# Optimization in higher dimensions 

- Quasi-Newton Methods
- Conjugate Gradient Method


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## A bit of history

[Nocedal, Wright, Numerical Optimization 06], Chapters 6-7
$\star$ in the 50s W.C. Davidon used "coodrdinate descent" method (GD on coordinates)

* the computer would always crash before the simulation was finished
$\star$ Davidon decided to find a way of accelerating the optimization process: he found one of the most creative ideas in nonlinear optimization
* Fletcher and Powell demonstrated that this algorithm was faster and more reliable than existing methods at the time
* paradoxically, Davidon's paper was not accepted for publication. It remained a technical report for more than thirty years until it appeared in SIAM Journal on Optimization in 1991!


## Motivation

Recall the Variable Metric Method and replace $A_{i}^{-1}$ by $S_{i}$ :

## Algorithm 1 (Generic Variable Metric method)

Choose the starting point $x_{0}$
Iteration $i$ :

- compute $f\left(x_{i}\right), \nabla f\left(x_{i}\right)$ and eventually $D^{2} f\left(x_{i}\right)$
- choose a symmetric positive-definite matrix $S_{i}$ : compute the new direction

$$
d_{i}=-S_{i} \nabla f\left(x_{i}\right)
$$

- perform a line-search from $x_{i}$ in the direction $d_{i}$ giving a new iterate

$$
x_{i+1}=x_{i}+t_{i} d_{i}=x_{i}-t_{i} S_{i} \nabla f\left(x_{i}\right)
$$

$\star$ in the modified Newton method $S_{i}$ is computed as follows: find the Hessian $D^{2} f\left(x_{i}\right)$, modify it to make it "well positive definite", then invert it or solve $S_{i} d_{i}=\nabla f\left(x_{i}\right)$

* in quasi-Newton method we try to skip all of this and compute $S_{i}$ recursively with one objective: $S_{i}-\left(D^{2} f\left(x_{i}\right)\right)^{-1} \rightarrow 0$
$\star$ minimize $f(x)=\frac{1}{2} x^{T} A x-b^{T} x$ with Steepest Descent line-search $\star$ denote $E\left(x_{i}\right)=f\left(x_{i}\right)-\min f$ : error in terms of objective function $\star x_{i+1}=x_{i}-t_{\text {opt }} S_{i} \nabla f\left(x_{i}\right)$ is equivalent to a change of coordinates $\xi=S_{i}^{1 / 2} x$ $\star$ the step $i$ in the VM method is just a Steepest-Descent step for the matrix $S_{i}^{1 / 2} A S_{i}^{1 / 2}$. Therefore we have the estimate

$$
E\left(x_{i+1}\right) \leq\left(\frac{Q-1}{Q+1}\right)^{2} E\left(x_{i}\right)
$$

where $Q$ is the condition number of $S_{i}^{1 / 2} A S_{i}^{1 / 2}$
$\star$ if $S_{i}$ is close to $D^{2} f\left(x_{i}\right)^{-1}=A^{-1}$ then $S_{i}^{1 / 2} A S_{i}^{1 / 2}$ is close to the identity matrix so $Q$ is close to 1 .
$\star$ Finally, if $Q$ converges to 1 , we eventually get that $E\left(x_{i+1}\right) / E\left(x_{i}\right) \rightarrow 0$, i.e. super-linear convergence

## Basic rules for updating $S_{i}$

$\star$ Taylor expansion formula tells us that

$$
\nabla f\left(x_{i+1}\right)-\nabla f\left(x_{i}\right) \approx D^{2} f\left(x_{i}\right)\left(x_{i+1}-x_{i}\right)
$$

