# Repulsive Deep Ensembles are Bayesian

Francesco D'Angelo ETH Zürich Zürich, Switzerland fdangelo@ethz.ch

Vincent Fortuin
ETH Zürich
Zürich, Switzerland
fortuin@inf.ethz.ch

### **Abstract**

Deep ensembles have recently gained popularity in the deep learning community for their conceptual simplicity and efficiency. However, maintaining functional diversity between ensemble members that are independently trained with gradient descent is challenging. This can lead to pathologies when adding more ensemble members, such as a saturation of the ensemble performance, which converges to the performance of a single model. Moreover, this does not only affect the quality of its predictions, but even more so the uncertainty estimates of the ensemble, and thus its performance on out-of-distribution data. We hypothesize that this limitation can be overcome by discouraging different ensemble members from collapsing to the same function. To this end, we introduce a kernelized repulsive term in the update rule of the deep ensembles. We show that this simple modification not only enforces and maintains diversity among the members but, even more importantly, transforms the maximum a posteriori inference into proper Bayesian inference. Namely, we show that the training dynamics of our proposed repulsive ensembles follow a Wasserstein gradient flow of the KL divergence with the true posterior. We study repulsive terms in weight and function space and empirically compare their performance to standard ensembles and Bayesian baselines on synthetic and real-world prediction tasks.

# 1 Introduction

There have been many recent advances on the theoretical properties of sampling algorithms for approximate Bayesian inference, which changed our interpretation and understanding of them. Particularly worth mentioning is the work of Jordan et al. [34], who reinterpret Markov Chain Monte Carlo (MCMC) as a gradient flow of the KL divergence over the Wasserstein space of probability measures. This new formulation allowed for a deeper understanding of approximate inference methods but also inspired the inception of new and more efficient inference strategies.

Following this direction, Liu and Wang [46] recently proposed the Stein Variational Gradient Descent (SVGD) method to perform approximate Wasserstein gradient descent. Conceptually, this method, that belongs to the family of particle-optimization variational inference (POVI), introduces a repulsive force through a kernel acting in the parameter space to evolve a set of samples towards high-density regions of the target distribution without collapsing to a point estimate. However, when dealing with neural networks, the Bayesian approach is not the only one capable of offering calibrated models that allow for uncertainty estimation; ensembles of neural networks (so-called *deep ensembles*) have indeed achieved great success recently, both in terms of predictive performance [38, 70] as well as uncertainty estimation [57], and constitute the main competitor to the Bayesian counterpart. That being said, while they might allow for the averaging of predictions over several hypotheses, they do not offer any guarantees for the diversity between those hypotheses nor do they benefit of the advantages of the probabilistic Bayesian framework.

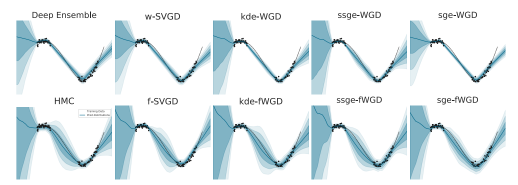


Figure 1: **BNN 1D regression.** The function-space methods (SVGD and WGD) approach the HMC posterior more closely, while the standard deep ensembles and weight-space methods fail to properly account for the uncertainty, especially the in-between uncertainty.

In this work, we show how the introduction of a repulsive term between the members in the ensemble, inspired by SVGD, not only naïvely guarantees the diversity among the members, avoiding their collapse in the parameter space, but also allows for a reformulation of the method as a gradient flow of the KL divergence in the Wasserstein space of distributions. It thus allows to reformulate deep ensembles with repulsion as a truly Bayesian method.

Nevertheless, BNN inference in weight space can lead to degenerate solutions, due to the over-parametrization of these models. That is, several samples could have very different weights but map to the same function, thus giving a false sense of diversity in the ensemble. This property, that we will refer to as *non-identifiability* of neural networks (see Appendix A), can lead to redundancies in the posterior distribution. It implies that methods like MCMC sampling, deep ensembles, and SVGD waste computation in local modes that account for equivalent functions. Predictive distributions approximated using samples from these modes do not improve over a simple point estimate and lead to a poor uncertainty estimation. Following this idea, Wang et al. [66] introduced a new method to extend POVI methods to function space, overcoming this limitation. Here, we also study an update rule that allows for an approximation of the gradient flow of the KL divergence in function space instead of weight space.

We make the following contributions:

- We derive several different repulsion terms that can be added as regularizers to the gradient updates of deep ensembles to endow them with Bayesian properties.
- We show that these terms approximate Wasserstein gradient flows of the KL divergence and can be used both in weight space and function space.
- We compare these proposed methods theoretically to standard deep ensembles and SVGD and highlight their different guarantees.
- We assess all these methods on synthetic and real-world deep learning tasks and show
  that our proposed repulsive ensembles can achieve competitive performance and improved
  uncertainty estimation.

### 2 Repulsive Deep Ensembles

In supervised deep learning, we typically consider a likelihood function  $p(\mathbf{y}|f(\mathbf{x};\mathbf{w}))$  (e.g., Gaussian for regression or Categorical for classification) parameterized by a neural network  $f(\mathbf{x};\mathbf{w})$  and training data  $\mathcal{D} = \{(\mathbf{x}_i, \mathbf{y}_i)\}_{i=1}^n$ . In Bayesian neural networks (BNNs), we are interested in the posterior distribution of all likely networks given by  $p(\mathbf{w}|\mathcal{D}) \propto \prod_{i=1}^n p(\mathbf{y}_i|f(\mathbf{x}_i;\mathbf{w})) p(\mathbf{w})$ , where  $p(\mathbf{w})$  is the prior distribution over weights. Crucially, when making a prediction on a test point  $\mathbf{x}^*$ , in the Bayesian approach we do not only use a single parameter  $\hat{\mathbf{w}}$  to predict  $\mathbf{y}^* = f(\mathbf{x}^*; \hat{\mathbf{w}})$ , but we marginalize over the whole posterior, thus taking all possible explanations of the data into account:

$$p(\mathbf{y}^*|\mathbf{x}^*, \mathcal{D}) = \int p(\mathbf{y}^*|f(\mathbf{x}^*; \mathbf{w})) p(\mathbf{w}|\mathcal{D}) d\mathbf{w}$$
(1)

While approximating the posterior of deep neural networks (or sampling from it) is a challenging task, performing maximum a posteriori (MAP) estimation, which corresponds to finding the mode of the posterior, is usually simple. Ensembles of neural networks use the non-convexity of the MAP optimization problem to create a collection of K independent—and possibly different—solutions. Considering n weight configurations of neural networks  $\{\mathbf{w}_i\}_{i=1}^n$  with  $\mathbf{w}_i \in \mathbb{R}^d$ , the dynamics of the ensemble under the gradient of the posterior lead to the following update rule at iteration t:

$$\mathbf{w}_{i}^{t+1} \leftarrow \mathbf{w}_{i}^{t} + \epsilon_{t} \phi(\mathbf{w}_{i}^{t})$$
with 
$$\phi(\mathbf{w}_{i}^{t}) = \nabla_{\mathbf{w}_{i}^{t}} \log p(\mathbf{w}_{i}^{t}|\mathcal{D}), \qquad (2)$$

with step size  $\epsilon_t$ . Ensemble methods have a long history [e.g., 41, 22, 5] and were recently revisited for neural networks [38] and coined *deep ensembles*. The predictions of the different members are combined to create a predictive distribution by using the solutions to compute the Bayesian model average (BMA) in Eq. (1). Recent works [57] have shown that deep ensembles can outperform some of the Bayesian approaches for uncertainty estimation. Even more recently, Wilson and Izmailov [72] argued that deep ensembles can be considered a compelling approach to Bayesian model averaging. Despite these ideas, the ability of deep ensembles to efficiently average over multiple hypotheses and to explore the functional landscape of the posterior distribution studied in [15] does not make the method necessarily Bayesian. Indeed, the additional Langevin noise introduced in [67], which is not considered in deep ensembles, is crucial to ensure samples from the correct posterior distribution.

