Exercise 4

Deadline: 13.06.2023, 16:00

This exercise is dedicated to linear regression and its efficient implementation for large sparse matrices. We will test this method on the relevant application of computer tomography.

Regulations

Please hand in your solution as a Jupyter notebook tomography.ipynb, accompanied with exported tomography.html. It is important that you stick to these file naming conventions! Zip all files into a single archive ex04.zip and upload this file to MaMPF before the given deadline. Moreover, please set your Anzeigename/display name and Name in Uebungsgruppen/name in tutorials in MaMPF to your real name, which should be identical to your name in muesli and make sure you join the submission of your team via the invitation code before the submission deadline. Check out https://mampf.blog/handing-in-homework-assignments for instructions.

Setting

You find yourself in a two-dimensional world in the role of a medical doctor. A patient (we will call him H.S. here for anonymity) walks into your hospital and complains about a headache. A close examination of the head provides no exterior clues as to the cause of the pain, and you soon realize that you will have to perform computer tomography. Unfortunately, the computer for processing recorded data is broken. Nevertheless, you let H.S. enter the tomograph and take X-ray images of his head from several angles. You send H.S. back to the waiting room and start to think about how you might process the scans using only your private laptop.

The required data files are available at https://tinyurl.com/HD-EML-hs-tomography-zip and as an external link on MaMPF.

Preliminaries

For the scans, H.S.'s 2D head was placed between an X-ray source and a 1D sensor array. The sensor elements record the accumulated signals of many parallel rays passing through the head (see left side of figure 1). To reconstruct all the internal structures, multiple measurements need to be taken from different angles α . The resulting tomogram is an image $\mu \in \mathbb{R}^{M \times M}$ where each pixel encodes the local X-ray absorption coefficient of the patient's head (bright for bones, gray for soft tissue, black for air). The least-squares reconstruction algorithm will create a flattened version $\beta \in \mathbb{R}^D$ of μ that needs to be reshaped into a 2D image to be displayed. The pixels μ_{j_a,j_b} are related to the vector elements β_j by the formula $j = j_a + M j_b$, as shown in figure 1.

The sensor consists of a 1-dimensional array of N_p sensor elements that register parallel X-rays, and the emitter-detector setup is rotated into N_o different orientations α_{i_o} . The measured intensities are commonly displayed as a 2D image called a *sinogram*, see figure 3. For least-squares reconstruction, we flatten the sinogram $\boldsymbol{s} \in \mathbb{R}^{N_o \times N_p}$ into the response vector $\boldsymbol{y} \in \mathbb{R}^N$. Its elements \boldsymbol{y}_i are defined by the relation $i = i_p + N_p i_o$, where i_p and i_o are the indices of the sensor elements and sensor orientations respectively.

We can now describe the relation between the tomogram β and the measured intensities \mathbf{y} as a linear projection with projection matrix $\mathbf{X} \in \mathbb{R}^{N \times D}$:

$$\mathbf{X} \cdot \boldsymbol{\beta} = \mathbf{y} \tag{1}$$

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Intuitively, entry \mathbf{X}_{ij} should be interpreted as a weight which encodes how much the sensor response \mathbf{y}_i is influenced by the absorption at pixel $\boldsymbol{\beta}_i$.

In the continuous domain, the measured intensity I_{sensor} of a single ray **r** depends on how much of the original radiation I_{source} is absorbed by the material between emitter and sensor. This can be modelled as

$$I_{\text{sensor}} = I_{\text{source}} \cdot \exp\left(-\int_{a,b \in \mathbf{r}} \mu(a,b) \,\mathrm{d}a \,\mathrm{d}b\right),\tag{2}$$

where μ describes the material's absorption properties at each spatial position (a, b) and can be seen as a continuous version of the discrete tomography image μ defined above. For simplicity, we assume that the raw sensor measurements are already subjected to preprocessing which computes the logarithm of this formula, so that the response is defined by

$$y = \ln\left(\frac{I_{\text{source}}}{I_{\text{sensor}}}\right) \tag{3}$$

$$= \int_{a,b \in \mathbf{r}} \mu(a,b) \,\mathrm{d}a \,\mathrm{d}b. \tag{4}$$

For practical computation, we need to discretize this formula. Each sensor element is essentially a bin collecting radiation from many parallel rays. We assume that a ray intersecting the sensor array inbetween two bins distributes its intensity among both. The ratio of distribution is determined in a linear fashion by how close the intersection is to the center of either bin.