$\star$ Therefore, it is reasonable to request that

$$
S_{i+1}\left(\nabla f\left(x_{i+1}\right)-\nabla f\left(x_{i}\right)\right)=x_{i+1}-x_{i}
$$

called the secant relation (make parallel with the 1D case)
$\star$ With the notations $g_{i}=\nabla f\left(x_{i}\right), p_{i}=x_{i+1}-x_{i}, q_{i}=g_{i+1}-g_{i}$ we have

$$
S_{i+1} q_{i}=p_{i},
$$

called the quasi-Newton equation

* this leaves us with infinitely many possibilities... another goal is that

$$
S_{i+1}-S_{i} \text { is as simple as possible! }
$$

* initialization? one may simply choose $S_{0}=\mathrm{Id}$
* idea: find $S_{i+1}=S_{i}+B_{i}$ where $B_{i}$ has low rank
$\star$ Rank 1 updates: $B_{i}=\alpha_{i} v_{i} v_{i}^{T}$ - one may find $B_{i}$ such that the quasi-Newton relation holds

$$
S_{i+1}=S_{i}+\alpha_{i} z_{i} z_{i}^{T}
$$

$\star$ the quasi-Newton relation $p_{i}=S_{i+1} q_{i}$ implies

$$
z_{i}=\omega_{i}\left(p_{i}-S q_{i}\right)
$$

$\star$ in the end we get

$$
S_{i+1}=S_{i}+\frac{1}{\left(p_{i}-S_{i} q_{i}\right)^{T} q_{i}}\left[p_{i}-S_{i} q_{i}\right]\left[p_{i}-S_{i} q_{i}\right]^{T}
$$

$\star$ not possible to guarantee that $S_{i+1}$ is positive definite if $S_{i}$ is

## Rank 2 updates: DFP

夫 Davidon-Fletcher-Powell: historically, the first "good" quasi-Newton method $\star$ use rank 2 updates: guarantee the positive-definiteness of $S_{i+1}$ under reasonable hypotheses

## Proposition 1

Let $S$ be a positive definite symmetric matrix and $p$ and $q$ be two vectors such that $p^{T} q>0$. Then the matrix

$$
S^{\prime}=S+\frac{1}{p^{T} q} p p^{T}-\frac{1}{q^{T} S q} S q q^{T} S
$$

is symmetric positive definite and satisfies $S^{\prime} q=p$.

* Proof: just compute $S^{\prime} q$ and $x S^{\prime} x$ and do a bit of linear algebra.
$\star$ How to get this idea? Just choose $S_{i+1}=S_{i}+\alpha u u^{T}+\beta v v^{\top}$ (rank 2 update)
$\star$ then choose $u=p_{i}$ and $v=S_{i} q_{i}$
* DFP update:

$$
S_{i+1}=S_{i}+\frac{1}{p_{i}^{T} q_{i}} p_{i} p_{i}^{T}-\frac{1}{q_{i}^{T} S_{i} q_{i}} S_{i} q_{i} q_{i}^{T} S_{i}
$$

$\star$ the condition $q_{i}^{T} p_{i}>0$ is equivalent to

$$
\left(\nabla f\left(x_{i+1}\right)-\nabla f\left(x_{i}\right)\right) \cdot\left(x_{i+1}-x_{i}\right)>0,
$$

which is true if $f$ is strictly convex: reasonable assumption near a minimum... $\star$ when using Wolfe line-search we can guarantee that $q_{i}^{\top} p_{i}>0$.
$\star$ for the quadratic case DFP becomes the conjugate gradient method $\star$ it turns out DFP is not the best method out there...

- it does not "self-correct" when $S_{i}$ gets far from the inverse Hessian


## Duality: quasi-Newton relation

$\star$ any quasi-Newton update can generate another one:

- $S_{i+1}=S_{i}+B_{i}\left(S_{i}, p_{i}, q_{i}\right)$ such that $S_{i+1} q_{i}=p_{i}$
- then $q_{i}=S_{i+1}^{-1} p_{i}$ where $S_{i+1}^{-1}=\left(S_{i}+B\left(S_{i}, p_{i}, q_{i}\right)\right)^{-1}$
- switching the roles of $p_{i}$ and $q_{i}$ we get a different update, called the dual update
$\star$ how to get the dual of DFP: replace $S_{i}$ with $S_{i}^{-1}$ and interchange $p_{i}$ and $q_{i}$

$$
S_{i+1}^{-1}=S_{i}^{-1}+\frac{1}{q_{i}^{T} p_{i}} q_{i} q_{i}^{T}-\frac{1}{p_{i}^{T} S_{i}^{-1} p_{i}} S_{i}^{-1} p_{i} p_{i}^{T} S_{i}^{-1}
$$