From a practical standpoint, since the quality of an ensemble hinges on the diversity of its members, many methods were recently proposed to improve this diversity without compromising the individual accuracy. For instance, Wenzel et al. [70] propose hyper-deep ensembles that combine deep networks with different hyperparameters. Similarly, cyclical learning-rate schedules can explore several local minima for the ensemble members [29]. Alternatively, Rame and Cord [59] proposed an information-theoretic framework to avoid redundancy in the members. However, the absence of a constraint that prevents particles from converging to the same mode limits the possibility of improvement by introducing more ensemble members. This means that any guarantees to converge to different modes must exclusively rely on:

- 1. The randomness of the initialization.
- 2. The noise in the estimation of the gradients due to minibatching.
- 3. The number of local optima that might be reached during gradient descent.

Moreover, the recent study of Geiger et al. [21] showed how the empirical test error of the ensemble converges to the one of a single trained model when the number of parameters goes to infinity, leading to deterioration of the performance. In other words, the bigger the model, the harder it is to maintain diversity in the ensemble and avoid collapse to the same solution. This is intuitively due to the fact that bigger models are less sensitive to the initialization. Namely, in order for them to get stuck in a local minimum, they must have second derivatives that are positive simultaneously in all directions. As the number of hidden units gets larger, this becomes less likely.

# 2.1 Repulsive force in weight space

To overcome the aforementioned limitations of standard deep ensembles, we introduce, inspired by SVGD [46], a deep ensemble with members that interact with each other through a repulsive component. Using a kernel function to model this interaction, the single models repel each other based on their position in the weight space, so that two members can never assume the same weights. Considering a stationary kernel  $k(\cdot,\cdot):\mathbb{R}^d\times\mathbb{R}^d\to\mathbb{R}$  acting in the parameter space of the neural networks, a repulsive term can be parameterized through its gradient:

$$\phi(\mathbf{w}_i^t) = \nabla_{\mathbf{w}_i^t} \log p(\mathbf{w}_i^t | \mathcal{D}) - \mathcal{R}\left(\left\{\nabla_{\mathbf{w}_i^t} k(\mathbf{w}_i^t, \mathbf{w}_j^t)\right\}_{j=1}^n\right). \tag{3}$$

To get an intuition for the behavior of this repulsive term and its gradients, we can consider the RBF kernel  $k(\mathbf{w}, \mathbf{w}') = \exp\left(-\frac{1}{h}||\mathbf{w}_i - \mathbf{w}_i||^2\right)$  with lengthscale h and notice how its gradient

$$\nabla_{\mathbf{w}_i^t} k(\mathbf{w}_i^t, \mathbf{w}_j^t) = \frac{\sum_{j=1}^n \frac{2}{h} (\mathbf{w}_i^t - \mathbf{w}_j^t) k(\mathbf{w}_i^t, \mathbf{w}_j^t)}{\sum_{j=1}^n k(\mathbf{w}_i^t, \mathbf{w}_j^t)}$$

drives  $\mathbf{w}_i$  away from its neighboring members  $\mathbf{w}_i$ , thus creating a repulsive effect.

### 2.2 Repulsive force in function space

To overcome the aforementioned overparameterization issue, the update in Eq. (3) can be formulated in function space instead of weight space. Let  $f: \mathbf{w} \mapsto f(\cdot; \mathbf{w})$  be the map that maps a configuration of weights  $\mathbf{w} \in \mathbb{R}^d$  to the corresponding neural network regression function and denote  $f_i := f(\cdot; \mathbf{w}_i)$  the function with a certain configuration of weights  $\mathbf{w}_i$ . We can now consider n particles in function space  $\{f_i\}_{i=1}^n$  with  $f \in \mathcal{F}$  and model their interaction  $\mathbf{w}$  with a general positive definite kernel  $k(\cdot, \cdot)$ . We also consider the implicit functional likelihood  $p(\mathbf{y}|\mathbf{x}, f)$ , determined by the measure  $p(\mathbf{y}|\mathbf{x}, \mathbf{w})$  in the weight space, as well as the functional prior p(f), which can either be defined separately (e.g., using a GP) or modeled as a push-forward measure of the weight-space prior  $p(\mathbf{w})$ . Together, they determine the posterior in function space  $p(f|\mathcal{D})$ . The functional evolution of a particle can then be written as:

$$\mathbf{f}_{i}^{t+1} \leftarrow \mathbf{f}_{i}^{t} + \epsilon_{t} \phi(\mathbf{f}_{i}^{t})$$
with 
$$\phi(\mathbf{f}_{i}^{t}) = \nabla_{\mathbf{f}_{i}^{t}} \log p(\mathbf{f}_{i}^{t}|\mathcal{D}) - \mathcal{R}\left(\left\{\nabla_{\mathbf{f}_{i}^{t}} k(\mathbf{f}_{i}^{t}, \mathbf{f}_{j}^{t})\right\}_{j=1}^{n}\right). \tag{4}$$

However, computing the update in function space is neither tractable nor practical, which is why two additional considerations are needed. The first one regards the infinite dimensionality of function space, which we circumvent using a canonical projection into a subspace:

**Definition 1** (Canonical projection). For any  $A \subset \mathcal{X}$ , let  $\pi_A : \mathbb{R}^{\hat{\mathcal{X}}} \to \mathbb{R}^A$  denote the canonical projection onto A, that is,  $\pi_A(f) = \{f(a)\}_{a \in A}$ .

In other words, the kernel will not be evaluated directly in function space, but on the projection  $k(\pi_B(f), \pi_B(f'))$ , with B being a subset of the input space given by a batch of training datapoints. The second consideration is to project this update back into the parameter space and evolve a set of particles there, because ultimately we are interested in representing the functions by parametrized neural networks. For this purpose, we can use the Jacobian of the i-th particle as a projector:

$$\phi(\mathbf{w}_{i}^{t}) = \left(\frac{\partial \mathbf{f}_{i}^{t}}{\partial \mathbf{w}_{i}^{t}}\right)^{\top} \left[ \nabla_{\mathbf{f}_{i}^{t}} \log p(\mathbf{f}_{i}^{t}|\mathcal{D}) - \mathcal{R}\left( \left\{ \nabla_{\mathbf{f}_{i}^{t}} k(\pi_{B}(\mathbf{f}_{i}^{t}), \pi_{B}(\mathbf{f}_{j}^{t})) \right\}_{j=1}^{n} \right) \right].$$
 (5)

### 2.3 Comparison to Stein variational gradient descent

Note that our update is reminiscent of SVGD [46], which in parameter space can be written as:

$$\phi(\mathbf{w}_i^t) = \sum_{j=1}^n k(\mathbf{w}_i^t, \mathbf{w}_j^t) \nabla_{\mathbf{w}_i^t} \log p(\mathbf{w}_i^t | \mathcal{D}) + \sum_{j=1}^n \nabla_{\mathbf{w}_j^t} k(\mathbf{w}_j^t, \mathbf{w}_i^t).$$
 (6)

It is important to notice that here, the gradients are averaged across all the particles using the kernel matrix. Interestingly, SVGD can be asymptotically interpreted as gradient flow of the KL divergence under a new metric induced by Stein operator [13, 45] (see Appendix D for more details). Moving the inference from parameter to function space [66], leads to the update rule

$$\phi(\mathbf{w}_i^t) = \left(\frac{\partial f_i^t}{\partial \mathbf{w}_i^t}\right)^{\top} \left(\frac{1}{n} \sum_{j=1}^n k(f_i^t, f_j^t) \nabla_{f_j^t} \log p(f_j^t | \mathcal{D}) + \nabla_{f_j^t} k(f_i^t, f_j^t)\right). \tag{7}$$

This way of averaging gradients using a kernel can be dangerous in high-dimensional settings, where kernel methods often suffer from the curse of dimensionality. Moreover, in Eq. (6), the posterior gradients of the particles are averaged using their similarity in weight space, which can be misleading in multi-modal posteriors. Worse yet, in Eq. (7), the gradients are averaged in function space and are then projected back using exclusively the *i*-th Jacobian, which can be harmful given that it is not guaranteed that distances between functions evaluated on a subset of their input space resemble their true distance. Our proposed method, on the other hand, does not employ any averaging of the posterior gradients and thus comes closest to the true particle gradients in deep ensembles.

