Goal  To introduce sparsity based regularization with emphasis on the problem of variable selection. To discuss its connection to sparse approximation and describe some of the methods designed to solve such problems.
sparsity based regularization: finite dimensional case
  - introduction
  - algorithms
  - theory
Idea: the target function depends only on few building blocks. Enforcing sparsity of the estimator can be a way to avoid overfitting.

- **Interpretability of the model**: a main goal besides good prediction is detecting the most discriminative information in the data.

- **Compression**: it is often desirable to have parsimonious models, that is models requiring a (possibly very) small number of parameters to be described.

- **Data driven representation**: one can take a large, redundant set of measurements and then use a data driven selection scheme.
A Useful Example

**Biomarker Identification**

**Set up:**
- $n$ patients belong to 2 groups (say two different diseases)
- $p$ measurements for *each* patient quantifying the expression of $p$ genes

**Goal:**
- learn a classification rule to predict occurrence of the disease for future patients
- detect which are the genes responsible for the disease

**High Dimensional learning**

$p \gg n$: typically $n$ is in the order of tens and $p$ of thousands....
Some Notation

Measurement matrix

Let \( X \) be the \( n \times p \) measurements matrix.

\[
X = \begin{pmatrix}
x_1^1 & \ldots & \ldots & \ldots & x_1^p \\
\vdots & \vdots & \vdots & \vdots & \vdots \\
x_n^1 & \ldots & \ldots & \ldots & x_n^p \\
\end{pmatrix}
\]

- \( n \) is the number of examples
- \( p \) is the number of variables
- we denote with \( X^j, j = 1, \ldots, p \) the columns of \( X \)

For each patient we have a response (output) \( y \in \mathbb{R} \) or \( y = \pm 1 \).

In particular we are given the labels for the training set

\[
Y = (y_1, y_2, \ldots, y_n)
\]
So far we still have to define what are "relevant" variables. Different approaches are based on different way to specify what is relevant.

- Filters methods.
- Wrappers.
- Embedded methods.

We will focus on the latter class of methods.

(see "Introduction to variable and features selection" Guyon and Elisseeff ’03)
The selection procedure is **embedded** in the training phase.

### An intuition

what happens to the generalization properties of empirical risk minimization as we discard variables?

- if we keep all the variables we probably overfit,
- if we take just a few variables we are likely to oversmooth (in the limit we have a single variable classifier).

We are going to discuss this class of methods in detail.
Suppose the output is a linear combination of the variables

\[ f(x) = \sum_{i=1}^{p} \beta_i x_i = \langle \beta, x \rangle \]

each coefficient \( \beta_i \) can be seen as a weight on the \( i \)-th variable.

**Sparsity**

We say that a function is *sparse* if most coefficients in the above expansion are zero.
In vector notation we can write the problem as a linear system of equation

\[ Y = X\beta. \]

The problem is *ill-posed.*
Define the $\ell_0$-norm (not a real norm) as

$$\|\beta\|_0 = \#\{i = 1, \ldots, p \mid \beta^i \neq 0\}$$

It is a measure of how "complex" is $f$ and of how many variables are important.
If we assume that a few variables are meaningful we can look for

$$\min_{\beta \in \mathbb{R}^p} \left\{ \frac{1}{n} \sum_{j=1}^{n} V(y_j, \langle \beta, x_j \rangle) + \lambda \| \beta \|_0 \right\}$$

this corresponds to the problem of best subset selection is hard.

This problem is not computationally feasible

⇒ It is as difficult as trying all possible subsets of variables.

Can we find meaningful approximations?
Approximate solutions

Two main approaches

Approximations exist for various loss functions usually based on:

2. Greedy schemes.

We mostly discuss the first class of methods (and consider the square loss).
A natural approximation to $\ell_0$ regularization is given by:

$$
\frac{1}{n} \sum_{j=1}^{n} V(y_j, \langle \beta, x_j \rangle) + \lambda \| \beta \|_1
$$

where $\| \beta \|_1 = \sum_{i=1}^{p} |\beta^i|$. 

If we choose the square loss

$$
\frac{1}{n} \sum_{j=1}^{n} (y_j - \langle \beta, x_j \rangle)^2 = \| Y - X\beta \|_n^2
$$

such a scheme is called **Basis Pursuit** or **Lasso** algorithms.
What is the difference with Tikhonov regularization?

- We have seen that Tikhonov regularization is a good way to avoid overfitting.
- Lasso provides sparse solution, Tikhonov regularization doesn’t.

Why?
We can go back to:

$$\min_{\beta \in \mathbb{R}^p} \left\{ \frac{1}{n} \sum_{j=1}^{n} V(y_j, \langle \beta, x_j \rangle) + \lambda \sum_{i=1}^{p} |\beta_i|^2 \right\}$$

How about sparsity?

