On a Neural Implementation of Brenier's Polar Factorization

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Abstract

In 1991, Brenier proved a theorem that generalizes the QR decomposition for square matrices – factored as PSD \times unitary – to any vector field $F: \mathbb{R}^d \to \mathbb{R}^d$. The theorem, known as the polar factorization theorem, states that any field F can be recovered as the composition of the gradient of a convex function u with a measure-preserving map M, namely $F = \nabla u \circ M$. We propose a practical implementation of this far-reaching theoretical result, and explore possible uses within machine learning. The theorem is closely related to optimal transport (OT) theory, and we borrow from recent advances in the field of neural optimal transport to parameterize the potential u as an input convex neural network. The map M can be either evaluated pointwise using u^* , the convex conjugate of u, through the identity $M = \nabla u^* \circ F$, or learned as an auxiliary network. Because Mis, in general, not injective, we consider the additional task of estimating the ill-posed inverse map that can approximate the pre-image measure M^{-1} using a stochastic generator. We illustrate possible applications of Brenier's polar factorization to non-convex optimization problems, as well as sampling of densities that are not log-concave.

1. Introduction

Brenier proved, through his seminal polar factorization theorem (1991), that *any* vector field can be decomposed into two simpler elements: Given a reference measure ρ supported on $\Omega \subset \mathbb{R}^d$, For any $F : \Omega \to \mathbb{R}^d$, there exists a convex potential $u : \mathbb{R}^d \to \mathbb{R}$ and a *measure-preserving* map $M : \Omega \to \Omega$ (i.e. on has $M_{\sharp}\rho = \rho$), such that $F = \nabla u \circ M$. The polar factorization theorem states that *any* vector field, no matter how irregular, can be reshuffled to match that of the gradient of a convex potential, and that this careful reshuffling in space is achieved by the measure-preserving map M. This paper aims to provide a practical approach to recover approximations of the potential u and vector-valued map M using exclusively samples $x_i \sim \rho$ and their associated images $F(x_i)$. We also highlight how a reliable polar factorization solver, coupled with an estimation of a stochastic generator that mimics the measure-valued inverse map M^{-1} , can be used to study the gradient field of non-convex landscapes. We consider, in particular, the case where the field F of interest is the gradient, with respect to the parameters of a neural architecture, of a learning loss. Note that the polar factorization theorem should not be confused with the major theorem from optimal transport closely associated with Brenier, which we recall in §2. That theorem states that the Monge formulation of the optimal transport (OT) problem, which seeks the push-forward map transporting a measure onto another with the least mean displacements (as measured with squared norms) is solved by the gradient of a convex potential.

Existing Implementations. Shortly after (Brenier, 1991), Benamou and Brenier (1994) proposed a numerical approach to decompose a vector field, with an explicit Eulerian (gridded) approach. Lagrangian approaches have been proposed by Gallouët and Mérigot (2018), while Mérigot and Mirebeau (2016) use a semidiscrete OT formulation. Both are used on low-dimensional manifolds as lower level subroutines to solve Euler's equation for incompressible and inviscid fluids (Arnold, 1966). More recently, Morel et al. (2023) proposed to use Brenier's insight to gradually refactor a normalizing flow as the gradient of a convex map, a.k.a a Monge (1781) map, by applying measure-preserving maps for the Gaussian distribution. Their approach does not, however, rely on neural OT solvers, and focuses instead on untangling an existing flow to turn it gradually into the gradient of a convex potential.

Contributions. We propose in this work a neural implementation of the polar factorization theorem that leverages recent advances in neural optimal transport. More precisely,

 After introducing the polar factorization theorem, as well as neural OT solvers, we show how the two blocks of Brenier's result can be recovered using input convex neural networks (ICNN) (Amos et al., 2017). We modify the ICNN architecture originally proposed in Amos et al.

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(2017) and Korotin et al. (2020), to propose quadratic (low-rank + diagonal) positive definite layers at each layer. Starting from an arbitrary field F, we use the modifications proposed by Amos (2023) to (Makkuva et al., 2020) to train the Brenier convex potential u_{θ} , that appears in the polar factorization of F.

- We study two alternative parameterizations for the measure-preserving map M: Either implicit, relying on the pointwise evaluation of the convex conjugate of u_θ composed with F, or explicit, through an additional network M_ξ trained to map samples x from ρ to ∇u^a_θ ∘ F(x).
- Because M is not, in general, injective, we consider the ill-posed problem of inverting M: we approximate a stochastic map I_ψ, parameterized as a generator, that can generate inputs x such as M(x) = y for a given y. We use bridge matching for this task (De Bortoli et al., 2023).
- We use our approach to factorize gradients of surfaces in low dimensions and show how to use our tools to study the critical points of a non-convex energy g. Factorizing G := ∇g as ∇u_θ ∘ M_ξ, and estimating the stochastic map I_ψ corresponding to M_ξ, our goal is to generate zeros of ∇g. The minimizer of u_θ being ∇u^{*}_θ(0) by definition of convex duality, the points generated as I_ψ(∇u^{*}_θ(0), z) where z is a Gaussian noise of suitable size should, in principle, result in points that are roots of ∇g. We use the cross-entropy loss of a small MNIST digits LeNet (LeCun et al., 1998) classifier and show the ability to sample new parameters with low gradient and good performance on the recognition task.

2. Background

This section introduces neural methods that have been proposed to learn Monge maps between two distributions and recalls the polar factorization theorem in its original form.

2.1. Neural Approaches to the Monge Problem

The Monge formulation of the OT problem between two probability measures μ and $\nu \in \mathcal{P}(\mathbb{R}^d)$ seeks a map $T : \mathbb{R}^d \to \mathbb{R}^d$ that transports μ onto ν , while minimizing the following transport cost:

$$\mathcal{W}_2^2(\mu,\nu) := \inf_{\substack{T:\mathbb{R}^d \to \mathbb{R}^d \\ T_\#\mu=\nu}} \int_{\mathbb{R}^d} \frac{1}{2} \|x - T(x)\|^2 \mathrm{d}\mu(x)$$
(1)

The existence of an optimal map T^* is guaranteed under fairly general conditions (Santambrogio, 2015, §1), when e.g. μ has a density w.r.t. the Lebesgue measure. In that case, Brenier's most famous theorem states that the Monge problem (1) has a unique solution, found at the gradient of a convex function f^* i.e. $T^* = \nabla f^*$. That convex function f^{\star} is itself the solution of the following dual objective:

$$f^{\star} \in \operatorname*{arg inf}_{f \in \mathrm{L}^{1}(\mu)} \int_{\mathbb{R}^{d}} f \mathrm{d}\mu + \int_{\mathbb{R}^{d}} f^{*} \mathrm{d}\nu \tag{2}$$

where the f^* is the convex conjugate of f,

$$f^*(y) := \sup_{x \in \mathbb{R}^d} \langle x, y \rangle - f(x) \,. \tag{3}$$

Note that the star symbol * used for convex-conjugacy should not be confused with the star symbol *, used throughout the paper to denote an optimal solution. The OT map from ν to μ is also given by the inverse of ∇f^{\star} when it exists, $\nabla(f^{\star})^{*}$. The goal of neural OT solvers is to estimate f^* using samples drawn from the source μ and the target distribution ν . Makkuva et al. (2020); Korotin et al. (2020) have proposed methods that build on input convex neural networks (ICNN), as originally proposed by Amos et al. (2017), to parameterize the potential f as an ICNN. The main difficulty in these methods lies in handling the Legendre transform in (2) of the ICNN variable. To address this difficulty, surrogate networks can be used to replace f^* , and we refer to Amos (2023) for the most recent proposal to refine these implementations using amortized optimization. Neural solvers have been used successfully in various applications, notably in single cell genomics (Bunne et al., 2023; 2022); see also (Huang et al., 2020; Cohen et al., 2021).