# 3 Repulsive deep ensembles are Bayesian

So far, we represented the repulsive force as a general function of the gradients of a kernel. In this section, we show how to determine the explicit form of the repulsive term, such that the resulting

update rule is equivalent to the discretization of the gradient flow dynamics of the KL divergence in Wasserstein space. We begin by introducing the concepts of particle approximation and gradient flow.

### 3.1 Particle approximation

A particle-based approximation of a target measure depends on a set of weighted samples  $\{(x_i, w_i)\}_{i=1}^n$ , for which an empirical measure can be defined as

$$\rho(x) = \sum_{i=1}^{n} w_i \, \delta(x - x_i) \,, \tag{8}$$

where  $\delta(\cdot)$  is the Dirac delta function and the weights  $w_i$  satisfy  $w_i \in [0,1]$  and  $\sum_{i=1}^n w_i = 1$ . To approximate a target distribution  $\pi(x)$  using the empirical measure, the particles and their weights need to be selected in a principled manner that minimizes some measure of distance between  $\pi(x)$  and  $\rho(x)$  (e.g., a set of N samples with weights  $w_i = 1/N$  obtained using an MCMC method).

### 3.2 Gradient flow in parameter space

Given a smooth function  $J: \mathbb{R}^d \to \mathbb{R}$  in Euclidean space, we can minimize it by creating a path that follows its negative gradient starting from some initial conditions  $x_0$ . The curve x(t) with starting point  $x_0$  described by that path is called *gradient flow*. The dynamics and evolution in time of a considered point in the space under this minimization problem can be described as the ODE<sup>1</sup>

$$\frac{dx}{dt} = -\nabla J(x) \,. \tag{9}$$

Analogously, we can consider a Riemannian manifold  $\mathcal{M}$  equipped with the metric tensor G(x) defined for all  $x \in \mathcal{M}$ . Here,  $G(x): \mathcal{T}_{\mathcal{W}} \times \mathcal{T}_{\mathcal{W}} \to \mathbb{R}$  defines a smoothly varying local inner product on the tangent space at each point of the manifold x. For manifolds over  $\mathbb{R}^d$ , the metric tensor is a positive definite matrix that defines local distances for infinitesimal displacements  $d(x, x + dx) = \sqrt{dx^{\top}G(x)dx}$ . Considering a functional  $J: \mathcal{M} \to \mathbb{R}$ , the evolution in Eq. (9) becomes

$$\frac{dx}{dt} = -G(x)^{-1}\nabla J(x) . {10}$$

We see that the metric tensor of the manifold acts like a perturbation of the gradient. We can now extend this concept to the space of probability distributions (*Wasserstein gradient flow*) [2]. Let us consider the space of probability measures  $\mathcal{P}_2(\mathcal{M})$ , that is the set of probability measures with finite second moments defined on the manifold  $\mathcal{M}$ :

$$\mathcal{P}_2(\mathcal{M}) = \left\{ \rho : \mathcal{M} \to [0, \infty) \middle| \int_{\mathcal{M}} d\rho = 1, \int_{\mathcal{M}} |x|^2 \rho(x) dx < +\infty \right\}.$$

Taking  $\Pi(\mu, \nu)$  as the set of joint probability measures with marginals  $\mu, \nu$ , we can define the Wasserstein metric on the space  $\mathcal{P}_2(\mathcal{M})$  as:

$$W_2^2(\mu,\nu) = \inf_{\pi \in \Pi(\mu,\nu)} \int |x-y|^2 d\pi(x,y).$$
 (11)

Considering the optimization problem of a functional  $J: \mathcal{P}_2(\mathcal{M}) \to \mathbb{R}$ , such as the KL divergence between the particle approximation in Eq. (8) and the target posterior  $\pi(x)$ ,

$$\inf_{\rho \in \mathcal{P}_2(\mathcal{M})} D_{KL}(\rho, \pi) = \int_{\mathcal{M}} (\log \rho(x) - \log \pi(x)) \rho(x) \, \mathrm{d}x \;,$$

the evolution in time of the measure  $\rho$  under the equivalent of the gradient, the Wasserstein gradient flow, is described by the *Liouville equation*:

$$\frac{\partial \rho(x)}{\partial t} = \nabla \cdot \left( \rho(x) \nabla \frac{\delta}{\delta \rho} D_{KL}(\rho, \pi) \right) 
= \nabla \cdot \left( \rho(x) \nabla \left( \log \rho(x) - \log \pi(x) \right) \right),$$
(12)

<sup>&</sup>lt;sup>1</sup>Together with the initial condition, this is know as the Cauchy problem.

where  $\nabla \frac{\delta}{\delta \rho} D_{KL}(\rho, \pi) =: \nabla_{\mathcal{W}_2} D_{KL}(\rho, \pi)$  is the Wasserstein gradient and the operator  $\frac{\delta}{\delta \rho}$ :  $\mathcal{P}_2(\mathcal{M}) \to \mathbb{R}$  represents the functional derivative or first variation (see Appendix C for more details). In the particular case of the KL functional, we can recover the Fokker-Planck equation,

$$\begin{split} \frac{\partial \rho(x)}{\partial t} &= \nabla \cdot \left( \rho(x) \nabla (\log \rho(x) - \log \pi(x)) \right) \\ &= -\nabla \cdot \left( \rho(x) \nabla \log \pi(x) \right) + \nabla^2 \rho(x) \end{split}$$

that admits as unique stationary distribution the posterior  $\pi(x)$ . The deterministic particle dynamics ODE [1] related to Eq. (12), namely mean-field Wasserstein dynamics, is then given by:

$$\frac{dx}{dt} = -\nabla \left(\log \rho(x) - \log \pi(x)\right). \tag{13}$$

Considering a discretization of Eq. (13) for a particle system  $\{x\}_{i=1}^n$  and small stepsize  $\epsilon_t$ , we can rewrite Eq. (13) as:

$$x_i^{t+1} = x_i^t + \epsilon_t \left( \nabla \log \pi(x_i^t) - \nabla \log \rho(x_i^t) \right). \tag{14}$$

Unfortunately, we do not have access to the analytical form of the gradient  $\nabla \log \rho$ , so an approximation is needed. At this point, it is crucial to observe the similarity between the discretization of the Wasserstein gradient flow in Eq. (14) and the repulsive update in Eq. (3) to notice how, if the kernelized repulsion is an approximation of the gradient of the empirical particle measure, the update rule minimizes the KL divergence between the particle measure and the target posterior. Different sample-based approximations of the gradient that use a kernel function have been recently studied. The simplest one is given by the kernel density estimation (KDE) (details in Appendix E)  $\tilde{\rho}_t(x) = \sum_{i=1}^n k(x, x_t^i)$ , where  $k(\cdot, \cdot) : \mathbb{R}^d \times \mathbb{R}^d \to \mathbb{R}$  and the gradient of its log density is given by [61]:

$$\nabla \log \rho(x_i^t) \approx \frac{\sum_{j=1}^n \nabla_{x_i^t} k(x_i^t, x_j^t)}{\sum_{j=1}^n k(x_i^t, x_j^t)} \,. \tag{15}$$

