Image Processing and Computer Graphics

Image Processing

Class 3
Energy Minimization

Formalize your model assumptions and cast the task as an optimization problem

$$E(x) = A_1(x) + \dots + A_n(x)$$

2. Solve the optimization problem

$$x^* = \operatorname{argmin}_x E(x)$$

- Objective function E(x) often called **energy** (motivated from physics)
- In machine learning it is often called loss function

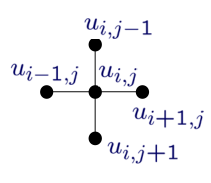
Example: image denoising

- First step: formulate the model assumptions
 - The outcome should be similar to the input image
 - The result should be smooth
- Second step: formalize these assumptions
 - Similarity to the input data (data term):

$$E_D(u_{i,j}) := \sum_{i,j} (u_{i,j} - I_{i,j})^2 \to \min$$

Similarity to neighboring values (smoothness term):

$$E_S(u_{i,j}) := \sum_{i,j} (u_{i+1,j} - u_{i,j})^2 + (u_{i,j+1} - u_{i,j})^2 \to \min$$



- Yields an energy minimization problem including a weighting parameter lpha

$$u_{i,j}^* = \operatorname{argmin}_{u_{i,j}} \left(E_D(u_{i,j}) + \alpha E_S(u_{i,j}) \right)$$

Third step: solve this optimization problem

Advantages

- All model assumptions are clearly stated
 → transparency
- Global approach: all variables are optimized in a joint manner; interdependencies are not lost by intermediate decisions → optimality
- Theoretical aspects can be analyzed:
 - Existence and uniqueness of solutions
 - Stability of solutions with respect of the input data
 - Difficulty of the problem class
- Usually fewer parameters than in heuristic multi-step methods
- Compatibility: approaches can be combined more easily

 Formalizing the assumptions appears rather ad-hoc: Why minimizing the sum of squared differences and not some other distance?

$$E_D(u_{i,j}) := \sum_{i,j} (u_{i,j} - I_{i,j})^2 \to \min$$

- → Probabilistic interpretations can usually answer this question.
- Choosing the weight parameter(s) is not easy and often depends on the data.
 - → Parameters can be optimized via a validation dataset.
- Global optimization is often hard.
 Heuristics can obliterate the initial transparency of the model.

Energy minimization in our denoising example

Here is our energy from the denoising example

$$E(u_{i,j}) := \sum_{i,j} \left(\underbrace{(u_{i,j} + I_{i,j})^2 + \alpha \left(\underbrace{(u_{i+1,j} + u_{i,j})^2 + \underbrace{(u_{i,j+1} + u_{i,j})^2}}_{} \right)}^2 + \underbrace{(u_{i,j} + I_{i,j})^2}_{} \right)$$

- How do we find the minimum?
- Necessary condition for a minimum: the first derivatives must be zero

$$\frac{dE}{du} = 0 \quad \Leftrightarrow \quad \frac{\partial E}{\partial u_{i,j}} = 0 \quad \forall i, j$$

Here we go:

$$\frac{\partial E}{\partial u_{i,j}} = 2(u_{i,j} - I_{i,j}) / +2\alpha(u_{i,j} - u_{i,j-1}) - 2\alpha(u_{i+1,j} - u_{i,j}) +2\alpha(u_{i,j} - u_{i,j-1}) - 2\alpha(u_{i,j+1} - u_{i,j}) = 0$$

At boundary pixels some terms are missing due to missing neighbors

Linear system of equations

Necessary conditions...

$$\frac{\partial E}{\partial u_{i,j}} = (u_{i,j} - I_{i,j}) + \alpha(u_{i,j} - u_{i-1,j}) - \alpha(u_{i+1,j} - u_{i,j}) + \alpha(u_{i,j} - u_{i,j-1}) - \alpha(u_{i,j+1} - u_{i,j}) = 0$$