2.2. Polar Factorization

Given a probability distribution ρ supported on a bounded set Ω , Brenier's polar factorization theorem states that any vector field $F: \Omega \to \mathbb{R}^d$ can be written as the composition of the gradient of a convex function $\nabla u: \Omega \to \mathbb{R}^d$ with a map $M: \Omega \to \Omega$ that preserves the distribution ρ (ie $M_{\#}\rho = \rho$). In that decomposition, ∇u is the unique OT map from Brenier's theorem that transports the measure ρ on $F_{\#}\rho$, since $F_{\#}\rho = (\nabla u \circ M)_{\#}\rho = \nabla u_{\#}(M_{\#}\rho) = \nabla u_{\#}\rho$.

Theorem 2.1 (Brenier polar factorization). Let ρ be a probability measure whose support, $\Omega \subseteq \mathbb{R}^d$, is a bounded set and $F : \Omega \to \mathbb{R}^d$ a square-integrable vector field being non degenerate i.e. $\int_{\mathbb{R}^d} ||F||^2 d\rho < \infty$ and $\rho(F^{-1}(A)) = 0$ on Lebesgue negligible subsets A of \mathbb{R}^d . Then, there exists a convex function $u : \Omega \to \mathbb{R}$ and a map $M : \Omega \to \Omega$ that is measure preserving, i.e. $M_{\#}\rho = \rho$, such that:

$$F = \nabla u \circ M \,. \tag{4}$$

Both M and ∇u are unique.

3. Neural Polar Factorization (NPF)

We describe our method to compute the approximate polar factorization of a field F, using i.i.d samples $(x_1, \ldots, x_n) \sim \rho$ and their evaluations $(F(x_i))_i$. We first estimate the convex potential u in the decomposition $F = \nabla u \circ M$ using an ICNN u_{θ} as an OT Brenier (1991) potential using an improved ICNN architecture. Next, we show that the measure-preserving map M can be defined implicitly for any x, by evaluating the convex conjugate of u_{θ} on F(x): This requires a call to a convex optimization routine at each evaluation. Thanks to our ICNN's strong convexity, the transform (3) is well-posed. To underline the link of that approach to estimate M using u_{θ} , we use the notation $M_{\theta}(x)$. Alternatively, we also propose to learn an amortized model for M, by learning a network M_{ξ} trained on a regression task using paired data samples $\{(x_i, M_{\theta}(x_i))\}$.

3.1. (Low-Rank + Diagonal) Quadratic Layers in ICNNs

ICNNs provide a neural network parameterization of convex functions. We propose a modification of the original architecture presented in (Amos et al., 2017). Our approach is inspired by the Gaussian initialization outlined in (Bunne et al., 2023) and the low-rank quadratic layers presented in (Korotin et al., 2020). The original ICNN was designed to re-inject the input vector x, transformed by an affine map, at every layer, as can be seen in (Amos et al., 2017, Equation 2, $y \rightarrow x$). Korotin et al., §B.2 proposed instead to modify x with multiple low-rank quadratic positive definite (PSD) forms. The PSD constraint ensures convexity of each entry, while the low-rank choice ensures a reasonable number of parameters. We propose quadratic PSD forms that incorporate a *positive* diagonal plus low-rank matrices (Saunderson et al., 2012; Liutkus and Yoshii, 2017):

$$Q_{A,\delta}(x) := \|\delta \circ x\|^2 + \|Ax\|^2 = x^T \left(\text{diag}(\delta) + A^T A \right) x.$$

The network has L+1 layers for $L \ge 1$; we have highlighted in blue the new PSD (diagonal + low-rank) terms:

$$z_{0} = \sigma_{0} \left([Q_{A_{0}^{i},\delta_{0}^{i}}(x)]_{i} + B_{0}x + c_{0} \right),$$

$$z_{\ell+1} = \sigma_{\ell} \left(W_{\ell} z_{\ell} + [Q_{A_{\ell}^{i},\delta_{\ell}^{i}}(x)]_{i} + B_{\ell} x + c_{\ell} \right),$$

$$z_{L+1} = \sigma_{L} \left(w_{L}^{T} z_{\ell} + Q_{A_{L},\delta_{L}}(x) + b_{L}^{T} x + c_{L} \right) \in \mathbb{R}$$

$$u_{\theta}(x) = z_{L+1}$$
(5)

In all layers above, the index *i* spans $1, \ldots, q$, where *q* is the size of the state vectors $z_{\ell} \in \mathbb{R}^{q}$. This augmented ICNN is parameterized with the following family of parameters,

$$\theta = \left(W_{1:L-1} \in (\mathbb{R}^{q \times q}_{+})^{L}, w_{L} \in \mathbb{R}^{q}_{+}, \\ \left(\delta^{i}_{0:L-1} \in (\mathbb{R}^{d}_{+})^{L}, A^{i}_{0:L-1} \in (\mathbb{R}^{r \times d})^{L} \right)_{i=1...q}, \\ \delta_{L} \in \mathbb{R}^{d}_{+}, A_{L} \in \mathbb{R}^{r \times d}, \\ B_{0:L-1} \in (\mathbb{R}^{q \times d})^{L}, b_{L} \in \mathbb{R}^{d}, \\ c_{0:L-1} \in (\mathbb{R}^{q})^{L}, c_{L} \in \mathbb{R} \right).$$

$$(6)$$

The activation functions σ_{ℓ} are convex, non-decreasing nonlinear and all parameters in red in addition to all diagonal vectors δ must be non-negative to ensure convexity.

3.2. Estimating the Convex Potential u

Starting from the existence result outlined in (4), we recover, by applying the push-foward map F on ρ , that

$$F_{\sharp}\rho = (\nabla u \circ M)_{\sharp}\rho = \nabla u_{\sharp}(M_{\sharp}\rho) = \nabla u_{\sharp}\rho.$$

Since u is a convex function, it optimally transports ρ on $\nabla u_{\#}\rho$ in the Monge sense. Therefore, the defining feature of u is that ∇u is the Monge map from ρ to $F_{\sharp}\rho$. We use Amos' solver (2023) to estimate the potential u that pushes ρ onto $F_{\#}\rho$, from the empirical measures $\rho_n := \frac{1}{n} \sum_{i=1}^{n} \delta_{x_i}$ and $F_{\#}\rho_n = \frac{1}{n} \sum_{i=1}^{n} \delta_{F(x_i)}$. Using this solver consists of parameterizing the convex function u as an ICNN u_{θ} following §3.1 and parameterizing ∇u^* directly by an auxiliary vector-valued network V_{ϕ} . The auxiliary network V_{ϕ} is learned by minimizing the objective:

$$\mathcal{L}_{\text{convex-dual}}(\phi) = \frac{1}{n} \sum_{i=1}^{n} \|V_{\phi}(F(x_i)) - \nabla u_{\theta}^*(F(x_i))\|^2$$

One can show, using Danskin's envelope theorem (1966), that $\nabla u_{\theta}^{*}(y)$ is the maximizer of the convex conjugate (3) problem for u at y,

$$\nabla u_{\theta}^{*}(y) = \arg \sup_{x} \langle x, y \rangle - u_{\theta}(x)$$