* a direct computation or Sherman-Morrison's formula gives:

$$
S_{i+1}=S_{i}-\frac{p_{i} q_{i}^{T} S_{i}+S_{i} q_{i} p_{i}^{T}}{p_{i}^{T} q_{i}}+\left(1+\frac{q_{i}^{T} S_{i} q_{i}}{p_{i}^{T} q_{i}}\right) \frac{p_{i} p_{i}^{T}}{p_{i}^{T} q_{i}}
$$

## The BFGS update

^ BFGS: Broyden, Fletcher, Goldfarb, Shanno

$$
S_{i+1}=S_{i}-\frac{p_{i} q_{i}^{T} S_{i}+S_{i} q_{i} p_{i}^{T}}{p_{i}^{T} q_{i}}+\left(1+\frac{q_{i}^{T} S_{i} q_{i}}{p_{i}^{T} q_{i}}\right) \frac{p_{i} p_{i}^{T}}{p_{i}^{T} q_{i}}
$$

* widely used in most of the codes implemented today
* since BFGS is the dual of DFP, and a matrix is positive-definite if and only if its inverse is positive-definite, the BFGS update maintains positive-definiteness if $p_{i}^{T} q_{i}>0$ (same hypothesis as for DFP to work...)
[Nocedal, Wright, Numerical Optimization 06], Chapters 6-7
* Local super-linear convergence: If an algorithm using BFGS with Wolfe's line-search converges to $x^{*}$ where $f$ is strongly convex with Lipschitz Hessian then the convergence rate is super-linear
$\star$ BFGS can also be found by minimizing a certain distance between the inverse Hessian and the rank 2 update $S_{i+1}$ among matrices verifying the secant condition!
* BFGS has effective self-correcting properties


## Extreme cases

## Dimension 1:

* the quasi-Newton relation is just $S_{i+1}=\frac{p_{i}}{q_{i}}$ and we get

$$
x_{i+1}=x_{i}-\frac{x_{i}-x_{i-1}}{f^{\prime}\left(x_{i}\right)-f^{\prime}\left(x_{i-1}\right)} f^{\prime}\left(x_{i}\right)
$$

which is the false position (or secant) method

Large dimension:
$\star$ same disadvantage as Newton methods - a $n \times n$ matrix may be too large to store in memory

* it is possible to store only the update vectors and compute matrix - vector products by doing only scalar - products

$$
\left(u v^{\top}\right) x=u\left(v^{\top} x\right)=\left(v^{\top} x\right) u
$$

$\star$ limited memory-BFGS (LBFGS): use only the last $m$ vectors $p_{i}, q_{i}$ in order to compute $S_{i+1}$ - good behavior in practice despite being an approximation of BFGS

## Computational cost per iteration

* after the function value, gradient and Hessian are computed (this is non-negligible in some applications)
- GD: $O(N)$
- Newton: $O\left(N^{3}\right)$ in worst case (solving a linear system) - it all depends on the structure of the Hessian
- BFGS, DFT: $O\left(N^{2}\right)$ - matrix vector products
- LBFGS: $O(m N)$ where $m$ is the fixed number of gradients to remember


## Practical example: the $N$-dimensional Rosenbrock

$$
f(x)=\sum_{i=1}^{N-1}\left[100\left(x_{i+1}-x_{i}^{2}\right)^{2}+\left(1-x_{i}\right)^{2}\right]
$$

with global minimum at $x^{*}=(1,1, \ldots, 1)$.
$\star$ ill conditioning: the optimization process wants to achieve $x_{i+1} \approx x_{i}^{2}$ rather than minimizing $\left(x_{i}-1\right)^{2}$ and go towards the global minimum!


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## Conclusion: quasi-Newton methods

- equivalent of the Secant method in higher dimensions
- achieve super-linear convergence without using the Hessian
- for extremely large $n$ BFGS may be costly from a memory point of view: if possible use L-BFGS instead
- BFGS and LBFGS are often available in standard optimization libraries: Example scipy.optimize.minimize


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## Motivation

$\star$ if $A$ is symmetric, positive-definite then solving the system $A x=b$ is equivalent to minimizing the quadratic function

$$
f: x \mapsto \frac{1}{2} x^{\top} A x-b \cdot x
$$

$\star$ the gradient of this quadratic function is $\nabla f(x)=A x-b$
$\star$ direct method: process details about the matrix $A$ (factorization) and then solve the system: complexity is between $O\left(n^{2}\right)$ and $O\left(n^{3}\right)$.
$\star$ in contrast to this, iterative algorithms produce an approximation of the solution, which might be good enough for very large $n$

