Geometric data analysis, beyond convolutions

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Joint work with B. Charlier, J. Glaunès (numerical foundations), T. Séjourné, F.-X. Vialard, G. Peyré (optimal transport theory), P. Roussillon, P. Gori, A. Trouvé (applications to computational anatomy), F. Sverrisson, B. E. Correia, M. Bronstein (applications to protein sciences). 2012–2016 ENS Paris, mathematics and applications.

2015 MVA thesis with **Siemens Healthcare** in Princeton.

2016–2019 PhD thesis with Alain Trouvé, **computational anatomy**; TA/tutor in applied maths at the ENS Paris.

2019–2022 PostDoc with Michael Bronstein, geometric deep learning.

Family of medical doctors (radiologist, haematologist, GPs...): strong motivation to work towards **clinical solutions**.

Make life easier for engineers and researchers in the field: two libraries (KeOps, GeomLoss) to speed up geometric methods, with new guarantees of robustness.

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Today, we will talk about:

- 1. KeOps: fast geometry with **symbolic matrices**.
- 2. Applications to machine learning, proteins, maths...
- 3. GeomLoss: fast, robust and scalable optimal transport.
- 4. Scientific context, future works.

Symbolic matrices?



Dense matrix Coefficients only

Dense matrices – large, contiguous arrays of numbers:

- + Convenient and well supported.
- Heavy load on the memories of our GPUs, with time-consuming transfers that take place between compute units.



Dense matrixSparse matrixCoefficients onlyCoordinates + coeffs

Sparse matrices – tensors that have few non-zero entries:

- + Represent large tensors with a small memory footprint.
- Outside of graph processing, few objects are sparse enough to really benefit from this representation.

Machine learning libraries represent most objects as tensors



Distance and kernel matrices, point convolutions, attention layers:

- + Linear memory usage: no more memory overflows.
- + We can optimize the use of registers for a $\times 10 \times 100$ speed-up vs. a standard PyTorch GPU baseline.

Our library comes with all the perks of a deep learning toolbox:

- + Transparent array-like interface.
- + Full support for automatic differentiation.
- + Comprehensive collection of **tutorials**, available online.

Under the hood: combines an optimized C++ engine with high-level binders for PyTorch, NumPy, Matlab and R (thanks to Ghislain Durif). (We welcome contributors for JAX, Julia and other frameworks!)

To get started: ⇒ pip install pykeops ← www.kernel-operations.io

A first example: efficient nearest neighbor search in dimension 50

Create large point clouds using standard PyTorch syntax:

import torch

```
N, M, D = 10**6, 10**6, 50
x = torch.rand(N, 1, D).cuda() # (1M, 1, 50) array
y = torch.rand(1, M, D).cuda() # ( 1, 1M, 50) array
```

Turn dense arrays into symbolic matrices:

from pykeops.torch import LazyTensor
x_i, y_j = LazyTensor(x), LazyTensor(y)

Create a large **symbolic matrix** of squared distances:

D_ij = ((x_i - y_j)**2).sum(dim=2) # (1M, 1M) symbolic

Use an .argmin() reduction to perform a nearest neighbor query: indices_i = D_ij.argmin(dim=1) # -> standard torch tensor

The KeOps library combines performance with flexibility

Script of the previous slide = efficient nearest neighbor query, on par with the bruteforce CUDA scheme of the FAISS library... And can be used with any metric!

KeOps supports arbitrary formulas and variables with:

- Reductions: sum, log-sum-exp, K-min, matrix-vector product, etc.
- **Operations:** +, ×, sqrt, exp, neural networks, etc.
- Advanced schemes: batch processing, block sparsity, etc.
- Automatic differentiation: seamless integration with PyTorch.

KeOps lets users work with millions of points at a time

Benchmark of a matrix-vector product with a N-by-N Gaussian kernel matrix between 3D point clouds.



Applications

KeOps is a good fit for machine learning research





K-Means.

Gaussian Mixture Model.

Use any kernel, metric or formula you like! \implies More tutorials coming up soon.

KeOps is a good fit for machine learning research





Spectral analysis.

UMAP in hyperbolic space.

Use any kernel, metric or formula you like! \implies More tutorials coming up soon.

