Geometric data analysis

Lecture 2/7 – Flat vector spaces

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Thursday, 9am-12pm - 7 lectures

Faculté de médecine, Hôpital Cochin, rooms 2001 + 2005

Validation: project + quizz

Remember this slide from lecture 1?



Supervised learning = Regression.

We look for a formula $F(x_1, ..., x_D)$ of the D variables that best approximates an important quantity (\heartsuit).

First thing you should do?



Working with **clients** < **colleagues** < **friends**.

Data **science** is never done in a vacuum. Our (big) spreadsheets are **partial projections** of a complex reality.

> What are we trying to achieve? What type of information is available? What do we already know?

To understand this **context**, you must break the ice with domain experts.

This is a continuous, time-consuming and enjoyable process.

Today: well-rounded methods for high-quality features

1. Decision trees - for heterogeneous data

• Greedy training and regularizations.

2. K-Nearest Neighbors – a first isotropic method

• Euclidean metrics and normalization.

3. Linear regression – to estimate global trends

• Linear, piecewise linear and polynomial regression.

4. Kernel methods - specify a custom prior

- · Smoothness, short- and long-range interactions.
- Nadaraya–Watson–Shepard and Ridge regressions.

Expert knowledge is often distilled as a tree



Transformation of the national breast cancer guideline into data-driven clinical decision trees, Hendriks et al., 2019

Tree models are easy to train with a greedy algorithm



Recursive splits that stop if improvements < T \iff **greedy** minimization of $\operatorname{Fit}_{x,y}(F) + \operatorname{Reg}(F) = \frac{1}{2} \sum_{i} \|F(x_i) - y_i\|^2 + T \cdot \#\operatorname{Leaves}(F)$.

Two toy regression problems



D=1 – 9 points **x** on the unit interval [0, 1]. **D=2** – 20 points **x** on the unit square $[0, 1]^2$.

In both cases: **scalar** output values **y**.





Depth 0: 1 constant value.





Depth 1:

2 distinct values. The model is **piecewise constant**.





Depth 2:

up to 4 distinct values. The model follows the **D axes** of the feature space.





Depth 3:

up to 8 distinct values. We may choose not to use them all to limit the **complexity** of the model.





Depth 4:

up to 16 distinct values. Starting to clearly **overfit**.





Depth 5:

up to 32 distinct values. Starting to clearly **overfit**.





Depth 10:

up to 1,024 distinct values. Full **overfit** on both datasets.



Decision trees are:

- Interpretable.
- Easy to train and deploy.
- Fast and CPU-friendly.
- Robust:
 - Only use a few columns at a time.
 - Work well with **heterogenous** information.
 - Only rely on the **ordering** of the features.

However, trees also **overfit** quickly and produce **blocky** results. Regularization methods mitigate these issues, at the cost of **interpretability**.



Iterative fits on the prediction residuals with shallow trees. Use a **small** learning rate for better regularization: $\label{eq:Residual_i} \texttt{Residual_i} = \texttt{y}_i - \textbf{0.1} \cdot \sum_k \texttt{Tree}_k(\texttt{x}_i) \; .$



1 tree of depth 3: a simple decision tree, with moderate complexity.





3 trees of depth 3: sum of three simple decision trees, fitted iteratively on **residuals**.





5 trees of depth 3: sum of five simple decision trees, fitted iteratively on **residuals**.





100 trees of depth 3: sum of a hundred simple decision trees. We reach a high training accuracy with a relatively smooth model.





Parallel fits on **bootstrap** samples of the original dataset. The final model is the **average** of a **forest** of independent trees.



1 tree of depth 3: a simple decision tree, computed on a bootstrap subset of the original sample.





2 trees of depth 3: average of two decision trees, fitted on two independent bootstrap samples.





5 trees of depth 3: average of five decision trees, fitted on five independent bootstrap samples.





100 trees of depth 3: a regularized decision rule. The model still follows the **axes** of the feature space, but is **smoother**.



Some features may require more work - understand the context! [Wit]



The **Body Mass Index** = weight / height² is a good indicator for many health problems.

Some features may require more work - understand the context!



Applying thresholds on **postal codes** is mostly useless. Other statistics may be much more informative.

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2:31:28 PM

Applying thresholds on **UNIX timestamps** is mostly useless. We must first apply periodic transforms to get hours-days-months.