Because u_{θ} is strictly convex, we compute the optimal solution solving $u^*(F(x))$ with a conjugate solver, e.g. gradient ascent, (L)BFGS (Liu and Nocedal, 1989) or ADAM (Kingma and Ba, 2014). We call a conjugate solver CS any algorithm that, for a given pair (u, y), outputs an approximation of $\nabla u^*(y)$. This results in the loss:

$$\mathcal{L}_{\text{convex-dual}}(\phi) = \frac{1}{n} \sum_{i=1}^{n} \|V_{\phi}(F(x_i)) - \mathbf{CS}(u_{\theta}, F(x_i))\|^2$$
(7)

In practice, the latter is initialized with the predictions of V_{ϕ} , which considerably reduces the number of iterations required for the solver to converge when V_{ϕ} starts making correct predictions. The parameters of the network u_{θ} are then updated alternatively, by taking steps along the gradients of the original dual objective of Makkuva et al. (2020):

$$\mathcal{L}_{\text{Monge}}(\theta) = \frac{1}{n} \sum_{i=1}^{n} u_{\theta}(x_i) + \langle V_{\phi}(F(x_i)), F(x_i) \rangle - u_{\theta}(V_{\phi}(F(x_i)))$$
(8)

3.3. Estimating the Measure-Preserving Map M

In the polar decomposition of F, ∇u is tasked with transporting ρ on $F_{\#}\rho$, the measure-preserving map M ensures then that $F = \nabla u \circ M$. To express M as a function of F

and u, one simply has to apply the inverse of ∇u on both sides. When u is strictly convex, we simply rely on the identity $\nabla u^* \circ \nabla u = \text{Id}$ to obtain:

$$M = \nabla u^* \circ F \tag{9}$$

Evaluating M using a Conjugate Solver. Given a conjugate solver CS and the estimate u_{θ} for the ground truth potential u, we can inject them in (9) to get an estimation $CS(u_{\theta}, F(x))$ of M(x) for a given x. Since this estimation depends on θ , we define that approximation as $M_{\theta}(x)$,

$$M_{\theta}(x) := \mathbf{CS}(u_{\theta}, F(x)), \qquad (10)$$

with a slight abuse of notation, since θ should not be understood as a parameter parameterizing M, but instead defining it implicitly through u_{θ} and CS.

Neural Estimation for M. While M_{θ} does indeed provide an estimate of M, it may be convenient to parameterize the measure-preserving map of interest as a neural network M_{ξ} , defined with an independent set of parameters ξ . Borrowing a page from amortized optimization (Amos et al., 2023), M_{ξ} can be used to initialize the conjugate solver used to estimate M_{θ} or even replace it when M_{ξ} is sufficiently accurate. Furthermore, the parameterization of M by M_{ξ} is sometimes necessary when, e.g., F is only given on a few samples, and one wishes to evaluate M at any point. The neural map M_{ξ} is then trained to minimize the following mean-squared error:

$$\mathcal{L}_{\text{preserving}}(\xi) = \frac{1}{n} \sum_{i=1}^{n} \|M_{\xi}(x_i) - \mathbf{CS}(u_{\theta}, F(x_i))\|^2 \,. \tag{11}$$

Note that while the loss in (11) resembles (7), the network V_{ϕ} takes the transported point $F(x_i)$ as an input, whereas M_{ξ} is only given x_i .

Evaluating The Measure Preservation of M. In both cases, M_{θ} , as evaluated with a conjugate solver, or its independently evaluated neural counterpart M_{ξ} should be measure-preserving. Indeed, we will use (as in Figure 1, bottom center plots) any departure from the identity

$$M_{\#}\rho = (\nabla u^* \circ F)_{\#}\rho = (\nabla u^*)_{\#}(F_{\#}\rho) = \rho$$

as a way to assess the quality of our factorization.

3.4. Sampling according to the pre-image measure M_{θ}^{-1}

Measure-preserving maps M are not invertible in general (Ryff, 1970), a well-known example in 1D being the doubling map defined as $M(x) = 2x \mod 1$ that preserves the Lebesgue measure rescaled to the interval [0, 1]. This non-invertibility is of particular interest in the optimization and sampling applications we propose. For a given y, our goal will therefore be to generate inputs x such that $M_{\theta}(x) = y$. To this end, we learn a generative process to sample according to the posterior density

$$\pi_{\theta}(x|y) = \frac{\mathbf{1}_{y=M_{\theta}(x)}\rho(x)}{\int_{x}\mathbf{1}_{y=M_{\theta}(x)}\rho(x)\mathrm{d}x}.$$
 (12)

We rely on the augmented bridge matching procedure presented in De Bortoli et al. (2023) to learn the drift of the stochastic differential equation (SDE) formulated in (13) so that, on input $X_0 = y$, the generated samples X_1 be distributed according to $\pi_{\theta}(x|y)$ (12).

$$dX_t = (X_{\psi}(X_0, X_t) - X_t) / (1 - t) dt + \sigma dB_t$$
(13)

De Bortoli et al.'s approach refines the bridge matching procedures that have been recently used to solve inverse problems (Somnath et al., 2023; Liu et al., 2023; Chung et al., 2024), by augmenting the learnable part of the drift X_{ψ} with the initial point X_0 of the SDE. This slight adjustment allows to correctly recover the coupling measure $(M_{\theta}, \text{Id})_{\#}\rho$ from the paired samples $\{(x_i, F(x_i))\}_{i=1}^n$ when X_{ψ} is parameterized using a multilayer perceptron trained according to Algorithm 1.

Alg	orithm 1 Training of X_{ψ}
1:	$u_{\theta} \leftarrow$ Trained ICNN s.t. $\nabla u_{\theta \#} \rho \approx F_{\#} \rho$
2:	Initialize X_{ψ}
3:	while not converged do
4:	Draw a sample $(x_i, F(x_i))$
5:	Compute $y_i = CS(u_\theta, F(x_i))$
6:	Sample $t \sim \mathcal{U}([0,1])$
7:	Sample $z_i \sim \mathcal{N}(0, I_d)$
8:	$x_t := (1-t)y_i + tx_i + \sigma(t(1-t))^{1/2}z_i$
9:	$\mathcal{L}_{\psi} \leftarrow rac{1}{n} \ X_{\psi}(y_i, x_t) - x_i \ ^2$
10:	Update X_{ψ} using $\nabla \mathcal{L}_{\psi}$
11:	end while

The optimized network X_{ψ} is then plugged in (13) that we solve with Heun's method as implemented in diffrax (Kidger, 2021) using S discretization steps. Given a sample y from $M_{\theta \#}\rho$, solving the SDE (13) using $X_0 = y$ allows to generate an output X_1 distributed according to the posterior density (12). To alleviate the notations, we call I_{ψ} the generative process such that $I_{\psi}(y, \mathbf{z})$ is the output X_1 returned by the differential equation solver associated to (13) on the input $X_0 = y$ when the injected gaussian noise z has been drawn from $\mathcal{N}(0, I_d)^{\otimes S}$

$$I_{\psi}(y, \mathbf{z}) = \text{SDE}(X_{\psi}, y, \mathbf{z}), \ y \in \mathbb{R}^d, \ \mathbf{z} \in \mathbb{R}^{d \times S}.$$

To generate several inputs x from $\pi_{\theta}(x|y)$, one only needs to inject different noises $\mathbf{z} \sim \mathcal{N}(0, I_d)^{\otimes S}$ in $I_{\psi}(y, \cdot)$, i.e.

$$I_{\psi}(y,\cdot)_{\#}\mathcal{N}(0,I_d)^{\otimes S} \approx \pi_{\theta}(\cdot|y).$$

4. NPF to Study Non-Convex Potentials g

In this section, we focus on the polar factorisation of the gradient field ∇g , where g is a non-convex function of interest. We show how computing the NPF of ∇g together with the inverse map I_{ψ} can be used to explore the space of critical points of g.