Applications to Kriging, spline, Gaussian process, kernel regression

A standard tool for regression [Lec18]:



Under the hood, solve a kernel linear system:

 $(\lambda \operatorname{Id} + K_{xx}) a = b$ i.e. $a \leftarrow (\lambda \operatorname{Id} + K_{xx})^{-1} b$

where $\lambda \ge 0$ and $(K_{xx})_{i,j} = k(x_i, x_j)$ is a positive definite matrix.

KeOps symbolic tensors:

- Can be fed to **standard solvers**: SciPy, GPytorch, etc.
- GPytorch on the 3DRoad dataset (N = 278k, D = 3): 7h with 8 GPUs \rightarrow 15mn with 1 GPU.
- Provide a fast backend for research codes: see e.g.
 Kernel methods through the roof: handling billions of points efficiently, by G. Meanti, L. Carratino, L. Rosasco, A. Rudi (2020).

Geometric deep learning w. F. Sverrisson, B. Correia and M. Bronstein

Data-driven methods on **point clouds** and **proteins**:

- + Fast K-NN search: local interactions.
- + Fast N-by-N computations: global interactions.
- + Heterogeneous **batches**, Octree-like acceleration.







Quasi-geodesic **convolutions**.

Applications to protein sciences [SFCB20]



(a) Raw protein data.



(b) Interface.



(c) Prediction.



(d) Chem. 1.

(e) Chem. 2.





(g) H at 10 Å.

Fast end-to-end learning on protein surfaces





 $\times 100$ - $\times 1,000$ faster, lighter and fully differentiable.

Some applications to **dynamical systems** [DM08, DFMAT17] and **statistics** [CDF19] with A. Diez, G. Clarté and P. Degond:



3D Vicsek model with orientation, interactive demo with 2k **flyers**.



2D Vicsek model on the torus, in real-time with 100k **swimmers**.

\implies Scale up to millions/billions of agents with Python scripts.





Packing problem in 2D with 10k repulsive balls.

Collective Monte Carlo **sampling** on the hyperbolic Poincaré disk.

Fast, scalable and robust optimal transport solvers











$$OT(\boldsymbol{\alpha}, \boldsymbol{\beta}) = \frac{1}{2N} \sum_{i=1}^{N} |\mathbf{x}_i - \mathbf{y}_{\sigma^*(i)}|^2$$



$$OT(\alpha, \beta) = \frac{1}{2N} \sum_{i=1}^{N} |\mathbf{x}_{i} - y_{\sigma^{*}(i)}|^{2} = \min_{\sigma \in S_{N}} \frac{1}{2N} \sum_{i=1}^{N} |\mathbf{x}_{i} - y_{\sigma(i)}|^{2}$$

Optimal transport generalizes sorting to ${\sf D}>1$



Minimize over N-by-M matrices (transport plans) π :

$$OT(\boldsymbol{\alpha}, \boldsymbol{\beta}) = \min_{\pi} \underbrace{\sum_{i,j} \pi_{i,j} \cdot \frac{1}{2} |\mathbf{x}_i - \mathbf{y}_j|^2}_{\text{transport cost}}$$



subject to $\pi_{i,j} \ge 0$, $\sum_{j} \pi_{i,j} = \alpha_{i}, \sum_{i} \pi_{i,j} = \beta_{j}.$

Key properties [Bre91]

The Wasserstein loss $OT(\alpha, \beta)$ is:

- Symmetric: $OT(\alpha, \beta) = OT(\beta, \alpha)$.
- Positive: $OT(\alpha, \beta) \ge 0$.
- Definite: $\operatorname{OT}(\alpha,\beta) = 0 \Longleftrightarrow \alpha = \beta$.
- Translation-aware: $OT(\alpha, Translate_{\vec{v}}(\alpha)) = \frac{1}{2} \|\vec{v}\|^2$.
- More generally, OT retrieves the unique gradient of a convex function T = ∇φ that maps α onto β:
 - $\begin{array}{ll} \text{In dimension 1,} & (\textbf{x}_i \textbf{x}_j) \cdot (\textbf{y}_{\sigma(i)} \textbf{y}_{\sigma(j)}) & \geqslant 0 \\ \text{In dimension D,} & \langle \textbf{x}_i \textbf{x}_j \ , \ \textbf{T}(\textbf{x}_i) \textbf{T}(\textbf{x}_j) \rangle_{\mathbb{R}^D} \ \geqslant \ 0 \ . \end{array}$