Sometimes, the input features are just not good enough [EPW11]



Tree models cannot process **raw pixel values**. Standard radiomic features only take you so far. Tree-based models are:

- Highly interpretable.
- Well suited to high-quality heterogeneous features.
- **Easy** to use: XGBoost, LightGBM, scikit-learn...

On the other hand, they produce **non-smooth** results and are biased along the **axes** of the feature space.

This is a major limitation if you work with **homogeneous** features: the 3D xyz coordinates, pixel values, audio signals...

K-Nearest Neighbors

The Euclidean metric is isotropic



Thresholding features promotes decisions along the **axes** of the feature space.



$$\begin{split} \text{The squared } \textbf{Euclidean} \ \text{metric} \\ \|(x_1,\ldots,x_D)\|^2 \ = \ x_1^2 + \cdots + x_D^2 \\ \text{is invariant to } \textbf{rotations}. \end{split}$$

Average value among the K-Nearest Neighbors



With **K** = **1** neighbor, we retrieve a simple nearest neighbor interpolation. This model is piecewise constant on the **Voronoi diagram** of x.



Average value among the K-Nearest Neighbors



With **K = 2** neighbors, the cells of the diagram become smaller.


Average value among the K-Nearest Neighbors



With **K = 3** neighbors, the cells of the diagram become smaller.



Average value among the K-Nearest Neighbors



With **K = 4** neighbors, the cells of the diagram become smaller.



Average value among the K-Nearest Neighbors



With **K = 5** neighbors, the model looks **smoother** and smoother but is still **piecewise constant**.



K-NN models are:

- Interpretable.
- Isotropic which may or may not be a good thing!
- **Easy** to deploy.
- Fast, parallel and GPU-friendly see our MVA Lecture 7 on algorithms.
- Well-packaged and scalable: FAISS, KeOps, (big-)ann-benchmarks.com...

Major weakness: K-NNs require a good scaling of the input features

Unlike tree-based models, the Euclidean distance is sensitive to the **precise values** of the features x.

Out-of-the-box, K-NNs are not even robust to the **choice of the units** for the columns of our dataset!

We must **normalize** the input features using:

- A feature-wise rescaling using e.g. the **standard deviation**.
- A **multivariate** normalization using e.g. Principal Component Analysis. The Euclidean distance with a normalized PCA is known as the **Mahalanobis** metric.
- Alternatively, a robust equalization of the feature histograms.

Tree-based and K-NN models are:

- Interpretable methods with heterogeneous / homogeneous features.
- Well-understood, well-packaged and easy to deploy.
- Excellent baselines for interpolation.

Unfortunately, both methods :

- Produce non-smooth, piecewise constant decision rules.
- Are local and do not estimate global trends.
 They are not a natural fit for extrapolation, forecasting.

Linear regression

A simple model: linear regression



We choose the weights **a**, **b**, ..., **f** by minimizing a least squares error.





































Linear regression



Linear regression models a **monotonic trend**.

It cannot handle complex relationships between the input x and the ouput y.

What should we do if the problem is complex?



Take a break :-)

Neural networks

Maybe, we could introduce some intermediate variables in our model?



Domain experts may have suggested a **step-by-step** process to compute the quantity of interest – say, the perimeter of an organ.

Let's complexify our model with intermediate variables...



Let's complexify our model with intermediate variables...










Let's complexify our model with intermediate variables... and non-linearities!































MLP with **1 hidden neuron**. This is a piecewise linear model with at most 1 hinge. The optimizer doesn't use them all.





MLP with **10 hidden neurons**. This is a piecewise linear model with at most 10 hinges. The optimizer doesn't use them all.





MLP with **20 hidden neurons**. This is a piecewise linear model with at most 20 hinges. The optimizer doesn't use them all.

0 0 0 C 0 0 0 0



MLP with **50 hidden neurons**. This is a piecewise linear model with at most 50 hinges. The optimizer doesn't use them all.

0 0 0 C 0 0 0 0 0



MLP with **100 hidden neurons**. This is a piecewise linear model with at most 100 hinges. The optimizer doesn't use them all.

0 0 0 C 0 \cap 0 0 0 0



Deeper MLP with 2 hidden layers and **100 + 100 hidden neurons**, i.e. at most 100 x 100 hinges. The **non-convex, stochastic** optimization is **unreliable** and not reproducible.

0 0 0 0 0 0



Deeper MLP with 3 hidden layers and **100 + 100 + 100 hidden neurons**, i.e. at most 100 x 100 x 100 hinges. The **non-convex, stochastic** optimization is **unreliable** and not reproducible.