4.1. On using the Inverse Map I_{ψ} of ∇g

NPF on $G = \nabla g$. Let $g : \mathbb{R}^d \to \mathbb{R}$ be a function of interest supported on a bounded set $\Omega \subset \mathbb{R}^d$. Assuming that $\nabla g : \mathbb{R}^d \to \mathbb{R}^d$ meets the requirements of (4), Brenier's polar factorization states the existence of a convex function u and a measure-preserving M that preserves the rescaled Lebesgue measure on Ω , \mathcal{L}_Ω such that:

$$\nabla g = \nabla u \circ M.$$

For a given vector v, the points in Ω whose gradient with respect to g is equal to v are all transported by M on the same point $\nabla u^*(v)$ i.e.

$$M\left(\{x\in\Omega:\nabla g(x)=v\}\right)=\left\{\nabla u^*(v)\right\}.$$

In particular, the critical points of g are all mapped by M onto the minimizer of the function u, which is $\nabla u^*(0)$.

On Extracting the Critical Points of g. When the NPF of ∇g is learned, resulting in u_{θ} , M_{ξ} and I_{ψ} , composing I_{ψ} with $\nabla u_{\theta} *$ provides an inversion process for ∇g . Generating an input point x_v whose gradient is v can in fact be done by first sampling $\mathbf{z} \sim \mathcal{N}(0, I_d)^{\otimes N}$ and successively applying $\nabla u_{\theta} *$ and I_{ψ} to v:

$$x_v = I_{\psi}(\nabla u_{\theta}^*(v), \mathbf{z}), \text{ where } \mathbf{z} \sim \mathcal{N}(0, I_d)^{\otimes S}$$

As a special case, sampling the critical points of g is done by taking v = 0 in the above procedure with different noises z. Note, however, that this convexification requires estimating the polar factorization of ∇g as well as the inverse map I_{ψ} over the entire space Ω , which is computationally expensive. To optimize the g function, we propose instead to combine this method with the Langevin Monte Carlo (LMC) algorithm to correctly estimate the polar factorization of ∇g around the minimums of g.

4.2. A LMC Method Assisted by NPF.

LMC algorithm Given a smooth log-concave density

$$\pi(x) = \frac{e^{-g(x)}}{\int_{x \in \mathbb{R}^d} e^{-g(x)} \mathrm{d}x},$$
(14)

with $g: \mathbb{R}^d \longrightarrow \mathbb{R}$, the Langevin Monte Carlo algorithm can sample from π by starting from $x^{(0)}$ to iterate

$$x^{(k+1)} = x^{(k)} - \gamma \nabla g(x^{(k)}) + \sqrt{2\gamma} z^{(k)}, \ z^{(k)} \sim \mathcal{N}(0, I_d)$$

When g is non-convex, the LMC algorithm lacks guarantees (Roberts and Tweedie, 1996; Cheng and Bartlett, 2018; Dalalyan and Karagulyan, 2019). In particular, when g has multiple local minima, the generated samples are highly correlated as the particles originating from the LMC algorithm often get stuck in some basins. For this reason, the LMC algorithm has been combined with methods enabling global jumps between modes (Pompe et al., 2020; Gabrié et al., 2022) to sample multi-modal distributions.

Sampling with Known Polar Factorization for ∇g . In this paragraph, we assume that the polar factorization $(\nabla u, M)$ and stochastic inverse map M^{-1} of ∇g are known. To sample the modes of $\pi(x) \propto e^{-g(x)}$ when g is nonconvex, one can run the LMC algorithm on the convex function u and sample back using an inverse generator M^{-1} :

$$y^{(k)} = M(x^{(k)})$$

$$y^{(k+1)} = y^{(k)} - \gamma \nabla u(y^{(k)}) + \sqrt{2\gamma} z^{(k)}$$

$$x^{(k+1)} = M^{-1}(y^{(k+1)}, z^{(k+\frac{1}{2})}).$$

The LMC step on u allows to move along a new descent direction or exploration direction to reach y_{k+1} while M^{-1} randomly generates a point $x^{(k+1)} \in \Omega$ whose gradient for g is $\nabla u(y_{k+1})$. This way, the neighborhoods of g's critical points are uniformly sampled, and a particle does not get stuck in one minimum as M^{-1} permits global moves between all the basins. Because it is difficult to differentiate a minimum from a saddle point or a maximum when sampling critical points using the polar factorization of ∇u , this procedure should be combined with Langevin steps on g to escape non-minimum critical points. The following paragraph details the sampling algorithm and complements it by showing how the polar factorization of ∇u can be learned while sampling.

Unknown Polar Factorization for ∇g When the polar factorization is unknown, we propose an algorithm that learns the polar factorization of ∇g as well as the inverse map I_{ψ} using the generated particle trajectories. The algorithm alternates between N Langevin steps on g and N Langevin steps on u_{θ} , while M_{θ} and I_{ψ} allow to transition between the two spaces. Algorithm 2 details the steps of the procedure. The notation LMC $(u_{\theta}, \gamma, y_i^{(k)}, N)$ means that N LMC steps are performed on the function u_{θ} with a time step of γ starting from the point $y_i^{(k)}$.

Algorithm 2 LMC-NPF

1: Initialize u_{θ} and I_{ψ} 2: Initialize the particles $\{x_i^{(0)}\}_{1 \le i \le n}$ 3: $k \leftarrow 0$ 4: while $k < k_{max}$ do $\begin{aligned} & \text{if } k \mod N = 0 \text{ then} \\ & y_i^{(k)} = M_{\theta}(x_i^{(k)}) \\ & y_i^{(k+1)} = \text{LMC} \left(u_{\theta}, \gamma, y_i^{(k)}, N \right) \\ & x_i^{(k+1)} = I_{\psi}(y_i^{(k+1)}, \mathbf{z}) \text{ with } \mathbf{z} \sim \mathcal{N}(0, I_d)^{\otimes S} \end{aligned}$ 5: 6: 7: 8: else $x_i^{(k+1)} = x_i^{(k)} - \gamma \nabla g(x_i^{(k)}) + \sqrt{2\gamma} z_i^{(k)}$ 9: 10: 11: Update u_{θ}, I_{ψ} with $\{(x_i^{(k)}, \nabla g(x_i^{(k)}))\}_{1 \leq i \leq n}$ 12: $k \leftarrow k + 1$ 13: 14: end while

The main insight of the proposed sampling algorithm is that LMC steps permit the exploration of the space locally, while NPF provides and stores a more global viewpoint, that is able to propose moves to potentially worthy areas.

5. Experiments

5.1. Accuracy Metrics

Assess NPF's Accuracy. When a field G is only available through samples, the following three criteria, evaluated on unseen samples (or test set) $\{(x_j, G(x_j))\}_{1 \le j \le m}$, are used to assess whether the estimated polar factorization is correct.