Example: Consider the ray passing through β_{19} at angle α_{i_o} , as illustrated in figure 1, which intersects the sensor array at position 4.35 (between elements 4 and 5). Sensor element 4 captures 65% of this ray's intensity and sensor element 5 captures 35%. This is encoded in the weight matrix **X** by setting $\mathbf{X}_{4+N_pi_o,19} = 0.65$ and $\mathbf{X}_{5+N_pi_o,19} = 0.35$.

Thanks to this definition, the integral in equation 4 can be discretized into a sum, and writing the sum in matrix notation gives the linear system of equation 1.

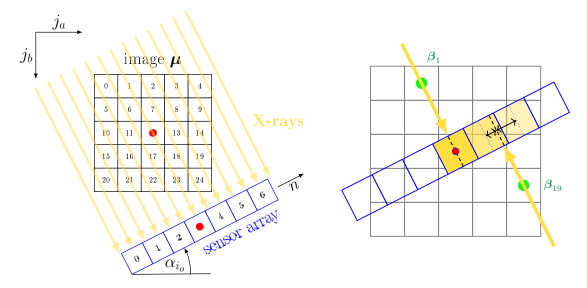


Figure 1: Sketch of the projection of a 5×5 -image μ onto a 7 pixel sensor array that has been rotated around the center of the image by the angle α_{i_o} (left). Small numbers reflect the indices j within β and i_p within one slice of \mathbf{s} , respectively. While both the X-ray source and the sensors are naturally situated outside the tissue, the projection can be made easier by virtually placing the sensor at the image center and considering the rays to come from both directions (right). In any case, for each individual coordinate in β , we trace the ray passing through that point and calculate how much of its intensity is collected by which sensor pixels.

1 Constructing the matrix X (22 points)

With only one projection, solving equation 1 is ill-posed and a useful reconstruction of β is impossible. The problem becomes well-posed by using sufficiently many projections of β under different angles α , whose recordings are just concatenated in the flattened measurement vector \mathbf{y} or the 2D sinogram (figure 3), respectively. Constructing the corresponding weight matrix \mathbf{X} is the main difficulty of this exercise.

Implement a function $X = \text{construct}_X(M, \text{alphas}, Np = None)$ which, given the desired tomogram size $D = M \times M$, a list of measurement angles $(\alpha_0, \alpha_1, \ldots)$ (in degrees) and an optional sensor resolution N_p , returns the matrix X, which completely describes the setup of our simplified CT scanner.

If Np is not given, choose a value large enough to fit the diagonal of the image β , like $N_p = \lceil \sqrt{2} \cdot M \rceil$. Choosing an odd number of sensor elements will make it easier to align the image coordinates with the coordinates of the sensor array: You can define a common coordinate origin and then find ray intersections by projecting the tomogram's pixel coordinates along the current ray orientation onto the rotated sensor array, see figure 1.

In the examination scenario for patient H.S., we have M = 195, i.e. the tomogram has D = 38025 pixels. The sensor has $N_p = 275$ bins, and intensities have been measured at $N_o = 179$ projection angles, resulting in a response vector of size N = 49225. Thus, matrix **X** has shape 49225×38025 (almost 2 billion entries), and you will run into trouble if you implement construct_X() naively.

Fortunately, the intensity of each ray is only distributed among the *two* sensor elements closest to the ray's intersection with the sensor. Consequently, each image pixel can contribute to no more than two sensor elements under any given measurement angle (see right side of figure 1) and the vast majority of entries of **X** are actually zero. This is called a *sparse matrix*, and the scipy module scipy.sparse¹ provides powerful tools to take advantage of this property: it allows you to save memory by offering spare matrix classes that store only the non-zero entries, and to skip unnecessary calculations by offering specialized implementations of linear algebra functionality. Specifically, the class coo_matrix ("COOrdinate" format) is recommended to create **X**, and the class csc_matrix ("Compressed Sparse Column" format) is best for solving the linear system. Converting between different matrix types is easy and results in faster code than using the same type throughout.

However, using a sparse matrix is not enough to make your code efficient. You should also vectorize the function construct_X() (cf. exercise 1). A good way of doing this is to create an array $\mathbf{C} \in \mathbb{R}^{2 \times D}$ holding the coordinates of the tomogram's pixel centers, i.e.