* for example: the gradient algorithm with Steepest-Descent will quickly converge to the optimum, but we can do better



## Conjugate directions

* A given symmetric positive-definite matrix $A$ defines a scalar product

$$
\langle x, y\rangle=x^{\top} A y
$$

$\star$ Two (non-zero) directions $d_{1}$ and $d_{2}$ are called conjugate with respect to $A$ if they are orthogonal w.r.t. the above scalar product:

$$
d_{1} \text { and } d_{2} \text { are conjugate } \Longleftrightarrow d_{1} A d_{2}=0
$$

$\star$ we may also call two directions which are conjugate w.r.t. $A$ as being $A$-orthogonal
$\star$ why is this useful? suppose $d_{1}, \ldots, d_{k}$ are mutually $A$-orthogonal and we have the decomposition

$$
d=\sum_{j=1}^{k} \alpha_{j} d_{j}
$$

Then, using the orthogonality property, we can find the coefficients $\alpha_{i}$ explicitly:

$$
d_{i}^{T} A d=\alpha_{i} d_{i}^{T} A d_{i} \Rightarrow \alpha_{i}=\frac{d_{i}^{T} A d}{d_{i}^{T} A d_{i}}=\frac{\left\langle d, d_{i}\right\rangle}{\left\langle d_{i}, d_{i}\right\rangle}
$$

$\star$ Consequence: If $d_{1}, \ldots, d_{k}$ are mutually orthogonal then they are linearly independent! (for a proof, use the above formula to see that $d=0 \Rightarrow \alpha_{i}=0$ )

## Why is this concept useful?

## Proposition 2 (Solve a system using Conjugate Directions)

Let $A$ be a symmetric positive-definite matrix and $d_{1}, \ldots, d_{n}$ a (complete) system of $n$ non-zero $A$-orthogonal vectors. Then the solution $x^{*}$ to the system $A x=b$ is given by the formula

$$
x^{*}=\sum_{j=1}^{n} \frac{b^{T} d_{j}}{d_{j}^{T} A d_{j}} d_{j}
$$

* An equivalent formulation:

$$
x^{*}=A^{-1} b=\sum_{j=1}^{n} \frac{b^{T} d_{j}}{d_{j}^{T} A d_{j}} d_{j}=\left(\sum_{j=1}^{n} \frac{1}{d_{j}^{T} A d_{j}} d_{j} d_{j}^{T}\right) b
$$

which gives us the explicit inverse of $A$

$$
A^{-1}=\sum_{j=1}^{n} \frac{1}{d_{j}^{T} A d_{j}} d_{j} d_{j}^{T}
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$\star$ All this is good when we know a complete family of $A$-orthogonal directions!

## Conjugate Directions: quadratic case

## Algorithm 2 (Conjugate Directions method)

Let $A$ be a $n \times n$ symmetric positive-definite matrix, $b$ a vector and $f(x)=\frac{1}{2} x^{\top} A x-b^{T} x$ the quad. form associated to $A$ and $b$.

Let $d_{0}, . ., d_{n-1}$ be a system of $A$-orthogonal vectors and $x_{0}$ a starting point.
Then, with the notation $g_{i}=\nabla f\left(x_{i}\right)=A x_{i}-b$, the iterative process

$$
x_{i+1}=x_{i}+\gamma_{i} d_{i}, \gamma_{i}=-\frac{d_{i}^{T} g_{i}}{d_{i}^{\top} A d_{i}}, i=1, \ldots, n
$$

converges to the unique minimizer $x^{*}$ of $f$ in $n$ steps.

* The step $\gamma_{i}$ is optimal in the direction $d_{i}$ : define $q(t)=f(x+t d)$ then

$$
q^{\prime}(t)=\nabla f(x+t d) \cdot d=d \cdot \nabla f(x)+t d^{\top} A d
$$

$\star$ Proof: just look at $x_{n}$ and see that it gives exactly the formula for $x^{*}$.
$\star$ Important idea: $d_{k} A\left(x_{k}-x_{0}\right)=0$ for any $k \geq 0$

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$\star$ Important idea: $d_{k} A\left(x_{k}-x_{0}\right)=0$ for any $k \geq 0$
$\star$ Again: all this is good when we know a complete family of $A$-orthogonal directions!