 \implies Appealing generalization of an **increasing mapping**.





$$t = .25$$



$$t = .50$$



$$t = 1.00$$



$$t = 5.00$$



$$t = 10.00$$

Robust optimal transport: softening the bijectivity constraints

Standard OT: minimize over N-by-M transport plans π ,

$$OT(\boldsymbol{\alpha}, \boldsymbol{\beta}) = \min_{\boldsymbol{\pi}} \left\langle \frac{1}{2} | \mathbf{x}_{i} - \mathbf{y}_{j} |^{2}, \boldsymbol{\pi} \right\rangle$$

s.t. $\boldsymbol{\pi} \ge 0$, $\boldsymbol{\pi} \mathbf{1} = \boldsymbol{\alpha}$, $\boldsymbol{\pi}^{\mathsf{T}} \mathbf{1} = \boldsymbol{\beta}$

When dealing with real-life data, we'd rather work with:

$$OT_{\sigma,\rho}(\boldsymbol{\alpha},\boldsymbol{\beta}) = \min_{\pi} \langle \frac{1}{2} | \mathbf{x}_{i} - \mathbf{y}_{j} |^{2}, \pi \rangle$$

+ $\underbrace{\sigma^{2} \operatorname{KL}(\pi \mid \boldsymbol{\alpha} \otimes \boldsymbol{\beta})}_{\pi \text{ is fuzzy at scale } \sigma} + \underbrace{\rho^{2} \operatorname{D}(\pi \mathbf{1} \mid \boldsymbol{\alpha}) + \rho^{2} \operatorname{D}(\pi^{\mathsf{T}} \mathbf{1} \mid \boldsymbol{\beta})}_{\pi \text{ tries to match } \boldsymbol{\alpha} \text{ with } \beta \dots \text{ up to a distance } \rho}$

In the formula above:

- KL is the relative entropy.
- D may be the relative entropy, the total variation, etc.

Robust optimal transport: fast algorithms, with guarantees

We define the Sinkhorn divergence:

$$\mathsf{S}_{\sigma,\rho}(\boldsymbol{\alpha},\boldsymbol{\beta}) = \mathsf{OT}_{\sigma,\rho}(\boldsymbol{\alpha},\boldsymbol{\beta}) - \frac{1}{2}\mathsf{OT}_{\sigma,\rho}(\boldsymbol{\alpha},\boldsymbol{\alpha}) - \frac{1}{2}\mathsf{OT}_{\sigma,\rho}(\boldsymbol{\beta},\boldsymbol{\beta})$$

$$\simeq$$
 OT"lazy- ρ "($k_{\sigma} \star \alpha$, $k_{\sigma} \star \beta$),

where k_{σ} is a Gaussian kernel of deviation σ and our "lazy" particles do not move beyond a distance ρ .

Theorem 1 (geometry): $S_{\sigma,\rho}$ is suitable for gradient descent. It is **positive**, definite, **convex** and metrizes the convergence in law.

Theorem 2 (algorithm): We can **implement** $S_{\sigma,\rho}$ efficiently, **on GPUs**. Two main ingredients: **log-convolution** with the Gaussian kernel k_{σ} and a **proximal operator** that is related to $\rho^2 D(\cdot | \cdot)$. Key dates for discrete optimal transport with N points:

- [Kan42]: Dual problem.
- [Kuh55]: Hungarian method in $O(N^3)$.
- [Ber79]: Auction algorithm in $O(N^2)$.
- [KY94]: **SoftAssign** = Sinkhorn + annealing, in $O(N^2)$.
- [GRL+98, CR00]: Robust Point Matching = Sinkhorn as a loss.
- [Cut13]: Start of the GPU era.
- [Mér11, Lév15, Sch19]: Multiscale solvers in $O(N \log N)$.
- Today: Multiscale Sinkhorn algorithm, on the GPU.

 \implies Generalized **QuickSort** algorithm.