- To measure that the distributions ∇u_{θ#}ρ and G_#ρ are close, we compute the Sinkhorn divergence S_ε (Ramdas et al., 2017; Genevay et al., 2018; Peyré et al., 2019) between the two point clouds (G(x_j))_{1≤j≤m} and (∇u_θ(x_j))_{1≤j≤m}. To quantify the scale of that measurement, we compare it with the distance between two batches of fixed size drawn from (G(x_j))_{1≤j≤m}. We also visualize this proximity by embedding the two point clouds using the TSNE algorithm (Van der Maaten and Hinton, 2008) and superimpose them.
- The second criterion assesses whether M_{ξ} is measurepreserving. Similarly, this is numerically estimated by computing the Sinkhorn divergence between the empirical measures associated with $(x_j)_{1 \le j \le m}$ and $(M_{\xi}(x_j))_{1 \le j \le m}$, and visualized with a TSNE embedding.
- Finally, we evaluate the L₂ distance between G and ∇u_θ ∘ M_ξ using the test set. Note that when M_θ is used (rather than M_ξ), that criterion is not useful since it only assesses the quality of the conjugate solver.

Assess the Generative Inverse Map I_{ψ} . Given y, we should be able to sample among the antecedents of y by M_{θ} using the multivalued map I_{ψ} . To quantify that, we esti-

mate the average distance between the probability associated to the density $\pi_{\theta}(x|y)$ (12) and $I_{\psi}(y, \cdot)_{\#}\mathcal{N}(0, I_d)^{\otimes S}$ from samples. Given the finite test set $\{(x_j, M_{\theta}(x_j))\}_{1 \leq j \leq m}$, it is unlikely to find a multitude of points with the same image. For this reason, we approximate $M_{\theta}^{-1}(M_{\theta}(x_k))$, by constructing the set

$$\mathcal{B}_{\alpha}(x_k) = \{x_j : \|M_{\theta}(x_j) - M_{\theta}(x_k)\|_2 \le \alpha\}$$

and choose α such that the cardinal of $\mathcal{B}_{\alpha}(x_k)$ is 100. We then compute the sinkhorn divergence between the predictions of I_{ψ} on $M_{\theta}(\mathcal{B}_{\alpha_k}(x_k))$ and $\mathcal{B}_{\alpha_k}(x_k)$ and average it over all the x_k . We compare the obtained value with the distance of two batches of fixed size drawn independently from the $(x_j)_{1 \leq j \leq m}$. We also evaluate the fact that the stochastic map $I_{\psi} \circ \nabla u_{\theta}^*$ approximates G^{-1} by computing the quantity, using MC samples for z,

$$\frac{1}{m} \sum_{j=1}^{m} \mathbb{E}_{\mathbf{z} \sim \mathcal{N}(0, I_d)^{\otimes S}} \| G \circ I_{\psi}(\nabla u_{\theta}^*(G(x_j)), \mathbf{z}) - G(x_j) \|^2$$
(15)

We also study the quantity where the cosine similarity replaces the norm in (15).

5.2. NPF of Topographical Data

Dataset. We use the Python package elevation to get the elevation of three regions of the world: Chamonix, London, and Cyprus. We estimate the gradients associated with the elevation in these regions with finite-differences, and obtain three datasets composed of (latitude, longitude) points paired with their gradients. We learn the polar factorization $(\nabla u_{\theta}, M_{\xi})$ of the underlying gradient field as well as the inverse map I_{ψ} . Because in these examples, G is only given through samples, we parameterized the measure-preserving map using a neural network M_{ξ} . To assess the quality of our method NPF, we used a 85% training / 15% test split. More details can be found in the appendix.

	Chamonix
$S_{\varepsilon}(\nabla u_{\theta \#} \rho_n, G_{\#} \rho_n)$	0.27
$S_{\varepsilon}(G_{\#}\rho_n, G_{\#}\rho'_n)$	1.55
$S_{\varepsilon}(M_{\xi\#}\rho_n,\rho_n)$	0.0029
$S_{arepsilon}(ho_n, ho_n')$	0.0034
$\mathbb{E}_{x \sim \rho_n} \left[\ G(x) - \nabla u_\theta \circ M_\xi(x) \ _2 \right]$	0.96
$\mathbb{E}_{x,\mathbf{z}}\left[S_{\varepsilon}((I_{\psi}(M_{\theta}(\mathcal{B}_{\alpha}(x)),\mathbf{z}),\mathcal{B}_{\alpha}(x)))\right]$	0.048
$S_{arepsilon}(ho_{64}, ho_{64}')$	0.077
$S_{arepsilon}(ho_{128}, ho_{128}')$	0.039

Table 1. Polar factorization and Inverse multivalued map metrics for learning the gradient of the elevation in Chamonix area. For these metrics, ρ_n and ρ'_n are two empirical measures created from n = 1024 samples drawn independently from the test set. The ε parameter is set equal to 0.1 which is the default parameter.

Polar Factorization Results. Table 1 shows that the estimated NPF is accurate: the Sinkhorn divergence between the predicted distribution $\nabla u_{\theta \#} \rho_n$ and the target distribution $G_{\#} \rho_n$ is lower than the divergence between two batches of size 1024 taken from the target. Similarly, the Sinkhorn divergence between ρ_n and its image by M_{ξ} is lower than that between two batches of size 1024 drawn from the source. The reconstruction of *G* is also quite satisfactory as corroborated visually (Figure 1).

Inverse Map Results. The data from Table 1 indicates that I_{ψ} generates the antecedents of the images by M_{θ} accurately: the estimated quantity $\mathbb{E}_{x,\mathbf{z}}[S_{\varepsilon}((I_{\psi}(M_{\theta}(\mathcal{B}_{\alpha}(x)),\mathbf{z}),\mathcal{B}_{\alpha}(x))]$ is comparable to the distance between two batches drawn from the source distribution whose size lies between 64 and 128. To visualize these performances, we transported the samples $(G(x_i))_{1 \le i \le m}$, that store gradients of the elevation, using $I_{\psi} \circ \nabla u_{\theta}^{*}$ that should estimate the inverse generative map G^{-1} . We expect very high gradients to be sent to points where the elevation varies rapidly, such as the sides of mountains in the Chamonix example. To visualize where a gradient was sent, we plot a point at this localization and color it according to the norm of the gradient from which it originates. We compare the image generated by this process with the one obtained by coloring directly the points $(x_j)_{1 \le j \le m}$ using their associated gradients. In the three cases (Chamonix, London, Cyprus), the two images are quite similar (Figure 2), showing the quality of our reconstruction.

5.3. Learn an NN Optimization Landscape using NPF

In this experiment, we consider a minimal neural architecture capable of classifying MNIST digits. Inspired by the LeNet architecture (LeCun et al., 1998), we use two convolutional layers, each followed by a Relu and a max pooling operation. A classification layer leads to an output layer of 10 neurons, followed by a softmax. The loss function is the cross entropy, computed with MNIST train dataset minibatches of size 128, and the vector field under study is the gradient of that loss for the d = 222 parameters of the neural network. The loss landscape of a non-linear neural network being very chaotic (Li et al., 2018), we do not expect to learn the polar factorization of the associated gradient field perfectly over the all optimization space Ω . The optimization space we are considering is $\Omega = [-1, 1]^{222}$.

