# U-Bootstrap percolation : critical probability, exponential decay and applications, by Ivailo HARTARSKY

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ENS Lyon M2 Advanced Mathematics

March 2022

## Motivation and principles of compressive learning

#### What is the setup?

We are confronted with a dataset which comes in form of n d-dimensional vectors  $\{x_i\}_{i=1}^n$ . We would like to perform some kind of learning on it but we are scared of the complexity when n is huge.

### What is compressive learning?

The principle of compressive learning consists of compressing (sketching) the dataset before applying any learning techniques. The sketch consists of a single vector  $\tilde{z}$  which is constructed by transforming each vector of the dataset and averaging the results :

$$\tilde{z} = \frac{1}{n} \sum_{i=1}^{n} \Phi(x_i)$$

## Motivation and principles of compressive learning

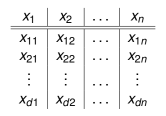
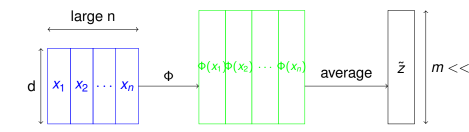


Table: Initial dataset



### Principal Component Analysis (PCA)

#### **PCA**

The goal is to find the linear subspace  $P_k$  that best fits the d-dimensional data  $\{x_i\}_{i=1}^N$  in the LS sense, i.e., find an orthogonal family of k vectors  $\{u_i\}_{i=1}^k$  that maximizes

$$\sum_{l=1}^{k} \sum_{i=1}^{N} |u_{l}^{T} x_{i}|^{2}.$$

A solution is the *k*-principal eigenvectors of the empirical autocorrelation matrix

$$\hat{R} = \frac{1}{N} \sum_{i=1}^{N} x_i x_i^T =: \frac{1}{N} \sum_{i=1}^{N} \Phi(x_i).$$

 $\hat{R}$  is a *sketch* of our data (of dim  $d^2$ ).

The sketch

$$\hat{R} = \frac{1}{N} \sum_{i=1}^{N} x_i x_i^T =: \frac{1}{N} \sum_{i=1}^{N} \Phi(x_i) \in \mathbb{R}^{d^2}$$

is a very compressed version of the data  $\{x_i\}_{i=1}^N$ , but, it still contains the geometry of the data.

#### CS inspired idea

Take m random measurements<sup>a</sup> of each sample and use the sketch defined by  $\Phi(x) = \mathcal{M}(xx^T)$ . Provided m > kd, the principal eigenvectors can be recovered.

 $<sup>{}^{</sup>a}\mathcal{M}: \mathbb{R}^{d \times d} \to \mathbb{R}^{m}$  satisfying RIP on matrices of rank at most 2k.

## Parallel with signal processing

- Recall that in signal processing our goal is to reconstruct a vector  $x \in \mathbb{R}^d$  from  $y = Ax + \epsilon$ , where y is its linear projection on a smaller subspace perturbed by an error
- At first glance compressive learning setup is rather different as we deal with a large collection of vectors, rather than just one
- The analogy becomes clearer if we assume (as it is often the case in ML) that our vectors  $\{x_i\}_{i=1}^n$  are modeled as i.i.d. random vectors having a probability measure  $\mathbb{P}$
- In this case we get

$$\lim_{n\to\infty}\frac{1}{n}\sum_{i=1}^n\mathbf{\Phi}(\mathbf{x}_i)\stackrel{a.s.}{=}\mathbb{E}_{\mathbb{P}}[\mathbf{\Phi}(X)]=\mathbb{A}(\mathbb{P}),$$

where  $\mathbb A$  is a linear operator matching a probability measure to the expectation over this measure of the feature map  $\Phi$ 

## Parallel with signal processing

In this manner we can write

$$\tilde{z} = \frac{1}{n} \sum_{i=1}^{n} \Phi(\mathbf{x}_i) \approx \mathbb{A}(\mathbb{P}) = \mathbb{A}(\mathbb{P}) + \epsilon$$

- Thus instead of considering a linear projection of a vector measured with noise as we did in signal processing, in compressive learning we consider a linear projection of the underlying probability measured with noise.
- If we take  $\Phi = x^k$  we get that  $\mathbb{E}_{\mathbb{P}}[\Phi(X)]$  is just the (uncentered) kth moment. In general, for any  $\Phi$  the quantity  $\mathbb{E}_{\mathbb{P}}[\Phi(X)]$  is called generalised moment.
- In signal processing the linear measurement matrix A can be chosen at random to ensure good reconstruction properties with high probability. By analogy, in compressive sensing Φ is also often randomised

#### k-means centroids

#### The problem

The goal is to recover k centroids  $\{c_l\}_{l=1}^k$  from some data  $\{x_i\}_{i=1}^N$  that minimize

$$\sum_{i=1}^{N} \min_{l} ||x_i - c_l||^2.$$

For N >> 1 traditional algorithms are not very efficient because they take the whole dataset at once...

But, N >> 1 allows to use the laws of large numbers and concentration. It is reasonable to consider that the data will accumulate on small portions of the space.

### The binning map

Assume the centroids are spaced by at least  $\varepsilon$  and have a norm smaller than r, then cover  $[-r,r]^d$  by,  $B=(\frac{2r}{\varepsilon})^d$ , d-dimensional cubes (*bins*). For each bin, count the average number of points that belong to it. This defines the binning map  $\hat{p} \in \mathbb{R}_+^B$ .

This gives us a sketch of the data, but in a large dimensional space.