Progresses of the last decade add up to a $\times 100 \cdot \times 1000$ acceleration: Sinkhorn GPU $\xrightarrow{\times 10}$ + KeOps $\xrightarrow{\times 10}$ + Annealing $\xrightarrow{\times 10}$ + Multiscale

With a precision of 1%, on a modern gaming GPU:



10k points in 30-50ms

100k points in 100-200ms

Our website: www.kernel-operations.io/geomloss

 \Rightarrow pip install geomloss \Leftarrow

```
# Large point clouds in [0,1]<sup>3</sup>
import torch
x = torch.rand(100000, 3, requires_grad=True).cuda()
y = torch.rand(200000, 3).cuda()
# Define a Wasserstein loss between sampled measures
```

from geomloss import SamplesLoss
loss = SamplesLoss(loss="sinkhorn", p=2)
L = loss(x, y) # By default, use constant weights

Soon: efficient support for images, meshes and generic metrics.

Ongoing work: computational anatomy





Fast OT-based registration with S. Joutard, X. Hao, A. Young from KCL, Z. Shen, M. Niethammer from UNC. Diffeomorphic and spline registration e.g. Deformetrica LDDMM software with the Aramis Inria team.

Scientific context, future works

Genuine team work



Alain Trouvé



Thibault Séjourné

F.-X. Vialard



Gabriel Peyré



Benjamin Charlier

Joan Glaunès



Pierre Roussillon



Pietro Gori

+ Freyr Sverrisson, Bruno Correia, Michael Bronstein, ...

Promoting cross-field interactions



Promoting cross-field interactions



The emergence of an open and **modular** ecosystem of scientific tools has been a **boon** to the community.

Deep learning frameworks have put **GPU computing** and **automatic differentiation** in the hands of every student. (Incredible!)

These libraries have attracted significant backing from **industry** players (Google, Facebook, ...) and allowed the field to **boom** over the last decade.

Interacting with other researchers, doctors and engineers has never been so **easy**.

But on the other hand, PyTorch and TensorFlow have also **biased** the field towards a **small set** of **well-supported** operations: convolutions and matrix-matrix products, mostly.

This design choice is **not** due to an intrinsic limitation of GPUs: our hardware is more than capable of **simulating** large, open **3D worlds** in real-time!

As academic researchers, we must strive to keep **other paths open**. Foster the development of a full range of methods, from **robust** convex baselines to **expressive** deep learning pipelines. KeOps and GeomLoss are:

- + Fast: $\times 10 \times 1,000$ speedup vs. naive GPU implementations.
- + Memory-efficient: O(N), not $O(N^2)$.
- + Versatile, with a transparent interface: freedom!
- + Powerful and well-documented: research-friendly.
- $-\,$ Slow with large vectors of dimension D > 100.

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First half of 2021:

- ightarrow Approximation strategies (Nyström, etc.) in KeOps.
- $\rightarrow\,$ Wasserstein <code>barycenters</code> and <code>grid</code> images in GeomLoss.

Roadmap for KeOps + GeomLoss:

- 2017–18 **Proof of concept** with conference papers, online codes. Get first feedback from the community.
- 2019–20 **Stable library** with solid theorems, a well-documented API. KeOps backends for high-level packages.
- 2021–22 **Mature library** with focused application papers, full tutorials. Works out-of-the-box for students and engineers.
 - 2022+ A standard toolbox, with genuine clinical applications? That's the target!

Conclusion

Key points

- Symbolic matrices are to geometric ML what sparse matrices are to graph processing:
 - $\longrightarrow~$ KeOps, x30 speed-up vs. PyTorch, TF and JAX.
 - $\longrightarrow~$ Useful in a wide range of settings.
- Optimal Transport = generalized sorting:
 - \longrightarrow Geometric gradients.
 - \longrightarrow Super-fast $O(N \log N)$ solvers.
- These tools open **new paths** for geometers and statisticians:
 - \longrightarrow GPUs are more **versatile** than you think.
 - $\longrightarrow~$ Ongoing work to provide fast GPU backends to researchers
 - $-\operatorname{going}$ beyond what Google and Facebook are ready to pay for.

Conclusion

We believe that KeOps and GeomLoss will stimulate research on:

- Clustering methods: fast K-Means and EM iterations.
- Data representation: UMAP, fast KNN graphs with any metric.
- Kernel methods: kernel matrices.
- Gaussian processes: covariance matrices.
- Geometric deep learning: point convolutions.
- Medical imaging: computational anatomy.
- Geometric statistics: going beyond Euclidean models.
- Natural language processing: transformer networks?

What do you think?

Documentation and tutorials are available online



www.jeanfeydy.com/geometric_data_analysis.pdf 35

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