Polar Factorization and Inverse Map Results. According to Table 2, we see that, overall, NPF manages to learn that vector field, but it lacks, as expected, accuracy in some parts of the space. This is, e.g., revealed visually using the TSNE plot from Figure 3. Similarly, I_{ψ} can be used to invert M_{θ} according to Table 3 and the histogram associated with the cosine similarity on Figure 3 confirms that we can

	n = 1024	n = 2048
$S_{\varepsilon}(\nabla u_{\theta \#}\rho_n, G_{\#}\rho_n)$	107.5 ± 8.2	93.5 ± 5.3
$S_{\varepsilon}(G_{\#}\rho_n, G_{\#}\rho'_n)$	110.3 ± 8.7	87.4 ± 4.2

Table 2. Polar factorization metrics for learning the gradient of the MNIST classifier loss function.

$\mathbb{E}_{k,\mathbf{z}}\left[S_{\varepsilon}((I_{\psi}(M_{\theta}(\mathcal{B}_{\alpha}(x_k)),\mathbf{z}),\mathcal{B}_{\alpha}(x_k)))\right]$	106.2 ± 0.5
$S_{arepsilon}(ho_{64}, ho_{64}')$	111.0 ± 0.6
$S_{arepsilon}(ho_{128}, ho_{128}')$	105.7 ± 0.4
$\mathbb{E}_{y \sim G_{\#}\rho_n, \mathbf{z}} \ G \circ I_{\psi}(\nabla u_{\theta}^*(y), \mathbf{z}) - y \ _2$	14.2 ± 13.0
$\mathbb{E}_{y \sim G_{\#}\rho_n, \mathbf{z}} \operatorname{cosine}(G \circ I_{\psi}(\nabla u_{\theta}^*(y), \mathbf{z}), y)$	0.83 ± 0.18

Table 3. Inverse multivalued map metrics for learning the gradient of the MNIST classifier loss function.

choose a certain gradient v and use $I_{\psi} \circ \nabla u_{\theta}^*$ to generate classifier weights whose gradient is approximately v. In particular, I_{ψ} allows the generation of correct critical points (Figure 12), which, however, have a low accuracy. This may be due to the fact that the gradient of good minimums varies greatly with the stochasticity of the loss function visible on Figure 13 and Figure 15, compared with the gradient of critical points with an accuracy of 10%, which is the performance of a classifier with random weights.

5.4. Learning NPF using Gradient Flow of Particles

In §5.3, we requested that NPF learn the entire gradient field. This of course limits the ability, given a certain budget of samples, to provide a good approximation of critical points through the inverse generative map. This is likely due to our uniform sampling procedure, which is unlikely to reveal interesting critical points. In this experiment, we train NPF using gradient descent trajectories to focus on those areas. To do this, we initialize 1024 particles randomly and have them follow a gradient flow. We use these trajectories' samples to learn NPF.

Polar Factorization Results. The gradients associated with particles in good accuracy basins vary greatly with loss stochasticity (see Figure 13). However, we can see that NPF is faithful in those areas: the Sinkhorn divergence between the distribution generated by ∇u_{θ} and the target is of the same order as that between two batches of size 1024.

Inverse Map Results. As for I_{ψ} , the gradients of the generated weights do have a norm close to 0, and the cosine similarity distribution reveals that the direction of the gradients is globally learned but stochasticity prevents us from performing better. Moreover, we can see in Figure 14 that the critical weights generated contain valid minima.



Figure 1. The g function under study is the elevation in the Chamonix area (France). The figures show the respective action of the vector fields involved in the polar factorization of ∇g on a sample measure ρ_n . We observe that $\nabla u_{\theta \#} \rho_n \approx G_{\#} \rho_n$. Both implicit and explicit measure-preserving maps M_{θ} (10) as well as the explicit network M_{ξ} trained with the loss (11) permutes the points of the distribution, ensuring that $G \approx \nabla u_{\theta} \circ M_{\xi}$ while $(M_{\xi})_{\#} \rho_n \approx (M_{\theta})_{\#} \rho_n \approx \rho_n$.



Figure 2. I_{ψ} 's ability to replace gradients in the original Ω space for the example of Chamonix region's elevation gradient. The figure on the right is generated by returning the gradients $\nabla g_{\#}\rho_n$ to their initial position in the image via $I_{\psi} \circ \nabla u_{\theta}^*$. This position is then colored according to the initial gradient norm (before transport). We can compare the result with the image on the left, generated by sampling uniformly in Ω space and colored according to the norm of their gradient.



Figure 3. Performance of the NPF and the inverse map I_{ψ} on the §5.3 MNIST classifier experiment where the loss gradient is learned over the entire space Ω . The TSNE allows to visualize in 2D the overlap of the predicted distribution $\nabla u_{\theta \#} \rho_n$ with the target distribution $G_{\#} \rho_n$ while the cosine similarity, mentioned in §5.1, shows that I_{ψ} permits to accurately generate weights associated with a given gradient.

5.5. LMC-NPF on MNIST-XE

We use LMC-NPF (Algorithm 2) to sample the loss of the MNIST classifier considered in §5.3. Our sampling algorithm is preceded by a warm-up containing particle descents to explore good minima before sampling as detailed in the supplementary files.

Sampling Algorithm Results. Following the warm-up, the TSNE (Figure 5) as well as Figure 4 show that our sampling algorithm proposes high-accuracy weights that are completely different from the minima found during the warm-up period. Since LMC-NPF alternates between Langevin steps on g and Langevin steps on u_{θ} , we demonstrate that these minima had indeed been discovered through the use of NPF by running an LMC algorithm initialized with the final warm-up particles. The latter was parameterized the same way as the one used in LMC-NPF, with the same number of iterations. We observe that the LMC algorithm samples around the warm-up particles but does not detach itself from them. This confirms that the use of PFNet in the sampling procedure permits the discovery of new local minima.

6. Conclusion

Brenier's polar factorization is arguably one of the most far-reaching results discovered in analysis in the last century, underpinning the better known Brenier theorem on the existence of solutions to the Monge (1781) problem. We proposed in this work the first implementation, to the best of our knowledge, of that factorization that is applicable to higher-dimensional settings. To do so, we have used the recently proposed machinery of neural optimal transport solvers. Beyond simply exploiting this result, we have also proposed to estimate a multivalued map that approximates the inverse of the measure-preserving map component in the polar factorization. We have shown that such an inverse map can be of potential use to sample the optimization land-



Figure 4. Characteristics of the points sampled using LMC-NPF for the §5.5 MNIST classifier experiment. In gray the classifier weights have been drawn uniformly in the Ω optimization space, while in purple the weights have been sampled using Algorithm 2. Generated samples are critical points which are good minima, as shown by the accuracy statistics.



Figure 5. Results of LMC-NPF applied to the MNIST classifier loss function. The TSNE is used to represent the final particles resulting from descent trajectories during the Warm-up period (in red), the particles sampled by the LMC algorithm (in grey), and the particles sampled by LMC-NPF (in purple).

scape of non-convex potentials. An interesting direction for perfecting the sampling algorithm would be to reweight the samples according to their probability, in the same vein as SMC samplers (Del Moral et al., 2006). This would require knowledge of the probability distribution generated by the generative model I_{ψ} , which is not possible with the current methodology.