Using this approximation in Eq. (14) we obtain

$$x_i^{t+1} = x_i^t + \epsilon_t \left( \nabla \log \pi(x_i^t) - \frac{\sum_{j=1}^n \nabla_{x_i^t} k(x_i^t, x_j^t)}{\sum_{j=1}^n k(x_i^t, x_j^t)} \right), \tag{16}$$

where, if we substitute the posterior for  $\pi$ , we obtain an expression for the repulsive force in Eq. (3). This shows that if the repulsive term is the normalized sum of the gradients  $\mathcal{R}(x_i^t) = \beta(x_i^t) \sum_{j=1}^n \nabla_{x_i^t} k(x_i^t, x_j^t)$ , we do not only encourage diversity of the ensemble members and thus avoid collapse, but surprisingly—in the asymptotic limit of  $n \to \infty$ , where the KDE approximation is exact [58] —also converge to the true Bayesian posterior!

Nevertheless, approximating the gradient of the empirical measure with the KDE can lead to suboptimal performance, as already studied by Li and Turner [42]. They instead introduced a new *Stein gradient estimator* (SGE) that offers better performance, while maintaining the same computational cost. Even more recently, Shi et al. [60] introduced a spectral method for gradient estimation (SSGE), that also allows for a simple estimation on out-of-sample points. These two estimators can be used in Eq. (14), leading to the following update rules. The one using the Stein estimator, that we will call SGE-WGD, is:

$$x_i^{t+1} = x_i^t + \epsilon_t \left( \nabla \log \pi(x_i^t) + \sum_{j=1}^n (K + \eta \mathbb{I})_{ij}^{-1} \sum_{k=1}^n \nabla_{x_k^t} k(x_k^t, x_j^t) \right), \tag{17}$$

where K is the kernel Gram matrix,  $\eta$  a small constant and  $\mathbb{I}$  the identity matrix. We can notice the important difference between KDE and SGE that the former one is only considering the interaction of the i-th particle being updated with all the others, while the latter is simultaneously considering also the interactions between the remaining particles. The spectral method, that we will call SSGE-WGD, leads to the following update rule:

$$x_{t+1}^{i} = x_{t}^{i} + \epsilon_{t} \left( \nabla \log \pi(x_{t}^{i}) + \sum_{j=1}^{J} \frac{1}{\lambda_{j}^{2}} \sum_{m=1}^{n} \sum_{k=1}^{n} u_{jk} \nabla_{x_{m}} k(x_{m}^{t}, x_{k}^{t}) \cdot \sum_{l=1}^{n} u_{jl} k(x_{i}^{t}, x_{l}^{t}) \right)$$
(18)

where  $\lambda_j$  is the j-th eigenvalue of the kernel matrix and  $u_{jk}$  is the k-th component of the j-th eigenvector.

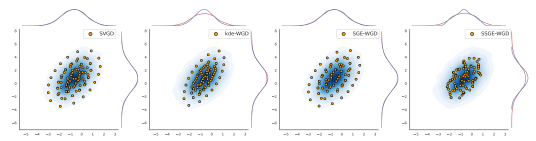


Figure 2: **Single Gaussian.** We show samples from SVGD, KDE-WGD, SGE-WGD, and SSGE-WGD (from left to right). The SGE-WGD and SVGD fit the target almost perfectly.

### 3.3 Gradient flow in function space

To theoretically justify the update rule introduced in function space in Eq. (5), we can rewrite the Liouville equation for the gradient flow in Eq. (12) in function space as

$$\frac{\partial \rho(\mathbf{f})}{\partial t} = \nabla \cdot \left( \rho(\mathbf{f}) \nabla \frac{\delta}{\delta \rho} D_{KL}(\rho, \pi) \right) 
= \nabla \cdot \left( \rho(\mathbf{f}) \nabla \left( \log \rho(\mathbf{f}) - \log \pi(\mathbf{f}) \right) \right).$$
(19)

Following this update, the mean field functional dynamics are

$$\frac{d\mathbf{f}}{dt} = -\nabla \left(\log \rho(\mathbf{f}) - \log \pi(\mathbf{f})\right). \tag{20}$$

Using the same KDE approximation as above, we can obtain a discretized evolution in function space and with it an explicit form for the repulsive force in Eq. (4) as

$$\mathbf{f}_{t+1}^{i} = \mathbf{f}_{t}^{i} + \epsilon_{t} \left( \nabla_{\mathbf{f}} \log \pi(\mathbf{f}_{t}^{i}) - \frac{\sum_{j=1}^{n} \nabla_{\mathbf{f}_{t}^{i}} k(\mathbf{f}_{t}^{i}, \mathbf{f}_{j}^{t})}{\sum_{j=1}^{n} k(\mathbf{f}_{t}^{i}, \mathbf{f}_{j}^{t})} \right). \tag{21}$$

The update rules using the SGE and SSGE approximations follow as for the parametric case.

# 4 Experiments

In this section, we compare the different proposed WGD methods with deep ensembles and SVGD on synthetic sampling, regression, and classification tasks and real-world image classification tasks. We also quantitatively assess the uncertainty estimation of the methods in terms of calibration and OOD detection. In our experiments, we report the test accuracy, negative log-likelihood (NLL), and the expected calibration error (ECE) [53]. To assert the robustness on out-of-distribution (OOD) data, we report the ratio between predictive entropy on OOD and test data points  $(H_o/H_t)$ , and the OOD detection area under the ROC curve AUROC(H) [43]. Moreover, to assess the diversity of the ensemble generated by the different methods in function space, we measure the functional diversity using the model disagreement (MD) (details in Appendix B). In particular, we report the ratio between the average model disagreement on the OOD and test data points  $(MD_o/MD_t)$  and additionally the AUROC(MD) computed using this measure instead of the entropy.

**Sampling from synthetic distributions** As a sanity check, we first assessed the ability of our different approximations for Wasserstein gradient descent (using KDE, SGE, and SSGE) to sample from a two-dimensional univariate Gaussian distribution (Figure 2). We see that our SGE-WGD and the SVGD fit the target almost perfectly. We also tested the different methods in a more complex two-dimensional Funnel distribution [54] and present the results in Figure F.1 in the Appendix. There, SGE-WGD and SVGD also perform best.

**BNN 1D regression** We then assessed the different methods in fitting a BNN posterior on a synthetically generated one-dimensional regression task. The results are reported in Figure 1, consisting of the mean prediction and  $\pm 1, 2, 3$  standard deviations of the predictive distribution. We

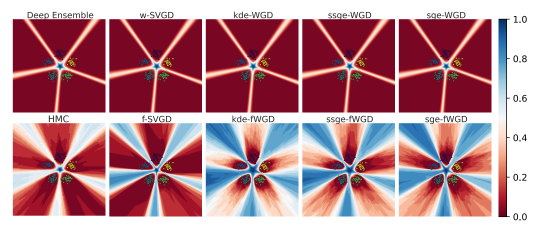


Figure 3: **BNN 2D classification.** Entropy of the predictive posteriors. Again, the function-space methods capture the epistemic uncertainty better than the weight-space ones, thus approaching the gold-standard HMC posterior.

can see that all methods performing inference in the weight space (DE, w-SVGD, WGD) are unable to capture the epistemic uncertainty between the two clusters of training data points. Conversely, the functional methods (f-SVGD, fWGD) are perfectly able to infer the diversity of the hypotheses in this region due to the lack of training evidence. They thereby achieve a predictive posterior that very closely resembles the one obtained with the gold-standard HMC sampling.