...can be written as a large linear system of equations (schematic view)

$$\begin{pmatrix} 1+2\alpha & -\alpha & & -\alpha & & & \\ -\alpha & 1+3\alpha & -\alpha & & -\alpha & & & \\ & -\alpha & 1+3\alpha & -\alpha & & -\alpha & & \\ -\alpha & & -\alpha & 1+4\alpha & -\alpha & & -\alpha & \\ & & -\alpha & & -\alpha & 1+3\alpha & -\alpha & \\ & & & -\alpha & & -\alpha & 1+3\alpha & -\alpha & \\ & & & & -\alpha & & -\alpha & 1+2\alpha & \end{pmatrix} \begin{pmatrix} u_1 \\ u_2 \\ \vdots \\ \vdots \\ u_{N-1} \\ u_N \end{pmatrix} = \begin{pmatrix} I_1 \\ I_2 \\ \vdots \\ \vdots \\ I_{N-1} \\ I_N \end{pmatrix}$$

- $N \times N$ system matrix (for N pixels) is symmetric and positive definite
- It contains one main diagonal (central pixels) and four off-diagonals (for each of the four neighbors of a pixel)

$$\begin{pmatrix} 1+2\alpha & -\alpha & -\alpha & & & & \\ -\alpha & 1+3\alpha & -\alpha & & -\alpha & & & \\ & -\alpha & 1+3\alpha & -\alpha & & -\alpha & & \\ -\alpha & & -\alpha & 1+4\alpha & -\alpha & & -\alpha & \\ & & -\alpha & & -\alpha & 1+3\alpha & -\alpha & \\ & & & -\alpha & & -\alpha & 1+3\alpha & -\alpha \\ & & & & -\alpha & & -\alpha & 1+2\alpha \end{pmatrix} \begin{pmatrix} u_1 \\ u_2 \\ \vdots \\ \vdots \\ u_{N-1} \\ u_N \end{pmatrix} = \begin{pmatrix} I_1 \\ I_2 \\ \vdots \\ \vdots \\ I_{N-1} \\ I_N \end{pmatrix}$$

- The system matrix A is **sparse** (almost all entries are 0)
- Positive definite \rightarrow the inverse A^{-1} exists and we can solve for \mathbf{u}
- Questions:
 - When do we generally get a linear/nonlinear system with a unique solution?
 - How can this system be solved (efficiently)?

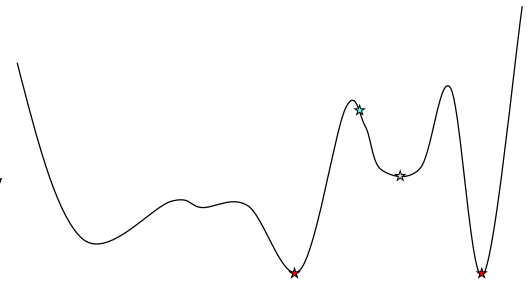
Convexity

Convex functions:

- Positive curvature
- No local minima
- Global minimum is unique
- Minimization by setting the derivative to 0 and solving the emerging linear or nonlinear system

Non-convex functions:

- Usually many local minima (and maxima)
- Global minimum may not be unique
- Global minimization is usually impossible, only heuristics exist

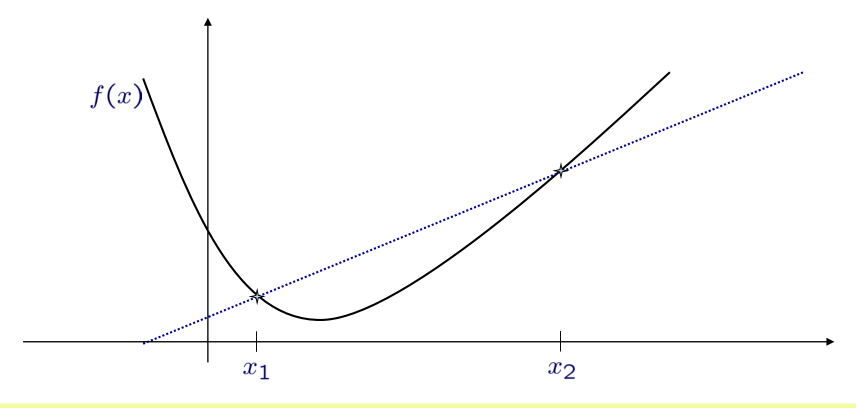


A function is convex if

$$f((1-\alpha)x_1+\alpha x_2) \le (1-\alpha)f(x_1)+\alpha f(x_2) \quad \forall x_1, x_2, \forall \alpha \in (0,1)$$