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A. Computation of the convex conjugate

Given a function convex function u, and a point y, the objective $J_u(x, y) = \langle y, x \rangle - u(x)$ is concave with respect to x and $u^*(y) = \sup_x J_u(x, y)$ can be computed using optimization algorithms like gradient ascent, (L)BFGS or Adam. As for $\nabla u^*(y)$, taking the gradient for y necessitates differentiate through a supremum. In our case, u is strictly convex a.e. and the supremum becomes a maximum :

$$u^*(y) = \max J_u(x, y)$$

Danskin's envelope theorem (1966) allows to differentiate through this maximum and to write:

$$\nabla_y u^*(y) = \nabla_y \max_x J_u(x, y)$$
$$= (\nabla_y J_u(x, y))(x^*(y))$$

where $x^*(y)$ is the optimal x that maximizes $J_u(x, y)$. Because $\nabla_y J_u(x, y) = x$, we get that

$$\nabla_y u^*(y) = x^*(y)$$

B. Preconditionned LMC

$$x^{(k+1)} = x^{(k)} - \gamma \nabla f(x^{(k)}) + \sqrt{2\gamma} z^{(k)}, \quad z^{(k)} \sim \mathcal{N}(0, I_d)$$

By replacing ∇f with its polar factorization, the procedure becomes:

$$x^{(k+1)} = x^{(k)} - \gamma \nabla u \circ M(x^{(k)}) + \sqrt{2\gamma} z^{(k)}$$

By studying $y^{(k)} = M(x^{(k)})$, one can see that the LMC procedure implies doing a preconditioned LMC algorithm on the convex function u.

$$y^{(k+1)} = M(x^{(k+1)})$$

= $M\left(x^{(k)} - \gamma \nabla u(y^{(k)}) + \sqrt{2\gamma}z^{(k)}\right)$
= $M(x^{(k)}) + J_M(x^{(k)}) \left[-\gamma \nabla u(y^{(k)}) + \sqrt{2\gamma}z^{(k)}\right]$
+ $\circ(\|\varepsilon\|)$
= $y^{(k)} - \gamma J_M(x^{(k)}) \nabla u(y^{(k)}) + \sqrt{2\gamma}J_M(x^{(k)})z^{(k)}$
+ $\circ(\|\varepsilon\|)$

with $\varepsilon = -\gamma \nabla u(y^{(k)}) + \sqrt{2\gamma} z^{(k)}$. One can note that the preconditioned matrix $H = J_M(x^{(k)})$ is not necessarily positive definite.

C. Augmented Bridge Matching

Given a coupling $\Pi_{0,1}$ and random variables (X_0, X_1) , the Augmented bridge matching algorithm (De Bortoli et al., 2023) aims at learning a stochastic dynamic mapping between X_0 and X_1 that preserves the coupling $\Pi_{0,1}$.

In the probability space of path measures $\mathcal{P}(\mathcal{C}([0,1],\mathbb{R}^d))$, let \mathcal{M} denotes the path measures associated to the SDE $dX_t = v_t(X_t)dt + \sigma_t dB_t$, the functions σ and v being locally Lipschitz. Given a path measure $\mathbb{Q} \in \mathcal{M}$, the diffusion bridge of \mathbb{Q} which is the distribution of \mathbb{Q} conditioned on both endpoint is denoted by $\mathbb{Q}_{|0,1}$. The set of path measures considered to bridge $\mathcal{P}(X_0)$ and $\mathcal{P}(X_1)$ according to the coupling $\Pi_{0,1}$ is $\Pi_{0,1}\mathbb{Q}_{|0,1} = \int_{\mathbb{R}^d \times \mathbb{R}^d} \mathbb{Q}_{|0,1}(.|x_0, x_1)\Pi_{0,1}(dx_0, dx_1)$. In De Bortoli et al. (2023), the authors showed that under mild conditions, $\Pi_{0,1}\mathbb{Q}_{|0,1}$ was associated to the following SDE:

$$dX_t = \{b_t(X_t) + \sigma_t^2 u_t\}dt + \sigma_t dB_t, \qquad X_0 \sim \mu$$

with $u_t = \mathbb{E}_{\mathbb{P}_{1|0,t}} \left[\nabla \log \mathbb{Q}_{1|t}(X_1|X_t) | X_0, X_t \right]$ where $\mathbb{Q}_{1|t}$ and $\mathbb{P}_{1|0,t}$ are respectively the conditional distribution of \mathbb{Q} at time 1 given the state at time t and the conditional distribution of \mathbb{P} at time 1 given the coupling state at time 0 and t.

This SDE gives a way to sample from $\Pi_{0,1}$ by first sampling $X_0 \sim \mu$ and then discretize the SDE to get X_1 . Because u_t is intractable, it is approximated by a neural network u_t^{θ} learned to minimize the regression loss:

$$\int_{0}^{1} \lambda_{t} \mathbb{E}[\|u_{t}^{\theta}(X_{0}, X_{t}) - \nabla \log \mathbb{Q}_{1|t}(X_{1}|X_{t})\|^{2}] d\mathbb{P}(X_{0}, X_{t}, X_{1})$$

A particular case of diffusion bridge is the Brownian bridge $\mathbb{Q}_{|0,1}$ for which v = 0 and $\sigma_t = \sigma$ which is the one usually used in practice.

D. Topography experiments

D.1. Creation of the dataset

We used the Python package elevation to get the elevation of three different regions of the globe: Chamonix, London, and Cyprus. Given the latitudes and longitudes of the desired area, elevation returns a grid of the area with the elevation value at each grid point. For the Chamonix example, we obtained 323932 points $(x, y) \in \mathbb{R}^2$ and their corresponding elevation. We dequantized the elevations by adding a uniform noise on [0,1] to them before using a Gaussian filter to make the gradients smoother. To do this, we used the function gaussian_filter from the scipy library. We then numerically estimate the gradients associated with the elevation and obtain a dataset of 323932 points in \mathbb{R}^2 and the associated gradients in \mathbb{R}^2 for the example of Chamonix. We obtained data for the Cyprus and London regions in the same way.

D.2. London and Cyprus results



Figure 6. Respective actions of the learned vector fields associated with the polar factorization of the elevation gradient in the London area.



Figure 7. I_{ψ} 's ability to replace gradients in the original Ω space for the example of London region's elevation gradient.



Figure 8. Respective actions of the learned vector fields associated with the polar factorization of the elevation gradient in the Cyprus neighborhood.



Figure 9. I_{ψ} 's ability to replace gradients in the original Ω space for the example of Cyprus region's elevation gradient.

	Chamonix	London	Chypre
$W_{\varepsilon}(\nabla u_{\theta \#}\rho_n, F_{\#}\rho_n)$	0.27	0.11	0.020
$W_{\varepsilon}(\nabla F_{\#}\rho_n, F_{\#}\rho'_n)$	1.55	0.33	0.036
$\mathbb{E}_{k,\mathbf{z}}\left[S_{\varepsilon}((I_{\psi}(M_{\theta}(\mathcal{B}_{\alpha}(x_k)),\mathbf{z}),\mathcal{B}_{\alpha}(x_k))\right]$	0.048	0.23	0.041

Figure 10. NPF and I_{ψ} performances for the topography experiments.

E. LeNet classifier Experiments

E.1. LeNet classifier architecture

The LeNet classifier architecture used for the experiments is composed of two convolutive layers followed by a relu activation function and a max pooling; it ends with a dense layer as described in Figure 11. The optimization space that we consider is $\Omega = [-1, 1]^{222}$.

E.2. Complementary graphs for the MNIST classifier experiments



Figure 11. Le Net classifier architecture used in experiments.



Figure 12. Characteristics of the generated critical points for the §5.3 MNIST classifier experiment. In gray the classifier weights have been drawn uniformly in the Ω optimization space, while in purple the weights have been drawn using $I_{\psi}(\nabla u_{\theta}^*(0), \mathbf{z})$.