BNN 2D classification Next, we investigated the predictive performance and quality of uncertainty estimation of the methods in a two-dimensional synthetic classification setting. The results are displayed in Figure 3. We can clearly observe that the weight-space methods are overconfident and do not capture the uncertainty well. Moreover, all the functions seems to collapse to the optimal classifier. These methods thus only account for uncertainty close to the decision boundaries and the origin region, for which the uncertainty is purely aleatoric. In this setting, f-SVGD suffers for a similar issues as the weight space methods, being overconfident away from the training data. Conversely, our fWGD methods are confident (low entropy) around the data but not out-of-distribution, thus representing the epistemic uncertainty better. This suggests that the functional diversity captured by this method naturally leads to a *distance-aware* uncertainty estimation, a property that translates into confident predictions only in the proximity of the training data, allowing for a principled OOD detection.

FashionMNIST classification In this section, the methods are tested in an image classification setting using the FashionMNIST dataset [73] for training and the MNIST dataset [39] as an out-of-distribution (OOD) task. The results are reported in Table 1 (top). We can see that all our methods improve upon standard deep ensembles and SVGD, highlighting the effectiveness of our proposed repulsion terms when training neural network ensembles. In particular, the sge-WGD offers the best accuracy, whereas the methods in function space all offer a better OOD detection. This is probably due to the fact that these methods achieve an higher entropy ratio and functional diversity measured via the model disagreement when compared to their weight space counterparts. Interestingly, they also reach the lowest NLL values. We can also notice how the model disagreement (MD) not only serves its purpose as a metric for the functional heterogeneity of the ensemble but also allows for a better OOD detection in comparison to the entropy. To the best of our knowledge, this insight has not been described before but similarly used in continual learning scenarios Henning et al. [27].

**CIFAR classification** In this experiment, we use a ResNet32 architecture [25] on CIFAR-10 [37] with the SVHN dataset [56] as OOD data. The results are reported in Table 1 (bottom). We can see that in this case the weight space methods achieve better performance in accuracy and OOD detection using the entropy than the ones in function space. Nevertheless, all our repulsive ensembles improve functional diversity, accuracy, and OOD detection when compared to standard SVGD whereas the standard deep ensemble achieves the best OOD detection using the entropy.

Table 1: **BNN image classification. AUROC(H)** is the AUROC computed using the entropy whereas **AUROC(MD)** is computed using the model disagreement.  $\mathbf{H_o/H_t}$  is the ratio of the entropies on OOD and test points respectively and  $\mathbf{MD_o/MD_t}$  is the ratio for model disagreement. We see that the best accuracy is achieved by our WGD methods, while our fWGD methods yield the best OOD detection and funtional diversity. All our proposed methods improve over standard deep ensembles in terms of accuracy and diversity, highlighting the effect of our repulsion.

		AUROC(H)	AUROC(MD)	Accuracy	$ m H_o/H_t$	$\mathrm{MD_o/MD_t}$	ECE	NLL
FashionMNIST	Deep ensemble [38]	0.958±0.001	0.975±0.001	91.122±0.013	6.257±0.005	6.394±0.001	$0.012 \pm 0.001$	0.129±0.001
	SVGD [46]	$0.960\pm0.001$	$0.973\pm0.001$	$91.134 \pm 0.024$	$6.315\pm0.019$	$6.395 \pm 0.018$	$0.014\pm0.001$	$0.127\pm0.001$
	f-SVGD [66]	$0.956\pm0.001$	$0.975\pm0.001$	$89.884 \pm 0.015$	$5.652\pm0.009$	$6.531 \pm 0.005$	$0.013\pm0.001$	$0.150\pm0.001$
	kde-WGD (ours)	$0.960\pm0.001$	$0.970\pm0.001$	$91.238 \pm 0.019$	$6.587 \pm 0.019$	$6.379\pm0.018$	$0.014\pm0.001$	$0.128\pm0.001$
	sge-WGD (ours)	$0.960\pm0.001$	$0.970\pm0.001$	$91.312 \pm 0.016$	$6.562 \pm 0.007$	$6.363 \pm 0.009$	$0.012 \pm 0.001$	$0.128\pm0.001$
	ssge-WGD (ours)	$0.968\pm0.001$	$0.979\pm0.001$	$91.198 \pm 0.024$	$6.522 \pm 0.009$	$6.610\pm0.012$	$0.012 \pm 0.001$	$0.130\pm0.001$
	kde-fWGD (ours)	$0.971\pm0.001$	$0.980 \pm 0.001$	$91.260 \pm 0.011$	$7.079\pm0.016$	$6.887 \pm 0.015$	$0.015\pm0.001$	$0.125 \pm 0.001$
	sge-fWGD (ours)	$0.969\pm0.001$	$0.978\pm0.001$	$91.192\pm0.013$	$7.076\pm0.004$	$6.900\pm0.005$	$0.015\pm0.001$	$0.125 \pm 0.001$
	ssge-fWGD (ours)	$0.971 \pm 0.001$	$0.980 \pm 0.001$	$91.240 \pm 0.022$	$7.129 \pm 0.006$	$6.951 \pm 0.005$	$0.016\pm0.001$	$0.124 \pm 0.001$
CIFAR10	Deep ensemble [38]	$0.843 {\pm} 0.004$	$0.736\pm0.005$	85.552±0.076	2.244±0.006	1.667±0.008	$0.049\pm0.001$	0.277±0.001
	SVGD [46]	$0.825 \pm 0.001$	$0.710\pm0.002$	$85.142 \pm 0.017$	$2.106\pm0.003$	$1.567 \pm 0.004$	$0.052\pm0.001$	$0.287 \pm 0.001$
	fSVGD [66]	$0.783\pm0.001$	$0.712\pm0.001$	$84.510\pm0.031$	$1.968\pm0.004$	$1.624\pm0.003$	$0.049\pm0.001$	$0.292\pm0.001$
	kde-WGD (ours)	$0.838\pm0.001$	$0.735\pm0.004$	$85.904\pm0.030$	$2.205\pm0.003$	$1.661\pm0.008$	$0.053\pm0.001$	$0.276\pm0.001$
	sge-WGD (ours)	$0.837\pm0.003$	$0.725\pm0.004$	$85.792 \pm 0.035$	$2.214\pm0.010$	$1.634\pm0.004$	$0.051\pm0.001$	$0.275\pm0.001$
	ssge-WGD (ours)	$0.832\pm0.003$	$0.731\pm0.005$	$85.638 \pm 0.038$	$2.182\pm0.015$	$1.655\pm0.001$	$0.049\pm0.001$	$0.276\pm0.001$
	kde-fWGD (ours)	$0.791\pm0.002$	$0.758 \pm 0.002$	$84.888 \pm 0.030$	$1.970\pm0.004$	$1.749 \pm 0.005$	$0.044 \pm 0.001$	$0.282 \pm 0.001$
	sge-fWGD (ours)	$0.795\pm0.001$	$0.754\pm0.002$	$84.766\pm0.060$	$1.984 \pm 0.003$	$1.729\pm0.002$	$0.047\pm0.001$	$0.288 \pm 0.001$
	ssge-fWGD (ours)	$0.792 \pm 0.002$	$0.752 \pm 0.002$	$84.762 \pm 0.034$	$1.970 \pm 0.006$	$1.723 \pm 0.005$	$0.046 \pm 0.001$	$0.286{\pm}0.001$

# 5 Related Work

The theoretical and empirical properties of SVGD have been well studied [36, 44, 12] and it can also be seen as a Wasserstein gradient flow of the KL divergence in the Stein geometry [13, 45] (see Appendix D for more details). Interestingly, a gradient flow interpretation is also possible for (stochastic gradient) MCMC-type algorithms [44], which can be unified under a general particle inference framework [9]. Moreover, our Wasserstein gradient descent using the SGE approximation can also be derived using an alternative formulation as a gradient flow with smoothed test functions [44]. A projected version of WGD has been studied in [65], which could also be readily applied in our framework. Besides particle methods, Bayesian neural networks MacKay [49], Neal [54] have gained popularity recently [69, 18, 16, 32], using modern MCMC [54, 69, 18, 20, 17] and variational inference techniques [4, 63, 14, 30]. Ensemble methods, on the other hand, have also been extensively studied [38, 15, 72, 19] and many variants have been proposed [70, 23, 28, 75, 68]. Moreover, providing Bayesian interpretations for deep ensembles has been previously attempted through the lenses of stationary SGD distributions [51, 7], ensembles of linear models [52], additional random functions [11, 23], approximate inference [72], and marginal likelihood lower bounds [48]. Furthermore, variational inference in function space has recently gained attention [62] and the limitations of the KL divergence have been studied in [6].