## Properties of the Conjugate Directions Method

$\star$ define for each $i \geq 1$ the linear space $\mathcal{B}_{i-1}=\operatorname{Span}\left\{d_{0}, \ldots, d_{i-1}\right\}$
$\star$ if we define the affine subspaces $M_{i}=x_{0}+\mathcal{B}_{i-1}$ then

$$
\left\{x_{0}\right\}=M_{0} \subset M_{1} \subset \ldots \subset M_{n}=\mathbb{R}^{n}
$$

* the Conjugate Directions method generate the minimizers of $f$ in each of the affine spaces $M_{i}$


## Proposition 3

For every $1 \leq i \leq n$ the vector $x_{i}$ is the minimizer of $f$ on the affine subspace $M_{i}=x_{0}+\mathcal{B}_{i-1}$. In particular, as shown previously, $x_{i}$ minimizes $f$ on the line $\left\{x_{i-1}+t d_{i-1}: t \in \mathbb{R}\right\}$.

Proof: $\star$ Compute the gradient $g_{i}=\nabla f\left(x_{i}\right)=A x_{i}-b$ and note that $g_{i}$ is orthogonal to $d_{0}, \ldots, d_{i-1}$.
$\star$ Then obtain that $\left\langle\nabla f\left(x_{i}\right), x-x_{i}\right\rangle=0$ for $x \in x_{0}+\mathcal{B}_{i-1}$.
$\star f$ is strictly convex so Euler's inequality tells us that $x_{i}$ is indeed the minimizer of $f$ in $x_{0}+\mathcal{B}_{i-1}$.

## Build a basis of conjugated directions

* recall the Gram-Schmidt procedure
$\star$ define the $A$-projection of $v$ on $u$ :

$$
\operatorname{proj}_{u}(v)=\frac{\langle u, v\rangle}{\langle u, u\rangle} u=\frac{u^{T} A v}{u^{T} A u} u
$$

## Algorithm 3 (Gram-Schmidt)

0. Take a basis $\left(v_{i}\right)$ of $\mathbb{R}^{n}$ : e.g. the canonical basis.
1. $u_{1}=v_{1}$
2. $u_{2}=v_{2}-\operatorname{proj}_{u_{1}}\left(v_{2}\right)$
3. $u_{3}=v_{3}-\operatorname{proj}_{u_{1}}\left(v_{3}\right)-\operatorname{proj}_{u_{2}}\left(v_{3}\right)$
n. $u_{n}=v_{n}-\operatorname{proj}_{u_{1}}\left(v_{n}\right)-\ldots-\operatorname{proj}_{u_{n-1}}\left(v_{n}\right)$

In the end normalize the vectors: $d_{i}=\frac{1}{\sqrt{u_{i}^{T} A u_{i}}} u_{i}$
$\star$ in this form the process is not numerically stable: due to rounding errors the vectors $u_{k}$ may not be exactly orthogonal...

## Conjugate Gradient Method

$\star$ we can compute the family of $A$-orthogonal directions during the optimization algorithm

## Algorithm 4 (Conjugate Gradient)

Choose arbitrary initialization point $x_{0}$ and set $d_{0}=-g_{0}=-\nabla f\left(x_{0}\right)=b-A x_{0}$
Loop on: $i=0, \ldots, n-1$

- if $\nabla f\left(x_{i}\right)=0$ then stop.
- $x_{i+1}=x_{i}+\gamma_{i} d_{i}$ with $\gamma_{i}=-\frac{d_{i}^{T} g_{i}}{d_{i}^{\top} A d_{i}}$
- Compute new gradient $g_{i+1}=\nabla f\left(x_{i+1}\right)=A x_{i+1}-b$
- Compute new direction $d_{i+1}=-g_{i+1}+\beta_{i} d_{i}$ with $\beta_{i}=\frac{g_{i+1}^{T} A d_{i}}{d_{i}^{T} A d_{i}}$
$\star$ as before $\gamma_{i}$ is the optimal step in the direction $d_{i}$
$\star$ the parameter $\beta_{i}$ is chosen such that $d_{i+1}^{T} A d_{i}=0$
$\star$ the new direction $d_{i+1}$ is given by the projection of the anti-gradient direction
$-g_{i+1}$ on the previous direction