*But*, if the model that generated the data is "structured", i.e., the data concentrates in a few centroids, then the problem is a *sparse* problem.

#### CS inspired idea

Use a Gaussian random matrix in  $\mathbb{R}^{m\times N}$  and define the sketch as  $\tilde{z}=A\hat{p}$  and solve

$$\tilde{p} = argmin_{p \in \Sigma_k^+} ||\tilde{z} - Ap||^2$$

## Sketching estimates the underlying data-distribution

We can do k-means clustering with m measures using:

- $m \ge k \log B$  for Gaussian sampling
- $m \ge k \log(k)^3 \log B$  for DFT of  $\hat{p}$

What if we consider the *continuous* Fourier Tranform (FT) ? We only know the empirical distribution  $\bar{p}_{\mathcal{X}} = \frac{1}{N} \sum_{i=1}^{N} \delta(x_i - x)$ , so

$$FT(\bar{p})(\omega) = \int_{\mathbb{R}^d} \bar{p}_{\mathcal{X}}(x) e^{-i2\pi \langle w, x \rangle} dx = \frac{1}{N} \sum_{i=1}^N e^{-i2\pi \langle w, x \rangle} =: \bar{\Psi}_{\bar{p}_{\mathcal{X}}}(\omega)$$
$$\longrightarrow^{\mathbb{E}} \bar{\Psi}_{p^*}(\omega) \quad \text{The "true" characteristic function at } \omega.$$

#### CS inspired idea

If the true distribution  $p^*$  is "simple", then interpolating the characteristic function, using simple models, on "few" of its samples should give the true distribution.

### Task driven distances

#### Task driven distance

Given a loss function L and probability distributions  $p_X$  and  $p_X^{'}$ , we consider the distance

$$\rho(p_X, p_X^{'}) = \sup_{\theta} |R^*(\theta|p_X) - R^*(\theta|p_X^{'})|$$

where  $R^*(\theta|p_X) = E_{X \sim p_X}(L(\theta)|X)$  is the expected risk under  $p_X$ .

- The loss function L is task specific
- Excess risk bounds  $0 \le R^*(\theta^{*'}|p_X) R^*(\theta^*|p_X) \le 2\rho(p_X, p_X^{'})$  where  $\theta^* = argmin_{\theta}R^*(\theta|p_X)$  and  $\theta^{*'} = argmin_{\theta}R^*(\theta|p_X^{'})$
- When  $\hat{p}_X$  is the empirical distribution on the data, and  $p_X$  is the true distribution, under certian conditions  $\rho(\hat{p}_X, p_X) = O(\frac{1}{\sqrt{n}})$

### LRIP and Excess risk control

In the compressive learning framework we are interested in an upper bound of the excess risk which is controlled by the task driven distance.

### The Lower Restricted Isometry Property (LRIP)

The operator A is said to have the LRIP with constant  $C_0$  and with respect to a parametric subfamily  $\Sigma_{\theta} = \{p_{\theta} | \theta \in \Theta\}$  if

$$\rho(p_{\theta}, p_{\theta'}) \leq C_0 ||\mathcal{A}(p_{\theta}) - \mathcal{A}(p_{\theta'})||$$

for all  $p_{\theta}, p_{\theta'} \in \Sigma_{\theta}$ 

Excess risk bound under LRIP: for all  $\theta' \in \Sigma_{\theta}$ 

$$R^*(\tilde{\theta}|p_X) - R^*(\theta'|p_X) \le 4C_0||A(p_X) - \tilde{z}|| + 4C_0||A(p_{\theta'}) - A(p_X)|| + 2\rho(p_{\theta'}, p_X)$$

• Choosing  $\theta' = \theta^*$ , this result is interpretable in terms of modeling and sampling error

### Expected and mean kernel, MMD

- Two sources of randomness: the data and the random features used for the sketch
- For the random feature map we have  $<\frac{1}{m}\Phi(x),\frac{1}{m}\Phi(x^{'})>=\frac{1}{m}\sum_{j=1}^{m}e^{-j2\pi w_{j}(x-x^{'})}$
- Averaging over the random features gives the *expected*  $kernel\ k(x,x')=E_w(\exp(-j2\pi < w,x-x'>))$
- We define the *mean kernel* as  $k(p,q) = E_{x \sim p, x' \sim q} k(x,x')$
- A quantity of interest is the maximum mean discrepancy  $MMD(p,q) = \sqrt{k(p,p) 2k(p,q) + k(q,q)}$
- It can be shown using concentration of measure that  $\frac{1}{\sqrt{2}}\textit{MMD}(p_{\theta},p_{\theta'}) \leq \frac{1}{\sqrt{m}}||\mathcal{A}(p_{\theta})-\mathcal{A}(p_{\theta'})||$  when  $\Sigma_{\theta}$  is a finite set
- When  $\Sigma_{\theta}$  is infinite additional assumptions are required to ensure that the LRIP property hodls

# Compressed clustering, fast sketching

- In practice computing the sketch using random Fourier samples might be problematic due to the difficulty in implementing accurately the complex valued function x → exp(-j2πx)
- Using the Fast Walsh-Hadamard transform it is possible to speed up the sketching process
- For blocks of 2<sup>q</sup> by 2<sup>q</sup> matrices, we make layers of alternatively sampling Radamacher diagonal layer matrices followed by Hadamard matrices of dimension 2<sup>q</sup> by 2<sup>q</sup>
- In general, we fit several blocks of this construction vertically and paddle with zeroes until we get proper dimension for W