# 6 Conclusion

We have presented a simple and principled way to improve upon standard deep ensemble methods. To this end, we have shown that the introduction of a kernelized repulsion between members of the ensemble not only improves the accuracy of the predictions but—even more importantly—can be seen as Wasserstein gradient descent on the KL divergence, thus transforming the MAP inference of deep ensembles into proper Bayesian inference. Moreover, we have shown that incorporating functional repulsion between ensemble members can improve the quality of the estimated uncertainties on simple synthetic examples and OOD detection on real-world data and can approach the true Bayesian posterior more closely.

In future work, it will be interesting to study the impact of the Jacobian in the fWGD update and its implications on the Liouville equation in more detail, also compared to other neural network Jacobian methods, such as neural tangent kernels [33] and generalized Gauss-Newton approximations [31]. Moreover, it would be interesting to derive explicit convergence bounds for our proposed method and compare them to the existing bounds for SVGD [36].

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# Supplementary Material: Repulsive Deep Ensembles are Bayesian

### A Non-identifiable neural networks

Deep neural networks are capable of successfully learning from examples in a wide variety of tasks. Although these networks are typically trained on large amounts of data, the number of free parameters in their architectures is often several orders of magnitude greater than the number of training examples. This overparametrization reflects the ability of DNNs to memorize entire datasets, even with randomized labels [74]. Moreover, this overparametrized regime makes the neural network approximation of a given function not unique in the sense that multiple configurations of weights might lead to the same function. Indeed, the output of a feedforward neural network given some fixed input remains unchanged under a set of transformations. For instance, certain weight permutations and sign flips in MLPs leave the output unchanged [8]. If a certain transformation of the weights leads to the same prediction and thus to the same function, then also the likelihood of the model is invariant under the same transformation. Such neural networks are also referred to as *non-identifiable*. Specifically, for any non-identifiable pair  $\{\mathbf{w}_i, \mathbf{w}_j\} \in \mathcal{W} \subseteq \mathbb{R}^d$  and  $\mathbf{f}_i, \mathbf{f}_j \in \mathcal{F}$  their respective functions:

$$f_i = f_j \implies p(\mathbf{w}_i | \mathcal{D}) = p(\mathbf{w}_j | \mathcal{D}) \implies \mathbf{w}_i = \mathbf{w}_j$$

In other words, the map  $g: \mathcal{X} \times \mathcal{W} \to \mathcal{F}$  is not injective (many to one). If we now call T the group of transformations on the parameter space that leaves the output unchanged, it can be shown that for each permutable feedforward neural network, there exists a cone  $K \subset \mathbb{R}^d$  such that for any parameter configuration  $\mathbf{w}$ , there exist  $\eta \in K$  and a transformation  $\tau \in T$  such that  $\tau \eta = \mathbf{w}$ . This means that every parameter configuration has an equivalent on the subset given by the cone [26].

Modern neural networks containing convolutional and max-pooling layers have even more symmetries than MLPs [3]. Given that in practice, we cannot constrain the support of the posterior distribution to be the cone of identifiable parameter configurations and given that the likelihood model is also invariant under those transformations that do not change the function, the posterior landscape includes multiple equally likely modes that—despite their different positions—represent the same function. It is important to notice that this is always true for the likelihood but not for the posterior. Indeed, for the modes to be equally likely, the prior should also be invariant under those transformations, a condition that is not true in general. Nevertheless, the fact that there are multiple modes of the posterior parametrizing for the same function remains true, but they might be arbitrarily re-scaled by the prior<sup>2</sup>. This redundancy of the posterior is problematic when we want to obtain samples from it. Moreover it is interesting to notice how this issue disappears when the Bayesian inference is considered in the space of functions instead of weights. In this case, indeed, every optimal function has a unique mode in the landscape of the posterior and redundancy is not present:

$$f_i \neq f_i \implies p(f_i|\mathcal{D}) \neq p(f_i|\mathcal{D})$$
.

In spite of that, performing inference over distributions of functions is prohibitive in practice due to the infinite dimensionality of the space in consideration. Only in very limited cases like the one of Gaussian process, Bayesian inference is exact. Interestingly, neural network models in the limit of infinite width are Gaussian processes with a particular choice of the kernel determined by the architecture [40, 71, 55]. In this limit, Bayesian inference over functions can be performed analytically.

# **B** Quantify functional diversity

As illustrated in Section 2, in the Bayesian context, predictions are made by doing a Monte-Carlo estimation of the BMA. Functional diversity, and so the diversity in the hypotheses taken in consideration when performing the estimation, determines the epistemic uncertainty and the confidence over the predictions. Importantly, the epistemic uncertainty allows for the quantification

<sup>&</sup>lt;sup>2</sup>Note that for the fully factorized Gaussian prior commonly adopted, the invariance under permutations is true.

of the likelihood of a test point to belong to the same distribution from which the training data points were sampled [57]. Following this, the uncertainty can be used for the problem of Out-of-distribution (OOD) detection [10] that is often linked to the ability of a model to "know what it doesn't know". A common way used in the literature to quantify the uncertainty is the Entropy<sup>3</sup>  $\mathcal{H}$  of the predictive distribution:

$$\mathcal{H}\left\{p(\mathbf{y}'|\mathbf{x}',\mathcal{D})\right\} = -\sum_{y} p(\mathbf{y}'|\mathbf{x}',\mathcal{D}) \log p(\mathbf{y}'|\mathbf{x}',\mathcal{D}). \tag{22}$$

Nevertheless, it has been argued in recent works [50] that this is not a good measure of uncertainty because it does not allow for a disentanglement of epistemic and aletoric uncertainty. Intuitively, we would like the predictive distribution of an OOD point to be uniform over the different classes. However, using the entropy and so the average prediction in the BMA, we are not able to distinguish between the case in which all the hypotheses disagree very confidently due to the epistemic uncertainty or are equally not confident due to the aleatoric uncertainty. To overcome this limitation, we can use a direct measure of the model disagreement computed as:

$$\mathcal{MD}^{2}(\mathbf{y}'; \mathbf{x}', \mathcal{D}) = \int_{\mathbf{w}} \left[ p(\mathbf{y}'|\mathbf{x}', \mathbf{w}) - p(\mathbf{y}'|\mathbf{x}', \mathcal{D}) \right]^{2} p(\mathbf{w}|\mathcal{D}) d\mathbf{w}.$$
 (23)

It is easy to see how the quantity in Eq. (23), measuring the deviation from the average prediction is zero when all models agree on the prediction. The latter can be the case of a training point where all hypotheses are confident or a noisy point where all models "don't know" the class and are equally uncertain. On the other side the model disagreement will be greater the zero the more the model disagree on a prediction representing like this the epistemic uncertainty. To obtain a scalar quantity out of Eq. (23) we can consider the expectation over the output space of y:

$$\mathcal{MD}^{2}(\mathbf{x}') = \mathbb{E}_{y} \left[ \int_{\mathbf{w}} \left[ p(\mathbf{y}'|\mathbf{x}', \mathbf{w}) - p(\mathbf{y}'|\mathbf{x}', \mathcal{D}) \right]^{2} p(\mathbf{w}|\mathcal{D}) d\mathbf{w} \right].$$
 (24)

# C Functional derivative of the KL divergence

In this section, we show the derivation of the functional derivative for the KL divergence functional. We start with some preliminary definitions.