⇒ in general all the $\beta^i$ in the solution will be different from zero.
Consider

\[ \min_{\beta} \left\{ \sum_{i=1}^{p} |\beta_i| \right\} \]

subject to

\[ \| Y - X\beta \|_2^2 \leq R. \]
Geometry of the Problem

\[ \beta_1 \]

\[ \beta_2 \]

\[ \ell_1 \]

\[ \ell_2 \]

\[ \ell_1 + \ell_2 \]

\[ R \]

\[ -R \]

\[ \theta \]
We focus on the square loss so that we now have to solve

$$\min_{\beta \in \mathbb{R}^p} \| Y - \beta X \|^2 + \lambda \| \beta \|_1.$$ 

- Though the problem is no longer hopeless it is nonlinear.
- The functional is convex but not strictly convex, so that the solution is not unique.
- Many possible optimization approaches (interior point methods, homotopy methods, coordinate descent...)
- **Proximal, Forward-Backward Methods**: Using convex analysis tools we can get a simple, yet powerful, iterative algorithm.
An Iterative Thresholding Algorithm

It can be proved that the following iterative algorithm converges to the solution $\beta^\lambda$ of $\ell_1$ regularization as the number of iteration increases.

Set $\beta_0^\lambda = 0$ and let

$$
\beta_t^\lambda = S_\lambda[\beta_{t-1}^\lambda + \tau X^T(Y - X\beta_{t-1}^\lambda)]
$$

where $\tau$ is a normalization constant ensuring that $\tau \|X\| \leq 1$ and the map $S_\lambda$ is defined component-wise as

$$
S_\lambda(\beta^i) = \begin{cases} 
\beta^i + \lambda/2 & \text{if } \beta^i < -\lambda/2 \\
0 & \text{if } |\beta^i| \leq \lambda/2 \\
\beta^i - \lambda/2 & \text{if } \beta^i < \lambda/2 
\end{cases}
$$
Thresholding Function

\[ S_{\lambda}(\beta)_{\gamma} \]

Figure 1. Balancing Principle.

\[ S_{\lambda}(\beta)_{\gamma} = \lambda w_{\gamma}/2 - \lambda w_{\gamma}/2 \]
Set $\beta^\lambda_0 = 0$

for $t=1:t_{\text{max}}$

$$\beta^\lambda_t = S^\lambda[\beta^\lambda_{t-1} + \tau X^T(Y - X \beta^\lambda_{t-1})]$$

- The regularization parameter controls the degree of sparsity of the solution.
- The number of iteration $t$ can be stopped when a certain precision is reached.
- The complexity of the algorithm is $O(tp^2)$ for each value of the regularization parameter.
- The algorithm we just described is very easy to implement but can be quite slow but speed-ups are possible using: 1) adaptive step-size, 2) continuation methods.
What about Theory?

The same kind of approaches are considered in different domains for different (though related) purposes.

- Machine Learning.
- Fixed design regression.
- Compressed sensing.

Similar theoretical questions but different settings (deterministic vs stochastics, random design vs fixed design).
Roughly speaking, the results in compressed sensing show that:
If \( Y = X\beta^* + \xi \), where \( X \) is an \( n \) by \( p \) matrix, \( \xi \sim \mathcal{N}(0, \sigma^2 I) \), \( \beta^* \) has at most \( s \) non zero coefficients and \( s \leq n/2 \) then

\[
\|\beta^\lambda - \beta^*\| \leq C\sigma^2 \log p.
\]

The constant \( C \) depends on the matrix \( X \) and is different from infinity, if the relevant variables are not correlated (RIP, restricted eigenvalue property, mutual coherence...).
Learning algorithms based on sparsity usually suffer from an excessive shrinkage effect of the coefficients. For this reason in practice a two-step procedure is usually used:

- Use Lasso (or Elastic Net) to select the relevant components
- Use ordinary least squares (in fact usually Tikhonov with $\lambda$ small...) on the selected variables.
Some Remarks

- **About Uniqueness**: the solution of $\ell_1$ regularization is not unique. Note that the various solution have the same prediction properties but different selection properties.

- **Correlated Variables**: If we have a group of correlated variables the algorithm is going to select just one of them. This can be bad for interpretability but maybe good for compression.
Consider a more general penalty of the form

$$\|\beta\|_q = \left( \sum_{i=1}^{p} |\beta_i|^q \right)^{1/q}$$

(called bridge regression in statistics).

It can be proved that:

- \( \lim_{q \to 0} \|\beta\|_q \to \|\beta\|_0 \),
- for 0 < q < 1 the norm is not a convex map,
- for q = 1 the norm is a convex map and is strictly convex for q > 1.
One possible way to cope with the previous problems is to consider

$$\min_{\beta \in \mathbb{R}^p} \| Y - \beta X \|^2 + \lambda (\alpha \| \beta \|_1 + (1 - \alpha) \| \beta \|_2^2).$$

- \( \lambda \) is the regularization parameter.
- \( \alpha \) controls the amount of sparsity and smoothness.

(Zhu. Hastie ’05; De Mol, De Vito, Rosasco ’07)
The $\ell_1$ term promotes sparsity and the $\ell_2$ term smoothness.

The functional is strictly convex: the solution is unique.

A whole group of correlated variables is selected rather than just one variable in the group.
Geometry of the Problem

\[ \beta_1 \]

\[ \beta_2 \]

\[ \ell_1 \]

\[ \ell_2 \]

\[ \ell_1 + \ell_2 \]

\[ R - R \ell_1 + \ell_2 \]

\[ \theta \]

L. Rosasco

Sparsity Based Regularization
Geometry of the Problem
Sparsity based regularization gives a way to deal with high dimensional problems...
while performing principled variable selection.
Very active field, connections with signal processing–compressed sensing, statistics and approximation theory.
Lasso is sparse but not stable (suffer where variables are correlated), elastic net is a way to solve this problem.

Current work: low rank matrix estimation, non linear variable selection.