A function is strictly convex if

$$f((1-\alpha)x_1+\alpha x_2) < (1-\alpha)f(x_1)+\alpha f(x_2) \quad \forall x_1, x_2, \forall \alpha \in (0,1)$$



Theorem: every convex combination of (strictly) convex functions is again (strictly) convex

Proof:

$$h(x) := \gamma f(x) + \delta g(x) \quad \gamma, \delta \in \mathbb{R}, \quad \gamma, \delta \ge 0$$

$$h((1 - \alpha)x_1 + \alpha x_2) = \gamma f((1 - \alpha)x_1 + \alpha x_2) + \delta g((1 - \alpha)x_1 + \alpha x_2)$$

$$\le \gamma (1 - \alpha)f(x_1) + \gamma \alpha f(x_2) + \delta (1 - \alpha)g(x_1) + \delta \alpha g(x_2)$$

$$= (1 - \alpha)(\gamma f(x_1) + \delta g(x_1)) + \alpha(\gamma f(x_2) + \delta g(x_2))$$

$$= (1 - \alpha)h(x_1) + \alpha h(x_2)$$

Our energy function

$$E(u_{i,j}) := \sum_{i,j} \left((u_{i,j} - I_{i,j})^2 + \alpha \left((u_{i+1,j} - u_{i,j})^2 + (u_{i,j+1} - u_{i,j})^2 \right) \right)$$

is a convex combination of strictly convex (quadratic) functions

- → It is strictly convex as well
- → It has a unique global minimum

Solving the linear system

$$\begin{pmatrix}
1 + 2\alpha & -\alpha & & -\alpha & & \\
-\alpha & 1 + 3\alpha & -\alpha & & -\alpha & & \\
& -\alpha & 1 + 3\alpha & -\alpha & & -\alpha & & \\
& -\alpha & -\alpha & 1 + 4\alpha & -\alpha & & -\alpha & \\
& -\alpha & & -\alpha & 1 + 3\alpha & -\alpha & & \\
& & -\alpha & & -\alpha & 1 + 3\alpha & -\alpha & \\
& & & -\alpha & & -\alpha & 1 + 2\alpha & & \\
\end{pmatrix}
\begin{pmatrix}
u_1 \\ u_2 \\ \vdots \\ \vdots \\ u_{N-2} \\ \vdots \\ \vdots \\ u_{N-1}
\end{pmatrix} = \begin{pmatrix}
I_1 \\ I_2 \\ \vdots \\ \vdots \\ I_{N-2} \\ I_{N-1}
\end{pmatrix}$$

- The two additional off-diagonals (together with the size of matrix) rule out Gauss-elimination (in 1D, however, Gauss-elimination is very efficient).
- An iterative solver is needed to preserve the sparsity of the matrix
- Simplest iterative solver: Jacobi method
- Converges if the matrix is strictly diagonal dominant

$$|a_{ii}| > \sum_{i \neq j} |a_{ij}| \quad \forall i$$

Decompose the matrix into its diagonal part D and its off-diagonal part M

$$A = D + M$$

For the linear system this means

$$Ax = b \Leftrightarrow (D+M)x = b \Leftrightarrow Dx = b-Mx$$

- D^{-1} can be computed very easily: just replace the diagonal elements by their inverse.
- Now we can compute the solution x iteratively. Starting with any initialization x^0 , iterate

$$x^{k+1} = D^{-1}(b - Mx^k)$$

Iterate until the norm of the **residual** $r^k := Ax^k - b$ is smaller than a threshold or the change in the solution $(x^{k+1} - x^k)^2$ becomes small. When the change is 0, the iterate has **converged**.

Jacobi method

- Advantages:
 - Simple
 - Can be implemented in parallel
- Disadvantages:
 - Slow
 - Convergence only for $k \to \infty$
 - Computation not in-place
- Alternatives:
 - Gauss-Seidel, Successive Over-relaxation (faster, in-place)
 - Conjugate gradient (convergence after finite number of iterations)
 - Multigrid methods (sometimes much faster)

 Split the off-diagonal part M into the lower triangle L and the upper triangle U

$$Ax = b \Leftrightarrow Dx = b - Lx - Ux$$

• During iteration, traverse the vector x from top to bottom and use already the new values for multiplication with the lower triangle

$$x^{k+1} = D^{-1}(b - Lx^{k+1} - Ux^k)$$

In our denoising example this reads

$$u_i^{k+1} = \frac{I_i + \alpha \sum_{j \in \mathcal{N}^-(i)} u_j^{k+1} + \alpha \sum_{j \in \mathcal{N}^+(i)} u_j^k}{1 + \sum_{j \in \mathcal{N}(i)} \alpha}$$