Given a manifold  $\mathcal{M}$  embedded in  $\mathbb{R}^d$ , let  $F[\rho]$  be a functional, i.e. a mapping from a normed linear space of function (Banach space)  $\mathcal{F} = \{\rho(x) : x \in \mathcal{M}\}$  to the field of real numbers  $F : \mathcal{F} \to \mathbb{R}$ . The functional derivative  $\delta F[\rho]/\delta \rho(x)$  represents the variation of value of the functional if the function  $\rho(x)$  is changed.

**Definition 2** (Functional derivative). *Given a manifold*  $\mathcal{M}$  *and a functional*  $F : \mathcal{F} \to \mathbb{R}$  *with respect to*  $\rho$  *is defined as:* 

$$\int \frac{\delta F}{\delta \rho(x)} \phi(x) dx = \lim_{\epsilon \to 0} \frac{F[\rho(x) + \epsilon \phi(x)] - F(\rho(x))}{\epsilon} = \frac{d}{d\epsilon} F[\rho(x) + \epsilon \phi(x)] \bigg|_{\epsilon = 0}$$
(25)

for every smooth  $\phi$ .

**Definition 3** (KL divergence). *Given*  $\rho$  *and*  $\pi$  *two probability densities on*  $\mathcal{M}$ , *the KL-divergence is defined as:* 

$$D_{KL}(\rho, \pi) = \int_{\mathcal{M}} \left( \log \rho(x) - \log \pi(x) \right) \rho(x) \, dx \,. \tag{26}$$

**Proposition 1.** The functional derivative of the KL divergence in Eq. (26) is:

$$\frac{\delta D_{KL}}{\delta \rho(x)} = \log \frac{\rho(x)}{\pi(x)} + 1 \tag{27}$$

<sup>&</sup>lt;sup>3</sup>The continuous case is analogous using the differential entropy

*Proof.* using the definition of functional derivative in Eq. (25):

$$\int \frac{\delta D_{KL}}{\delta \rho(x)} \phi(x) dx = \frac{d}{d\epsilon} D_{KL}(\rho + \epsilon \phi, \pi) \Big|_{\epsilon=0}$$

$$= \int \frac{d}{d\epsilon} \left[ (\rho(x) + \epsilon \phi(x)) \log \frac{(\rho(x) + \epsilon \phi(x))}{\pi(x)} \right]_{\epsilon=0} dx$$

$$= \int \left[ \phi(x) \log \frac{(\rho(x) + \epsilon \phi(x))}{\pi(x)} + \frac{d(\rho(x) + \epsilon \phi(x))}{d\epsilon} \right]_{\epsilon=0} dx$$

$$= \int \left[ \log \frac{\rho(x)}{\pi(x)} + 1 \right] \phi(x) dx$$
(28)

# D SVGD as Wasserstein gradient flow

To understand the connection between the Wasserstein gradient flow and the SVGD method, the update equation in 6 needs to be reformulated in the continuous time, as the following ODE:

$$\frac{dx_i}{dt} = \frac{1}{n} \sum_{j=1}^{n} [k(x_j, x_i) \nabla_{x_j} \log p(x_j) + \nabla_{x_j} k(x_j, x_i)]$$
 (29)

that in the mean field limit becomes:

$$\frac{dx}{dt} = \int \left[ k(x', x) \nabla_{x'} \log \pi(x') + \nabla_{x'} k(x', x) \right] \rho(x') dx'$$

$$= \int k(x', x) \nabla_{x'} \log \pi(x') \rho(x') dx' + \int \nabla_{x'} k(x', x) \rho(x') dx'.$$
(30)

Due to the boundary condition of a Kernel in the Stein class (see Liu et al. [47] for more details), without loss of generality, we can rewrite the integrals in the previous equation as:

$$\frac{dx}{dt} = \int k(x', x) \nabla_{x'} \log \pi(x') \rho(x') dx' + \int \nabla_{x'} k(x', x) \rho(x') dx' - \underbrace{k(x', x) \rho(x')}_{||x'|| \to \infty}$$

and notice how the second and the third terms are the result of an integration by parts:

$$\frac{dx}{dt} = \int k(x', x) \nabla_{x'} \log \pi(x') \rho(x') dx' - \int \nabla_{x'} k(x', x) \nabla_{x'} \rho(x') dx'$$

$$= \int k(x', x) \nabla_{x'} \log \pi(x') \rho(x') dx' - \int \nabla_{x'} k(x', x) \nabla_{x'} \log \rho(x') \rho(x') dx'$$

$$= \int k(x', x) \nabla_{x'} \left[ \log \pi(x') - \log \rho(x') \right] \rho(x') dx'$$

$$= \mathbb{E}_{x' \sim \rho} \left[ k(x', x) \nabla_{x'} (\log \pi(x') - \log \rho(x')) \right]$$
(31)

which is exactly the functional derivative of the KL divergence in 12 approximated in the RKHS of the kernel [13, 45]. Following this, the Liouville equation of the SVGD dynamics in the mean field limit is:

$$\frac{\partial \rho(x)}{\partial t} = -\nabla \cdot \left( \int k(x', x) \nabla_{x'} \left[ \log \pi(x') - \log \rho(x') \right] \rho(x') dx' \right) 
= \nabla \cdot \left( \int k(x', x) \nabla_{x'} \frac{\delta}{\delta \rho} D_{KL}(\rho, \pi) \rho(x') dx' \right).$$
(32)

Defining the linear operator  $(\mathcal{K}_{\rho}\phi)(x) := \mathbb{E}_{x'\sim\rho}[k(x',x)\phi(x')]$  the ODE in Eq. (31) becomes:

$$\frac{dx}{dt} = \mathcal{K}_{\rho} \nabla_{x'} \left( \log \pi(x') - \log \rho(x') \right). \tag{33}$$

And the Liouville equation describing the evolution of the empirical measure of the particles:

$$\frac{\partial \rho}{\partial t} = \nabla \cdot \left( \rho(x) \mathcal{K}_{\rho} \nabla_{x'} (\log \rho(x') - \log \pi(x')) \right) 
= \nabla \cdot \left( \rho(x) \mathcal{K}_{\rho} \nabla_{x'} \frac{\delta}{\delta \rho} D_{KL}(\rho, \pi) \right).$$
(34)

Notice how the only solution of the previous equation is  $\rho = \pi$ . If we now compare Eq. (34) and Eq. (12) we can see how they only differ for the application of the operator  $\mathcal{K}_{\rho}$ ; moreover the operator seems to act like a perturbation of the gradient and defines a local geometry in the same way the Riemannian metric tensor G(x) is doing in 10. The intuition suggest then that the SVGD can be reinterpreted as a gradient flow in the Wasserstein space under a particular *Stein geometry* [13] induced on the manifold by the kernel and where  $\mathcal{K}_{\rho}$  is the metric.