## Proposition 4 (CG is a Conjugate Direction method)

If the algorithm does not terminate at step $i$ then:

- the gradients $g_{0}, \ldots, g_{i-1}$ at $x_{0}, \ldots, x_{i-1}$ are non-zero and $\operatorname{Span}\left\{g_{0}, g_{1}, \ldots, g_{i-1}\right\}=\operatorname{Span}\left\{g_{0}, A g_{0}, \ldots, A^{i-1} g_{0}\right\}$
- The directions $d_{0}, \ldots, d_{i-1}$ are non-zero and $\operatorname{Span}\left\{d_{0}, d_{1}, \ldots, d_{i-1}\right\}=\operatorname{Span}\left\{g_{0}, A g_{0}, \ldots, A^{i-1} g_{0}\right\}$
- The directions $d_{0}, \ldots, d_{i-1}$ are $A$ orthogonal
- Alternative formulas for $\gamma_{i}$ and $\beta_{i}$ :

$$
\gamma_{i}=\frac{g_{i}^{T} g_{i}}{d_{i}^{T} A d_{i}} \text { and } \beta_{i}=\frac{g_{i+1}^{T} g_{i+1}}{g_{i}^{T} g_{i}} .
$$

$\star$ A sequence of the type $g_{0}, A g_{0}, A^{2} g_{0}, \ldots$ is called a Krylov sequence

## Consequences and convergence

$\star x_{i}$ is the minimizer of $f$ in the affine subspace

$$
x_{0}+\operatorname{Span}\left\{d_{0}, \ldots, d_{i-1}\right\}=x_{0}+\operatorname{Span}\left\{g_{0}, A g_{0}, \ldots, A^{i-1} g_{0}\right\}
$$

$\star x_{i}$ is the minimizer of $f$ in the affine subspace generated by $x_{0}$ and polynomials of $A$ of degree at most $i-1$ times $g_{0}$ (denote this polynomial space by $\mathcal{P}_{i-1}$ )

$$
x_{0}+\left\{p(A) g_{0}: p(z)=\sum_{i=0}^{i-1} p_{i} z^{i}\right\}
$$

$\star$ error in terms of the objective function: $E(x)=f(x)-\min f=\frac{1}{2}\left(x-x^{*}\right)^{\top} A\left(x-x^{*}\right)$

## Proposition 5 (Error for CG)

$$
E\left(x_{i}\right)=\min _{p \in \mathcal{P}_{i-1}} \frac{1}{2}\left(x_{0}-x^{*}\right) A(\operatorname{ld}-A p(A))^{2}\left(x_{0}-x^{*}\right)
$$

$\star$ Proof: write $x_{i}=x_{0}+p(A) g_{0}$ and recall that $\nabla f\left(x_{i}\right)=A\left(x_{i}-x^{*}\right)$

## Corollary

Let $\Sigma$ be the spectrum of $A$. Then

$$
E\left(x_{i}\right) \leq E\left(x_{0}\right) \min _{p \in \mathcal{P}_{i}^{*}} \max _{\lambda \in \Sigma} p^{2}(\lambda),
$$

where $\mathcal{P}_{i}^{*}$ is the set of polynomials $p$ of degree at most $i$ such that $p(0)=1$.
Another estimate is

$$
E\left(x_{i}\right) \leq \frac{1}{2}\left|x^{*}-x_{0}\right|^{2} \min _{p \in \mathcal{P}_{i}^{*}} \max _{\lambda \in \Sigma} \lambda p^{2}(\lambda),
$$

* Proof: use an orthonormal basis made of eigenvectors of $A$
$\star$ denote by $Q$ the condition number of $A$. Then there exists a polynomial $q \in \mathcal{P}_{s}^{*}$ such that

$$
\max _{\lambda \in \Sigma} q_{s}(\lambda)^{2} \leq 4\left(\frac{\sqrt{Q}-1}{\sqrt{Q}+1}\right)^{2 s}
$$