0 0 0 0 0 0 0



Deeper MLP with 4 hidden layers and **100 + 100 + 100 + 100 hidden neurons**, i.e. at most 100 x 100 x 100 x 100 hinges.

Starting to look like a smooth origami ;-)

0 0 0 0 0 C 0

(Vanilla, fully connected) neural networks - strengths and weaknesses

- Modular and easy to extend.
- Simplest way of implementing high-dimensional piecewise linear models.
- Extremely well-supported on CPU and GPU: PyTorch, TensorFlow...

Unfortunately, the optimization of the "neural" weights corresponds to a **non-convex** optimization problem.

We must rely on **non-deterministic**, stochastic solvers. Performance and smoothness are **not** simply correlated to the number of neurons and layers.

In most applications, this lack of reproducibility and interpretability is a **deal-breaker**.



Constant polynomials of **degree 0**: **D=1** – 1 constant. **D=2** – 1 constant.





Linear polynomials of **degree 1**: **D=1** – 1, x. **D=2** – 1, x, y.





Quadratic polynomials of **degree 2**: $D=1-1, x, x^2$. $D=2-1, x, y, x^2, xy, y^2$.



Cubic polynomials of **degree 3**: **D=1** – 1, x, x², x³. **D=2** – 1, x, y, x², xy, y², x³, x²y, xy², y³.





Quartic polynomials of **degree 4**: **D=1** – 1, x, x², x³, x⁴. **D=2** – 1, x, y, x², xy, y², x³, x²y, xy², y³, $x^4, x^3y, x^2y^2, xy^3, y^4$. Starting to **overfit** in dimension D=2.





Polynomials of **degree 5**: **D=1** – 1, x, x², x³, x⁴, x⁵. **D=2** – 1, x, y, x², xy, y², x³, x²y, xy², y³, x⁴, x³y, ..., y⁴, x⁵, x⁴y, ..., y⁵. Full **overfit** in dimension D=2.





Polynomials of **degree 10**: **D=1** – 1, x, x^2 , x^3 , x^4 , x^5 , ..., x^{10} . **D=2** – 1, x, y, x^2 , xy, y^2 , x^3 , x^2y , xy^2 , y^3 , x^4 , x^3y , ..., y^4 , x^5 , x^4y , ..., y^5 , ..., y^{10} . Full **overfit** in both examples.



"Non-parametric" methods:

- Tree-based models robust, but with a bias along the axes.
- K-Nearest Neighbors models isotropic, but requires a good scaling.

"Parametric" methods:

- Linear regression useful, but often too simplistic.
- Neural networks expressive, but unreliable.

Polynomial regression:

- Linear regression with polynomial features.
- Quadratic regression is fine but we badly **overfit** beyond degree 4-5.

Kernel interpolation

Let's specify directly a linear parametric form for the model:

$$F(a_1,\ldots,a_J;x) \ = \ a_1F_1(x) \ + \ \cdots \ + \ a_JF_J(x).$$

In practice, we often use:

$$F(a_j;x) \;=\; \sum_j a_j \, k(x-x_j)$$

and say that k(x - y) is the **kernel** of our method.

Some common kernels



Two main criteria: is the kernel **smooth** or **peaky**? Does the kernel have **compact support** or a **heavy tail**? First method – just use a fraction instead of a linear combination:

$$F(\mathbf{x}) = \frac{\sum_{j} \mathbf{k}(\mathbf{x} - \mathbf{x}_{j}) \mathbf{y}_{j}}{\sum_{j} \mathbf{k}(\mathbf{x} - \mathbf{x}_{j})}$$

The Nadaraya–Watson method assumes that k(x - y) takes positive values.

It corresponds to a **barycentric interpolation** between the values y_j , with weights that are proportional to $k(x - x_j)$.


$$\begin{split} \textbf{Smooth, local Gaussian kernel with } \boldsymbol{\sigma} &= \textbf{0.2} \\ k(x,y) &= exp(-\|x-y\|^2/2\sigma^2). \end{split}$$

Smooth local averaging on the unit interval and square.





Smooth, local Gaussian kernel with σ = 0.1 $k(x,y) = exp(-\|x-y\|^2/2\sigma^2).$

Sharper, K-NN-like decision boundaries.