Figure 13. Performances of I_{ψ} for the §5.4 MNIST classifier experiment and stochasticity of the MNIST loss. The distance and cosine plots demonstrate the ability of I_{ψ} to correctly generate weights with a fixed gradient. For the stochasticity plot, the classifier weights are fixed to the weights obtained after gradient descent and different minibatches of MNIST images are used to compute the gradient of the loss function. The stochasticity plot shows the distribution of the sinkhorn divergence between two gradient batches computed from the same weights.



Figure 14. Characteristics of the generated critical points for the §5.4 MNIST classifier experiment. In gray the classifier weights have been drawn uniformly in the Ω optimization space, while in purple the weights have been drawn using $I_{\psi}(\nabla u_{\theta}^*(0), \mathbf{z})$.



Figure 15. Performances of I_{ψ} for the §5.5 sampling MNIST classifier experiment. The distance and cosine plots demonstrate the ability of I_{ψ} to correctly generate weights with a fixed gradient. For the stochasticity plot, the classifier weights are fixed to the final sampled particles and different minibatches of MNIST images are used to compute the gradient of the loss function. The stochasticity plot shows the distribution of the sinkhorn divergence between two gradient batches computed from the same weights.



Figure 16. Characteristics of the points sampled using Algorithm 2 for the \$5.5 MNIST classifier experiment. In gray the classifier weights have been drawn uniformly in the Ω optimization space, while in purple the weights have been sampled using Algorithm 2. The generated samples are critical points which are good minima, as shown by the accuracy statistics.

F. Hyperparameters

F.1. Parameterize u_{θ}

In all experiments, the convex function u is parameterized using an ICNN u_{θ} whose architecture is detailed in §3.1. The rank of the quadratic term $Q_{A,\delta}(x)$ is always taken equal to 1, which means that A is a row matrix. We noted that it was necessary to choose smooth activation functions in u's parameterization to avoid convergence problems with the conjugate solvers, that occur especially in high dimension. This is why we have favored the use of ELU (Clevert et al., 2015) activations in the u parameterization rather than Relu activations.

F.2. Computation of u_{θ}^*

The use of a conjugate solver is necessary to compute the loss functions of V_{ϕ} , M_{ξ} , I_{ψ} and to estimate M_{θ} . In all cases, the objective is to estimate the gradient of u_{θ} 's conjugate at a given point y: $\nabla(u_{\theta})^*(y)$. For a given experiment, that justifies the use of the same conjugate solver parameters for these different applications. We relied on ADAM solver for the computation of the convex conjugate as it runs faster than LBFGS on our examples and use Amos implementation. The two hyperparameters that remain to be set are the maximum number of iterations given to the solver to converge and the tolerance factor at which the norm of the gradient is considered small enough for the solver to have converged. These two hyperparameters are strongly dependent on the dimension of the problem as well as on the function u_{θ} and, therefore, on the distributions ρ and $F_{\#}\rho$. To amortize the number of iterations required for the solver to converge, it is always initialized with the prediction of the V_{ϕ} network that is trained in conjunction with u_{θ} .

F.3. Parameterize V_{ϕ}

We use an MLP with 2 hidden layers of size 512 and Relu activation functions to parameterize V_{ϕ} in all our experiments.

F.4. Parameterize M_{ξ}

The measure-preserving map M is parameterized by a neural network only when the vector field under study is available through samples only. This is the case in the topography examples where M_{ξ} is parameterized by an MLP with 2 hidden layers of size 512 and Relu activation functions.

F.5. Parameterize X_{ψ}

The learned part of the drift X_{ψ} is parameterized using an MLP, and we use Silu activation functions which is the classic choice for parameterizing the drift X_{ψ} .

The same hyperparameters have been used for the Chamonix, London, and Cyprus cases.

model	hyperparameter	value
$u_{ heta}$	activation function architecture b1 b2 scheduler initial learning rate α scheduler steps training steps steps	elu [64, 64, 64, 64] 0.50 0.50 cosine decay 0.001 0.10 50000 dual step ICNN 50000
I_ψ	activation function architecture scheduler initial learning rate α scheduler steps training steps steps σ	silu [256, 256, 256] cosine decay 0.001 0.010 50001 flow matching 50000 0.10
V_{ϕ}	activation function architecture b1 b2 scheduler initial learning rate α scheduler steps steps	relu [512, 512] 0.90 0.999 cosine decay 0.0005 0.010 51001 50001
conjugate solver	name max iteration gtol	Adam 200 0.0010

Figure 17. Hyperparameters used for the topography experiments (the same hyperparameters have been used for Chamonix, London, and Cyprus).

model	hyperparameter	value
$u_{ heta}$	activation function architecture b1	elu [128, 128, 128, 128] 0.50
	b2	0.50
	scheduler	cosine decays
	initial learning rate	0.001
	α	0.010000
	scheduler steps	10000
	training steps	dual step ICNN
	steps	10000
I_{ψ}	activation function	silu
	architecture	[512, 512]
	scheduler	cosine decay
	initial learning rate	0.0005
	α	0.010
	scheduler steps	4000°
	training steps	flow matching
	steps	50000
	σ	1.0
V_{ϕ}	activation function	relu
	architecture	[512, 512]
	b1	0.90
	b2	0.999
	scheduler	cosine decay
	initial learning rate	0.0005
	α	0.010
	scheduler steps	11000
	steps	10001
conjugate solver	name	Adam
	max iterations	700

Figure 18. Hyperparameters used for experiment 6.3.

model	hyperparameter	value
$u_{ heta}$	activation function architecture b1	elu [128, 128, 128, 128] 0.50
	b2	0.50
	scheduler	cosine decay
	initial learning rate	0.0001
	α	0.10
	scheduler steps	30000
	training steps	dual step ICNN
	steps	1002
I_{ψ}	activation function	silu
	architecture	[512, 512]
	scheduler	cosine decay
	initial learning rate	0.0005
	α	0
	scheduler steps	61003
	training steps	flow matching
	σ	0.10
V_{ϕ}	activation function	relu
,	architecture	[512, 512]
	b1	0.90
	b2	0.999
	scheduler	cosine decay
	initial learning rate	0.0005
	α	0
	scheduler steps	62005
	steps	1002
conjugate solver	name	Adam
	max iteration	1000
	gtol	0.001
particles	steps	60000
	coefficient LMC f	1
	coefficient LMC u	1
	particules	1024
	warming steps	30000
	consecutive LMC steps on f	200
	consecutive LMC steps on u	200
	$ au_f$	0.10
	$ au_u$	0.10

Figure 19. Hyperparameters used for experiment 6.4.

model	hyperparameter	value
$u_{ heta}$	activation function architecture b1 b2 scheduler initial learning rate α scheduler steps training steps steps	elu [128, 128, 128, 128] 0.50 0.50 cosine decay 0.0001 0.10 30000 dual step 1002
I_{ψ}	activation function architecture scheduler initial learning rate α scheduler steps training steps σ	silu [512, 512] cosine decay 0.000500 0 61003 flow matching 0.10
V_{ϕ}	activation function architecture b1 b2 scheduler initial learning rate α scheduler steps steps	relu [512, 512] 0.90 0.999 cosine decay 0.0005 0 62005 1002
conjugate solver	name max iteration gtol	Adam 1000 0.001
particles	steps coefficient LMC f coefficient LMC u particules warming steps LMC steps on f LMC steps on u τ_f τ_u	60000 1000 1000 1024 30000 200 200 0.10 0.10

Figure 20. Hyperparameters used for experiment 6.5.