## Error estimate in terms of the condition number

* for the Conjugate Gradient algorithm we have

$$
E\left(x_{N}\right) \leq 4\left(\frac{\sqrt{Q}-1}{\sqrt{Q}+1}\right)^{2 N} E\left(x_{0}\right)
$$

where $Q$ is the condition number of $A$.
$\star$ compare this with the error estimate for the Steepest-Descent

$$
E\left(x_{N}\right) \leq\left(\frac{Q-1}{Q+1}\right)^{2 N} E\left(x_{0}\right)
$$

* in order to reduce the initial error by a factor of $\varepsilon$ one needs to do $O(Q)$ steps with Steepest Descent compared to $O(\sqrt{Q})$ steps with CG. This is a big difference!
$\star$ CG is supposed to converge in $n$ iterations, however rounding errors may prevent the convergence!
$\star$ moreover, if $A$ has $k \leq n$ distinct eigenvalues then CG converges in $k$ iterations!
* Often, for $n$ large, the process is stopped before reaching $n$ iterations, when the error estimate is small enough


## Example: Hilbert matrices

$A=(1 /(i+j-1))_{1 \leq i, j \leq n}$, ill conditioned

* below you can see a comparison between GD with optimal step and CG. The residual $|A x-b|$ is plotted at every iteration
$\star$ the residual decreases slowly for GD: the algorithm tends to go multiple times in the same direction! CG optimizes once and for all in the current direction.
$\star$ small residual does not mean that $x$ is close to $x^{*}: A x-b=A\left(x-x^{*}\right)$ !

GD vs CG: Hilbert matrix $N=4 Q=1.6 e+04$


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- Consider Laplace's equation

$$
\text { Find } u \in H_{0}^{1}(D) \text { such that }\left\{\begin{array}{rll}
-\Delta u & =f & \text { in } D \\
u & =0 & \text { on } \partial D
\end{array}\right.
$$

where $f \in L^{2}(D)$ is a given source.

- It is possible to associate to this a variational formulation:

Find $u \in V$ such that $\forall v \in V$ we have $a(u, v)=\ell(v)$
where

- The Hilbert space $V$ is a Sobolev space $H_{0}^{1}(D)$
- $a(\cdot, \cdot)$ is a bilinear form on $V$ given by $a(u, v)=\int_{D} \nabla u \cdot \nabla v d x$
- $\ell(\cdot)$ is a linear form on $V$ given by $\ell(v)=\int_{D} f v d x$
- Lax-Milgram's theorem assures us that such a problem has a solution on $V$.
- The finite element method proposes to search for an approximation $u_{h}$ in a finite dimension subspace $V_{h} \subset V$.
- the variational formulation is replaced by:

Find $u_{h} \in V_{h}$ such that $\forall v_{h} \in V_{h}$ we have $a\left(u_{h}, v_{h}\right)=\ell\left(v_{h}\right)$

- Advantage : $V_{h}$ being of finite dimension, we can choose a basis $\mathcal{B}=\left\{\varphi_{i}\right\}_{i=1}^{N}$ and the variational formulation becomes a linear system $A \bar{u}=b$ with

$$
A=\left(a\left(\varphi_{i}, \varphi_{j}\right)\right), b=\left(\ell\left(\varphi_{i}\right)\right)
$$

where $\bar{u}$ are the coordinates of $u_{h}$ in the basis $\mathcal{B}$.

- The choice of the basis is important: one objective is to have a system given by a sparse matrix


## Construct a finite element space

- The domain $D$ is discretized using a mesh $\mathcal{T}_{h}$ which consists of a partitions in triangles in 2D or tetrahedra in 3D.
- The parameter $h$ which indicates the convergence of the method is typically related to the size of the mesh elements.



## Construct a finite element space (2)

A basis $\left\{\varphi_{1}, \ldots, \varphi_{N_{h}}\right\}$ of finite element functions is introduced on the mesh $\mathcal{T}_{h}$ Example

- $N_{h}$ is the number of vertices $a_{1}, \ldots, a_{N_{h}}$ of the mesh
- For each $i=1, \ldots, N_{h}, \varphi_{i}$ is affine on each triangle $T \in \mathcal{T}_{h}$ and

$$
\varphi_{i}\left(a_{j}\right)=1 \text { et } \varphi_{i}\left(a_{j}\right)=0 \text { pour } i \neq j
$$