- Converges if A is positive or negative definite
- For already updated neighbors take the new value → in-place computation
- Recursive propagation of information → faster

Successive over-relaxation (SOR)

• Emphasize the Gauss-Seidel idea by over-relaxing the new solution

$$x^{k+1} = (1 - \omega)x^k + \omega D^{-1}(b - Lx^{k+1} - Ux^k)$$

- For $\omega = 1$ this is the Gauss-Seidel method
- Converges for positive- or negative-definite matrices (all eigenvalues positive or negative, respectively), if $\omega \in (0,2)$
- Over-relaxation for $\omega > 1$: faster convergence
- Under-relaxation for $\omega < 1$: can help establish convergence in case of divergent iterative processes
- Optimal ω must be determined empirically

Conjugate gradient (CG)

- Two non-zero vectors u and v are **conjugate** with respect to A if the inner product $\langle u,v\rangle_A:=u^\top Av=0$. This means the two vectors are orthogonal with respect to this special scalar product.
- A set of n conjugate vectors $\{p_k\}$ forms a basis of \mathbb{R}^n , so the solution x^* of Ax = b can be expanded as $x^* = \alpha_1 p_1 +, ..., +\alpha_n p_n$
- The coefficients α_k are derived as follows:

$$Ax^* = \alpha_1 A p_1 + \ldots + \alpha_n A p_n = b$$

$$p_k^\top A x^* = p_k^\top \alpha_1 A p_1 + \ldots + p_k^\top \alpha_n A p_n = p_k^\top b \quad \text{(expansion with } p_k \text{)}$$

$$\alpha_k = \frac{p_k^\top b}{p_k^\top A p_k}$$

- After n computations we obtain the exact solution x^*
- Good choice of $\{p_k\}$ \rightarrow few coefficients approximate the solution well

Conjugate gradient: iteratively assembling the basis

- Start with some initial point x^0
- Let p_0 be the residual $r_0 = b Ax^0$. This is the gradient of

$$E(x) = \frac{1}{2}x^{\top}Ax - b^{\top}x$$

the minimizer of which is x^* . Therefore the name conjugate gradient.

Iteratively compute:

$$\alpha_{k} = \frac{r_{k}^{\top} r_{k}}{p_{k}^{\top} A p_{k}} \qquad x^{k+1} = x^{k} + \alpha_{k} p_{k} \qquad r_{k+1} = r_{k} - \alpha_{k} A p_{k}$$

$$\beta_{k} = \frac{r_{k+1}^{\top} r_{k+1}}{r_{k}^{\top} r_{k}} \qquad p_{k+1} = r_{k+1} + \beta_{k} p_{k}$$

Stop when residual is small. Guaranteed solution after n iterations.

Conjugate gradient: convergence and preconditioning

- Matrix A must be symmetric and positive definite
- Usually in image processing, computing the exact solution is not an option since n is the number of pixels
- Number of iterations needed to get a good approximate solution depends on the **condition number** of *A* (largest vs. smallest eigenvalue). The same holds for the other iterative methods (Gauss-Seidel, etc.)
- Sometimes so-called **preconditioners** P^{-1} are used to have a small condition number for $P^{-1}A$

$$Ax = b \Leftrightarrow P^{-1}Ax = P^{-1}b$$

Simplest preconditioner: Jacobi preconditioner

$$P = D \quad \Leftrightarrow \quad P^{-1} = D^{-1}$$

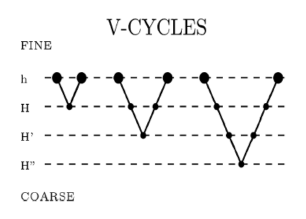
$$Ax = b \quad \Leftrightarrow \quad D^{-1}Ax = D^{-1}b$$