Heavy-tail Cauchy kernel with $\sigma = 0.1$ $k(x, y) = 1 / (1 + ||x - y||^2 / \sigma^2).$

Dampened towards the global average.





Pointy exponential kernel with $\boldsymbol{\sigma}$ = 0.1 $k(x,y) = exp(-\|x-y\|/\sigma). \label{eq:k}$

Closer fit to the training data.





 $\label{eq:singular} \begin{array}{l} \textbf{Singular} \ \text{Shepard kernel} \\ k(x,y) = 1 \, / \, \|x-y\|. \end{array}$

Perfect fit to the training data.



Singular and heavy-tail Shepard kernel $k(x,y) = 1 \, / \, \sqrt{\|x-y\|}.$

Perfect fit to the training data, dampening to the average value elsewhere.





Highly singular Shepard kernel $k(x, y) = 1 / ||x - y||^2.$

Perfect fit to the training data, close to a **linear** interpolation.



Very highly singular Shepard kernel $k(x,y) = 1 \, / \, \|x-y\|^4.$

Perfect fit to the training data, close to a **nearest neighbor** interpolation.

Second method: solve a linear system

$$\begin{split} \mathsf{F}(\mathsf{x}_1) \ &= \ \mathbf{a}_1 \, \varphi(\mathsf{x}_1 - \mathsf{x}_1) \, + \, \cdots \, + \, \mathbf{a}_N \, \varphi(\mathsf{x}_1 - \mathsf{x}_N) \, \simeq \, \mathsf{y}_1 \\ \mathsf{F}(\mathsf{x}_2) \ &= \ \mathbf{a}_1 \, \varphi(\mathsf{x}_2 - \mathsf{x}_1) \, + \, \cdots \, + \, \mathbf{a}_N \, \varphi(\mathsf{x}_2 - \mathsf{x}_N) \, \simeq \, \mathsf{y}_2 \\ &\vdots \ &= \ \vdots \ &+ \, \ddots \, + \quad \vdots \ &\simeq \ \vdots \\ \mathsf{F}(\mathsf{x}_N) \ &= \ \mathbf{a}_1 \, \varphi(\mathsf{x}_N - \mathsf{x}_1) \, + \, \cdots \, + \, \mathbf{a}_N \, \varphi(\mathsf{x}_N - \mathsf{x}_N) \, \simeq \, \mathsf{y}_N \end{split}$$

Linear system $\Phi \mathbf{a} = \mathbf{y}$

Enforcing a **perfect fit** to the data may not be reasonable.

Instead, we target a trade-off between accuracy and smoothness:

$$\min_{\mathbf{a}} \|\Phi \mathbf{a} - \mathbf{y}\|^2 + \operatorname{Reg}(\mathbf{a}).$$

Popular regularization terms are **convex**:

- Ridge: $\alpha \|\mathbf{a}\|^2 = \alpha (\mathbf{a}_1^2 + \dots + \mathbf{a}_N^2).$
- Lasso: $\lambda \|\mathbf{a}\|_1 = \lambda (|\mathbf{a}_1| + \dots + |\mathbf{a}_N|).$
- Elastic Net: $\lambda \|\mathbf{a}\|_1 + \alpha \|\mathbf{a}\|^2$.

$$\begin{split} \min_{\mathbf{a}} \, \|\Phi \, \mathbf{a} - \mathbf{y}\|^2 \, + \, \alpha \, \|\mathbf{a}\|^2 \ &= \ (\mathbf{a}^\top \Phi^\top - \mathbf{y}^\top) (\Phi \mathbf{a} - \mathbf{y}) + \alpha \, \mathbf{a}^\top \mathbf{a} \\ &= \ \mathbf{a}^\top (\Phi^\top \Phi + \alpha \, \mathsf{Id}_{\mathsf{N}}) \, \mathbf{a} - 2 \, \mathbf{y}^\top \Phi \mathbf{a} + \mathbf{y}^\top \mathbf{y} \end{split}$$

$$\implies \mathbf{a} = (\Phi^{\top} \Phi + \alpha \operatorname{Id}_{\mathsf{N}})^{-1} \Phi^{\top} \mathbf{y}$$
$$= \Phi^{\top} (\Phi \Phi^{\top} + \alpha \operatorname{Id}_{\mathsf{N}})^{-1} \mathbf{y}$$

$$\implies \mathsf{F}(\mathsf{x}) \;=\; \Phi \,\mathsf{a} \;=\; \Phi \Phi^\top (\Phi \Phi^\top + \alpha \,\mathsf{Id}_\mathsf{N})^{-1}\,\mathsf{y}\,.$$