## Formulation of a matrix system

Decompose the solution $u_{h}$ in the basis of finite elements

$$
u_{h}=\sum_{i=1}^{N_{h}} u_{j} \varphi_{i}
$$

and the variational problem becomes a linear system of size $N_{h} \times N_{h}$

$$
K U=f
$$

where

- $U=\left(\begin{array}{c}u_{1} \\ \vdots \\ u_{N_{h}}\end{array}\right)$ is the vector of coefficients
- $K$ is the rigidity matrix given by $K_{i j}=a\left(\varphi_{i}, \varphi_{j}\right)$
- $F$ is the vector $F=\left(\ell\left(\varphi_{i}\right)\right)_{i=1, \ldots, N_{h}}$.
$\star$ The matrix $K$ will be symmetric and positive-definite so we are in the good framework where CG works!
* when $N_{h}$ is large (a few tens of thousands of elements) direct methods will fail to work (computation time, memory limitations)
$\star$ CG will work well even for $N_{h}>10^{5}$


## Some results



## CG for general functions

## Algorithm 5 (Fletcher-Reeves CG on $\mathbb{R}^{n}$ )

Choose a starting point $x_{0}$. Set cycle counter $k=1$.
Cycle $k$ : Initialization of the cycle: Given $x_{0}$ compute $g_{0}=\nabla f\left(x_{0}\right), d_{0}=-g_{0}$ Inner Loop: for $i=0, \ldots, n-1$

- if $g_{i}=0$ terminate, otherwise set $x_{i+1}$ as the minimizer of $f\left(x_{i}+t d_{i}\right)$
- compute $g_{i+1}=\nabla f\left(x_{i+1}\right)$
- set $d_{i+1}=-g_{i+1}+\beta_{i} d_{i}$ with $\beta_{i}=\frac{g_{i+1}^{T} g_{i+1}}{g_{i}^{T} g_{i}}$

When the loop is finished replace $x_{0}$ with $x_{n}$ and restart.

* note that in the inner loop we have a Steepest Descent line-search: this is not applicable in general. A line-search procedure should be used instead! $\star$ It can be proved that in the non-degenerate case the convergence is quadratic in the number of cycles i.e.

$$
\left|x^{k+1}-x^{*}\right| \leq C\left|x^{k}-x^{*}\right|^{2}
$$

where $x^{k}$ is the sequence of starting points for cycles

## Comparison with previous methods

$\star$ again on the Rosenbrock function for $N=100$

* in general nonlinear-CG converges faster than GD but not necessarily faster than quasi-Newton methods



## Conclusion on Conjugate Gradient method

- when a complete system of $A$-orthogonal directions is known everything is explicit
- it can be made into an iterative algorithm with a convergence ratio way better than Steepest Descent
- it converges in $n$ iterations (theoretically). In practice, for large $n$, we usually stop the process once the error estimate

$$
E\left(x_{N}\right) \leq 4\left(\frac{\sqrt{Q}-1}{\sqrt{Q}+1}\right)^{2 N} E\left(x_{0}\right)
$$

is satisfying.

- cost of a step in CG:

$$
O(n)+\text { cost of a matrix-vector multiplication } d \rightarrow A d
$$

This is particularly efficient when $A$ is sparse (has few non-zero elements)

- Disadvantage: sensitivity to the condition number!


## Conclusions: unconstrained optimization in ND

- Gradient Descent algorithms: sensitive to conditioning!
- Newton methods: fast convergence under right hypotheses. Major practical inconveniences:
- compute Hessian matrix and (possibly) store it
- doesn't necessarily decrease the function value
- solve a linear system at every iteration
- variable metric methods: compute an approximation of the inverse Hessian
- BFGS: rank 2 updates, standard in available implementations
- even better for large $n$ : L-BFGS - limit memory by using only information from the previous $m$ iterations
- Conjugate Gradient methods: less sensitive to conditioning than Steepest Descent
- Newton-Gauss: non-linear least squares
- Nedler-Mead: gradient free method
* get used to the structure of algorithms which are already implemented: in the practical session you will play with tools from scipy.optimize
* keep in mind to minimize the number of function evaluations in your codes: not all functions to be optimized are computed in a cheap way
- when the value of a function or its gradient are used multiple times store them in some variables
- in some computations involving physical simulations the gradient can often be computed using existing information from the solution given by the model: there is no point computing it multiple times