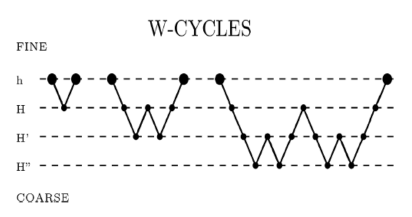
- All previous linear solvers have the drawback that they only act locally.
- This is due to the sparsity of the matrix: one iteration only distributes information at a pixel to its four neighbors (the recursive nature of Gauss-Seidel relaxes this statement a bit).
- Idea of multigrid solvers: shorten distances by regarding the system from a coarser point of view
- Additional effect: coarse versions of the system have fewer entries → iterations are faster at coarse levels
- Different types of multigrid solvers
 - Unidirectional (cascadic) multigrid
 - Bidirectional (correcting) multigrid
 - Full multigrid (a combination of both)

Unidirectional (cascadic) multigrid

- Create downsampled versions of the linear system (usually by downsampling the image and deriving the linear system from this)
- Compute first approximate solution at the coarse grid (e.g. with SOR)
- Take upsampled result as initial guess for the next finer grid
- Refine result there (again with SOR)
- Advantages:
 - Iterations at the coarse level are very fast
 - Simple implementation
- Disadvantage:
 - Coarse level systems often do not approximate the original system well
 Iterations at coarse levels do not really help

- Basic idea: do not downsample the image <u>but the error</u>
- Compute first solution at fine grid
- Correct error at coarse grid
- Refine result at finer grid

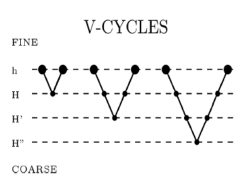




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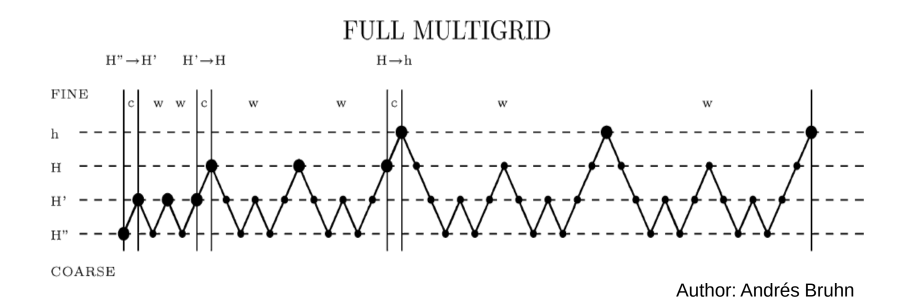
Correcting multigrid in detail

- 1. Presmoothing relaxation step $A^h x^h = b^h$
 - Run some iterations at the fine grid
 - Yields approximate solution $ilde{x}^h$
 - Remaining error: $e^h = x^h \tilde{x}^h$



- Correction step:
 - Goal: compute error $A^h e^h = r^h$ $r^h = b^h A^h \tilde{x}^h$
 - Local part of error already removed
 → Solve this system at coarser grid A^He^H = r^H
 - Transfer error to fine grid and correct the solution $\tilde{\tilde{x}}^h = \tilde{x}^h + \tilde{e}^h$
- 3. Postsmoothing relaxation step
 - Apply some further iterations at fine grid to remove local errors introduced by \tilde{e}^h

- Combination of cascadic and correcting multigrid
- Start at coarse grid with downsampled image
- At each finer level apply a W-cycle



Integer problems

- Here we were considering problems with continuous variables (each vector component of the solution is a real number)
- Segmentation and matching typically leads to integer problems
- These are combinatorial problems, only few of them being solvable in polynomial time
- Some typical problems arising in computer vision are:
 - Linear programs (LP)
 - Integer quadratic programs (IQP) including special cases like min-cut
 - Second order cone programs (SOCP)
- More in the Computer Vision course or in Algorithm Theory

Summary

- The energy minimization framework is a sound way to model and solve image processing problems
- All model assumptions are clearly stated
- Global optimization is "easy" if the energy function is convex
- The necessary condition for a minimum is that the gradient is zero
- Leads to a large, but sparse, linear or nonlinear system of equations
- There are several methods to solve sparse linear systems iteratively, some are easier to implement, others are faster

Literature

- D. Young: Iterative Solution of Large Linear Systems, Academic Press, 1971.
 Reprint by Dover 2003.
- J. R. Shewchuk: An introduction to the Conjugate Gradient method without the agonizing pain. CMU Technical Report, 1994.