 $\mathbf{C}_{0j} = a_0 + j_a \cdot h \text{ (the } a \text{ coordinate of pixel } j)$ $\mathbf{C}_{1j} = b_0 + j_b \cdot h \text{ (the } b \text{ coordinate of pixel } j)$

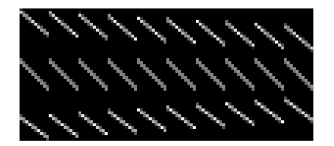
with h being the pixel distance and $j = j_a + M j_b$ as defined earlier (numpy.mgrid is useful here). In this case, the pixel distance for both image and sensor is simply h = 1. The current orientation can be expressed by a unit vector \mathbf{n} along the rotated sensor, and the projection \mathbf{p} of each pixel onto the sensor is simply a linear projection according to

$$\mathbf{p} = \mathbf{n}^\top \mathbf{C} + s_0,$$

where s_0 is the distance between the first sensor element and the sensor's coordinate origin. The weights **X** for each angle can now be computed from **p** by numpy's vector functions. In the end, the valid elements of **X** are represented by their indices (i, j) and the corresponding weights. The constructor of class coo_matrix expects this information to be provided by three 1-dimensional arrays i_indices, j_indices, and weights, resulting in the call

X = coo_matrix((weights, (i_indices, j_indices)), shape=(N, D), dtype = numpy.float32),

¹Install scipy via conda as usual.



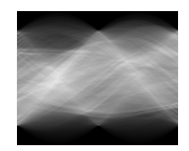


Figure 2: Visualization of matrix X for M = 10, $N_p = 15$ and three projections at angles (-33, 1, 42). You can also find this matrix in the data-zip as 'hs_tomography/X_example.npy'.

Figure 3: Sinogram visualization of y. Each column shows the sensor response for a different angle $\alpha \in [-90, \ldots, 90]$.

where we used element type float32 to save even more memory. Check the scipy documentation for more details about how this works. In our example implementation, constructing the sparse 49225×38025 matrix takes about 5 seconds on a standard laptop.

As in real-world scenarios, there is some ambiguity left to this task: Where is the origin of the coordinate system, which direction is $\alpha = 0$, and how is **y** oriented? The answers to these questions determine the appropriate indexing order and sign of the expressions, and figuring out these details is part of your task (emulating what often happens in the real world). To check whether your matrix construction is correct, visualize the result of construct_X(10, [-33, 1, 42]) with pyplot.imshow() and compare it to the matrix shown in figure 2, which you find in the data-zip under 'hs_tomography/X_example.npy'². Note that you will have to convert your sparse matrix to a dense numpy-array for visualization.

2 Recovering the image (9 points)

The material for this sheet contains the list of angles and the measured sensor data **y** for two versions of the experiment. The smaller one was created with M = 77, $N_p = 109$ and 90 projection angles. The larger one was created with M = 195, $N_p = 275$ and 179 projection angles. The exact set of angles and the corresponding response vectors **y** can be found in the data-zip as 'alphas_77.npy' and 'y_195.npy' resp. 'alphas_195.npy' and 'y_77.npy' (in folder 'hs_tomography'). We recommend the smaller version of the data for debugging your code. However, its resolution is insufficient to diagnose the cause for your patient's headache. If you can recognize the head in the small tomogram, your code is probably correct, and you should switch to the larger version.

Exactly how many non-zero entries does \mathbf{X} have? Use scipy's tools to find out and report the *sparsity* of \mathbf{X} .

With the ability to construct the matrix \mathbf{X} and having obtained a set of projections \mathbf{y} , you are now able to reconstruct the original image $\boldsymbol{\beta}$. In theory, you could try to solve the equation system directly via the pseudo-inverse

 $\boldsymbol{\beta} = (\mathbf{X}^{\top} \mathbf{X})^{-1} \cdot \mathbf{X}^{\top} \cdot \mathbf{y},$

but this is not recommended because it ignores sparsity and is therefore very slow.

Fortunately, scipy already gives you access to efficient solvers for sparse linear equation systems! Find out how to use scipy.sparse.linalg.lsqr() to obtain the least-squares solution to your problem. The solver's low default tolerance parameters atol = 1e-08 and btol = 1e-08 lead to high quality solutions, but also long computation times on the large version of the data. A good

² .npy' is numpy's matrix file format. You can load it with X_ex = np.load('hs_tomography/X_example.npy')

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trade-off might be atol = btol = 1e-05, but you can reduce the tolerance once your are confident that your code is correct.

Reconstruct the tomogram and plot it as a 2D image. Give a diagnosis on what causes H.S.'s headache and propose a treatment.

If you didn't manage to correctly construct X in task 1, you can find a precomputed weight matrix for the smaller version of the experiment in 'hs_tomography/X_77.npy' and use it for this task. You can load it with $X = np.load('hs_tomography/X_77.npy', allow_pickle=True)$ and convert it to a sparse matrix with $X = scipy.sparse.csc_matrix(X.all())$.

3 Minimizing the radiation dose (9 points)

As a doctor, you do not want to expose your patients to unnecessary radiation. Try to reduce the number of projection angles in a sensible way and visualize how this changes the quality of the reconstruction. In the case of H.S., what would you say is the minimal number of projections that still allows you to resolve the cause of his headache?