLS. Table 8 shows a comparison of the runtimes of DOT, DDLS, and, DG flow on the 25Gaussians dataset under same conditions. As expected, these refinement methods have similar runtimes in practice. The wall-clock time required for DG $\text{flow}_{(\text{KL})}$ to refine 100 samples from different base models on the CIFAR10 and STL10 datasets is reported in tables 9 and 10.



(a) 25 Gaussians



(b) 2D Swissroll

Figure 6: Latent space recovered by DG flow (right) for the 2D datasets is same as the one derived by Che et al. (2020) (left).

Table 9: Runtime of $DGflow_{(KL)}$ for models that do not require density-ratio correction on a single GeForce RTX 2080 Ti GPU. The runtime is averaged over 100 runs with standard deviation reported in parentheses.

	Model	Runtime (s) per 100 samples
CIFAR10	WGAN-GP	0.897 (0.017)
	SN-DCGAN (hi)	0.952 (0.008)
	SN-DCGAN (ns)	0.952 (0.007)
	SN-ResNet-GAN	1.982 (0.013)
STL10	WGAN-GP	1.376 (0.025)
	SN-DCGAN (hi)	1.413 (0.015)
	SN-DCGAN (ns)	1.415 (0.013)

Table 10: Runtime of $DGflow_{(KL)}$ for models that require density-ratio correction on a single GeForce RTX 2080 Ti GPU. The runtime is averaged over 100 runs with standard deviation reported in parentheses.

	Model	Runtime (s) per 100 samples
CIFAR10	MMDGAN	1.192 (0.007)
	OCFGAN-GP	1.186 (0.011)
	VAE	1.186 (0.012)
STL10	MMDGAN	1.036 (0.004)
	OCFGAN-GP	1.029 (0.010)
	VAE	1.028 (0.011)

Table 11: Comparison of different variants of $DGflow$ with DOT on the CIFAR10 and STL10 datasets. Higher scores are better.

Model	Inception Score				
	Base Model	DOT	$DGflow_{(KL)}$	$DGflow_{(JS)}$	$DGflow_{(\log D)}$
CIFAR10	WGAN-GP	6.51 (.02)	7.45	7.99 (.02)	7.71 (.02)
	SN-DCGAN (hi)	7.35 (.03)	8.02	8.13 (.02)	7.98 (.01)
	SN-DCGAN (ns)	7.38 (.03)	7.97	8.14 (.03)	7.98 (.04)
	SN-ResNet-GAN	8.38 (.03)	–	9.35 (.03)	9.13 (.04)
STL10	WGAN-GP	8.72 (.02)	9.31	10.41 (.02)	8.85 (.06)
	SN-DCGAN (hi)	8.77 (.03)	9.35	9.74 (.04)	9.50 (.05)
	SN-DCGAN (ns)	8.61 (.04)	9.45	9.66 (.01)	9.49 (.03)

Table 12: Comparison of different variants of $DGflow$ applied to MMDGAN, OCFGAN-GP, VAE, and Glow models. Higher scores are better.

Model	Inception Score			
	Base Model	$DGflow_{(KL)}$	$DGflow_{(JS)}$	$DGflow_{(\log D)}$
CIFAR10	MMDGAN	5.74 (.02)	6.27 (.05)	5.99 (.03)
	OCFGAN-GP	6.52 (.02)	7.21 (.05)	6.93 (.03)
	VAE	3.20 (.01)	3.85 (.01)	3.21 (.02)
	Glow	3.64 (.02)	4.57 (.02)	3.91 (.03)
CIFAR10	MMDGAN	6.07 (.02)	6.16 (.01)	6.12 (.03)
	OCFGAN-GP	7.09 (.01)	7.46 (.04)	7.10 (.03)
	VAE	3.25 (.01)	3.72 (.04)	3.27 (.01)

Table 13: Comparison of different variants of DG f low without diffusion (i.e., $\gamma = 0$) on the CIFAR10 and STL10 datasets. Lower scores are better.

		Fréchet Inception Distance				
Model		Base Model	DOT	DG f low _(KL)	DG f low _(JS)	DG f low _(log D)
CIFAR10	WGAN-GP	28.34 (.11)	24.14	24.64 (.13)	23.30 (.11)	24.42 (.19)
	SN-DCGAN (hi)	20.67 (.09)	17.12	15.79 (.07)	16.79 (.09)	17.79 (.05)
	SN-DCGAN (ns)	20.94 (.12)	15.78	15.47 (.11)	16.32 (.11)	16.97 (.08)
STL10	WGAN-GP	51.34 (.21)	44.45	38.96 (.08)	50.44 (.09)	39.35 (.12)
	SN-DCGAN (hi)	40.82 (.16)	34.85	35.18 (.09)	36.53 (.13)	36.75 (.13)
	SN-DCGAN (ns)	41.83 (.20)	34.84	34.81 (.08)	35.75 (.10)	37.68 (.08)

Table 14: Comparison of DDLS with DG f low (with and without diffusion) on the CIFAR10 dataset. Higher scores are better.

Model	Inception Score
SN-ResNet-GAN (Miyato et al., 2018)	8.22 (.05)
SN-ResNet-GAN + DDLS (cal) (Che et al., 2020)	9.09 (.10)
SN-ResNet-GAN (our evaluation)	8.38 (.03)
SN-ResNet-GAN + DG f low _(KL) ($\gamma = 0$)	9.35 (.04)
SN-ResNet-GAN + DG f low _(KL) ($\gamma = 0.01$)	9.35 (.03)
BigGAN	9.22

Table 15: Comparison of different variants of DG f low applied to MMDGAN, OCFGAN-GP, and VAE models without density-ratio correction. Lower scores are better.

		Fréchet Inception Distance			
Model		Base Model	KL	JS	log D
CIFAR10	MMDGAN	42.03 (.06)	39.06 (.08)	39.68 (.06)	39.47 (.07)
	OCFGAN-GP	31.95 (.07)	27.92 (.08)	29.25 (.06)	28.82 (.10)
	VAE	129.49 (.19)	127.50 (.15)	128.24 (.11)	128.3 (.14)
	Glow	100.7 (.14)	93.47 (.09)	97.50 (.11)	97.78 (.14)
STL10	MMDGAN	47.22 (.04)	45.75 (.10)	45.96 (.07)	46.26 (.13)
	OCFGAN-GP	36.60 (.15)	34.17 (.18)	34.42 (.04)	34.99 (.07)
	VAE	150.49 (.07)	151.76 (.01)	152.03 (.05)	151.88 (.11)



Figure 7: Samples from different models for the CIFAR10 dataset before (left) and after (right) refinement using DGflow.



Figure 8: Samples from different models for the CIFAR10 dataset before (left) and after (right) refinement using DGflow.



Figure 9: Samples from different models for the STL10 dataset before (left) and after (right) refinement using DG^flow.



Figure 10: Samples from different models for the STL10 dataset before (left) and after (right) refinement using DGflow.