# **E** Kernel density estimation

Kernel Density Estimation (KDE) is a nonparametric density estimation technique [58]. When an RBF kernel is used, it can be thought as a smoothed version of the empirical data distribution. Given some training datapoints  $\mathcal{D} = \{\mathbf{x}_1, ..., \mathbf{x}_n\}$  with  $\mathbf{x}_i \sim p(\mathbf{x})$  and  $\mathbf{x} \in \mathbb{R}^D$  their empirical distribution  $q_0(\mathbf{x})$  is a mixture of n Dirac deltas centered at each training data:

$$q_0(\mathbf{x}) = \frac{1}{N} \sum_{i=1}^{N} \delta(\mathbf{x} - \mathbf{x}_i).$$
 (35)

We can now smooth the latter by replacing each delta with an RBF kernel:

$$k_{\epsilon}(\mathbf{x}, \mathbf{x}_i) = \frac{1}{\epsilon} \exp\left(\frac{||\mathbf{x} - \mathbf{x}_i||^2}{\epsilon}\right)$$
 (36)

where  $\epsilon > 0$ . The kernel density estimator is then defined as:

$$q_{\epsilon}(\mathbf{x}) = \frac{1}{N} \sum_{i=1}^{N} k_{\epsilon}(\mathbf{x}, \mathbf{x}_i)$$
(37)

In the limit of  $\epsilon \to 0$  and  $N \to \infty$  the kernel density estimator is unbiased: it is equal to the true density. Indeed  $k_{\epsilon \to 0}(\mathbf{x}, \mathbf{x}_i) \to \delta(\mathbf{x} - \mathbf{x}_i)$  and so  $q_{\epsilon \to 0}(\mathbf{x}) \to q_0(\mathbf{x})$  and:

$$\lim_{N \to \infty} q_0(\mathbf{x}) = \lim_{N \to \infty} \frac{1}{N} \sum_{i=1}^{N} (\delta(\mathbf{x} - \mathbf{x}_i))$$

$$= \mathbb{E}_{p(\mathbf{x}')} \left[ \delta(\mathbf{x} - \mathbf{x}') \right]$$

$$= \int_{\mathbb{R}^D} \delta(\mathbf{x} - \mathbf{x}') p(\mathbf{x}') d\mathbf{x}' = p(\mathbf{x})$$
(38)

# F Additional experiments

In this section, we report the additional results for the different methods when sampling from the Funnel distribution  $p(x,y) = \mathcal{N}(y|\mu=0,\sigma=3)\mathcal{N}(x|0,\exp(y/2))$ , the results are reported in Figure F.1.

# **G** Implementation details

In this section, we report details on our implementation in the experiments we performed. The code is partially based on von Oswald et al. [64]. All the experiments were performed on an internal cluster with NVIDIA GTX 1080 Ti and took roughly 150 GPU hours.

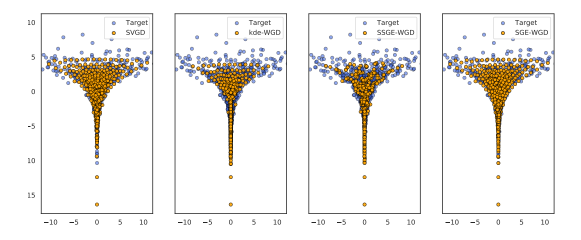


Figure F.1: Neal's funnel. The SGE-WGD and SVGD again fit the distribution best.

### G.1 Sampling from synthetic distributions

Single Gaussian: we created a two-dimensional Gaussian distribution with mean  $\mu=(-0.6871,0.8010)$  and covariance  $\Sigma=\begin{pmatrix} 1.130 & 0.826 \\ 0.826 & 3.389 \end{pmatrix}$ . We used a normal initialization with zero mean and standard deviation  $\sigma^2=3$ . We sampled 100 initial particles and optimized them for

5000 iterations using Adam with a fixed learning rate of 0.1. The kernel bandwidth was estimated using the median heuristic for all methods. For the SSGE we used all the eigenvalues. The random seed was fixed to 42.

**Funnel:** the target distribution followed the density  $p(x,y) = \mathcal{N}(y|\mu=0,\sigma=3)\mathcal{N}(x|0,\exp(y/2))$ . We used a normal initialization with zero mean and standard deviation  $\sigma^2=3$ . We sampled 500 initial particles and optimized them for 2000 iterations using Adam with a fixed learning rate of 0.1. The kernel bandwidth was fixed to 0.5 for all methods. For the SSGE we used all the eigenvalues. The random seed was fixed to 42.

### **G.2** 1D regression

We generated the training data by sampling 45 points from  $x_i \sim \text{Uniform}(1.5, 2.5)$  and 45 from  $x_i \sim \text{Uniform}(4.5, 6.0)$ . The output  $y_i$  for a given  $x_i$  is then modeled following  $y_i = x_i \sin(x_i) + \epsilon_i$  with  $\epsilon_i \sim \mathcal{N}(0, 0.25)$ . We use a standard Gaussian likelihood and standard normal prior  $\mathcal{N}(0, \mathbb{I})$ . The model is a feed-forward neural network with 2 hidden layers and 50 hidden units with ReLU activation function. We use 50 particles initialized with random samples from the prior and optimize them using Adam [35] with 15000 gradient steps, a learning rate of 0.01 and batchsize 64. The kernel bandwidth is estimated using the median heuristic. We tested the models on 100 uniformly distributed points in the interval [0,7]. The random seed was fixed to 42.

### G.3 2D classification

We generate 200 training data points sampled from a mixture of 5 Gaussians with means equidistant on a ring of radius 5 and unitary covariance. The model is a feed-forward neural network with 2 hidden layers and 50 hidden units with ReLU activation function. We use a softmax likelihood and standard normal prior  $\mathcal{N}(0,\mathbb{I})$ . We use 100 particles initialized with random samples from the prior and optimize them using Adam [35] with 10,000 gradient steps, a learning rate of 0.001 and batchsize 64. The kernel bandwidth is estimated using the median heuristic. The random seed was fixed to 42.

### **G.4** Classification on FashionMNIST

On this dataset, we use a feed-forward neural network with 3 hidden layers and 100 hidden units with ReLU activation function. We use a softmax likelihood and standard normal prior  $\mathcal{N}(0,\mathbb{I})$ . We use 50 particles initialized with random samples from the prior and optimize them using Adam [35] for 50000 steps, a learning rate was 0.001 for sge-WGD,kde-WG,ssge-WGD and 0.0025 for kde-fWGD,ssge-fWGD, sge-fWGD, Deep ensemble, fSVGD, SVGD, and batchsize was 256. The kernel bandwidth is estimated using the median heuristic for all different methods. The learning rates were searched over the following values (1e-4, 5e-4, 1e-3, 5e-3, 25e-4) we tested for 50000 and 30000 total number of iterations, 50 and 100 particles and batchsize 256 and 128. All results in Table 1 are averaged over the following random seeds (38, 39, 40, 41, 42).

### G.5 Classification on CIFAR-10

On this dataset, we used a residual network (ResNet32) with ReLU activation function. We use a softmax likelihood and standard normal prior  $\mathcal{N}(0,0.1\mathbb{I})$ . We use 20 particles initialized using He initialization [24] and optimize them using Adam [35] for 50000 steps, a learning rate was 0.00025 for sge-fWGD,kde-fWGD,ssge-fWGD,fSVGD and 0.0005 for kde-WGD,ssge-WGD,sge-fWGD, Deep ensemble and SVGD, and batchsize was 128. The kernel bandwidth is estimated using the median heuristic for all different methods. The learning rates were searched over the following values (1e-4, 5e-4, 1e-3, 5e-3, 25e-4, 5e-5) we tested for 50000 and 30000 total number of iterations, 20 and 10 particles. All results in Table 1 are averaged over the following random seeds (38, 39, 40, 41, 42).