A fundamental object appears: the symmetric, positive, semidefinite **kernel matrix** $K = \Phi \Phi^{\top}$.

$$\mathsf{K} = \Phi \Phi^\top \quad \text{i.e.} \quad \mathsf{K}(\mathsf{x}_\mathsf{i},\mathsf{x}_\mathsf{j}) \ = \ \langle \ \Phi(\mathsf{x}_\mathsf{i}), \Phi(\mathsf{x}_\mathsf{j}) \ \rangle \ = \ \sum_{\mathsf{s}=\mathsf{1}}^\mathsf{N} \varphi(\mathsf{x}_\mathsf{i}-\mathsf{x}_\mathsf{s}) \ \varphi(\mathsf{x}_\mathsf{j}-\mathsf{x}_\mathsf{s})$$

This may be **expensive**: N terms for every coefficient of K.

Fortunately, we may use the continuous limit instead:

$$\mathsf{k}(\mathsf{x}_{\mathsf{i}},\mathsf{x}_{\mathsf{j}}) \; = \; \int_{\mathsf{x}} \varphi(\mathsf{x}_{\mathsf{i}}-\mathsf{x}) \, \varphi(\mathsf{x}_{\mathsf{j}}-\mathsf{x}) \, \mathsf{d}\mathsf{x}$$

This dot product between **two translated copies** of φ is often known in **closed form**.

We consider functions $k(x_i - x_j)$ that can be written as the previous integral for a suitable function φ .

Criterion: if the **Fourier** transform
$$\hat{k}(\omega)$$
 is real-valued and positive,
then $\widehat{\varphi}(\omega) = \sqrt{\hat{k}(\omega)}$ works.

Then, **kernel ridge regression** simply relies on the model: $F(x) = K (K + \alpha Id_N)^{-1} y.$

On GPUs, we may solve this linear system efficiently using:

- **KeOps** bruteforce methods scale to N = 1,000,000 in seconds.
- FalkonML approximate methods scale to N = 1,000,000,000 in hours.

Kriging, Gaussian process regression [Lec18]



Kernel ridge regression has a **rich history** in applied mathematics. It is especially popular in geostatistics to estimate smooth terrain models: the approximation parameter α controls the **nugget** effect.

This theory is also behind **Sobolev** norms and **Gaussian** processes... More about this in the MVA Lecture 6 on probability distributions!



Smooth, **global** Gaussian kernel with $\pmb{\sigma}$ = 1.0 $k(x,y) = exp(-\|x-y\|^2/2\sigma^2).$

Only models a **global** linear trend.





$$\begin{split} \textbf{Smooth} & \text{Gaussian kernel with } \boldsymbol{\sigma} = \textbf{0.5} \\ & k(x,y) = exp(-\|x-y\|^2/2\sigma^2). \end{split}$$

Starts to discern different regions.





$$\begin{split} \textbf{Smooth} \mbox{ Gaussian kernel with } \boldsymbol{\sigma} &= \textbf{0.2} \\ k(x,y) &= exp(-\|x-y\|^2/2\sigma^2). \end{split}$$

Well-suited to the **sampling density**.





Smooth, local Gaussian kernel with σ = 0.1 $k(x,y) = exp(-\|x-y\|^2/2\sigma^2).$

Overfits on individual sample values.





$$\label{eq:heavy-tail} \begin{split} \mbox{Heavy-tail} \mbox{Cauchy kernel with } \sigma \mbox{= 0.2} \\ k(x,y) &= 1 \,/ \, (1 + \|x-y\|^2 / \sigma^2). \end{split}$$

Extrapolates with more confidence.





$$\label{eq:pointy} \begin{split} \text{Pointy} \ & \text{exponential kernel with } \sigma \text{ = } 0.2 \\ & k(x,y) = exp(-\|x-y\|/\sigma). \end{split}$$

Closer fit to the training data.





Pointy, global distance kernel $\mathbf{k}(\mathbf{x},\mathbf{y}) = -\|\mathbf{x} - \mathbf{y}\|.$

Models both **local** and **global** trends. **Excellent parameter-free baseline.**



Conclusion

Numerous regression models... But what about the curse of dimensionality?



Decision tree.



Random forest.



Nearest neighbors.



Neural network.



Linear.



Cubic.



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Kernel.

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