

# HUMBOLDT UNIVERSITÄT ZU BERLIN SEMINAR PAPER

# Numerical Methods for solving Eigenvalue-Problems

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Numerical Introductory Course

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#### 1 Motivation

#### Abstract

Eigenvalues and eigenvectors are often the solution to multidimensional optimization problems, however computing them by hand for anything but trivial matrices is most of the time infeasible or inpractical. To this extend we would like to deploy an automated procedure which yields the correct eigenvectors and eigenvalues. We demonstrate the relevance of eigenvalues and eigenvectors by revising two applications from statistics, Principal Component Analysis and Fisher's Linear Discriminant Analysis, which we follow up by investigating four algorithms suited for eigenvalue problems. Finally we provide a compound solution that takes advantage of each algorithms strengths.

For many statistical applications eigenvectors provide a formidable solution. Be it dimensionality reduction in terms of a Principal Component Analysis or classification by Fisher's Linear Discriminant Analysis, both come in the guise of optimization problems. But what are eigenvalues and eigenvectors?

If A is an  $n \times n$  matrix, v is a non-zero vector and  $\lambda$  is a scalar, such that

$$Av = \lambda v \tag{1}$$

then v is called an *eigenvector* and  $\lambda$  is called an *eigenvalue* of the matrix A. An eigenvalue of A is a root of the characteristic equation,

$$det (A - \lambda I) = 0. (2)$$

Geometrically speaking, we require a vector which, when multiplied by matrix A, will not get rotated but only elongated by a factor  $\lambda$ .

When confronted with a high-dimensional data matrix  $X \in \mathbb{R}^{n \times m}$  an analyst often wishes to find a lower-dimensional representation, while conserving as much of the structure as possible. One way of achieving this goal is to choose a standardized linear combination of features that aim to maximize the variance of the projection  $\delta' X$ . We can formalize this as

$$\max \delta' Var(X) \delta s.t. \sum \delta_i^2 = 1.$$
 (3)

where  $X \in \mathbb{R}^{n \times m}$ ;  $m, n \in \mathbb{N}$ ;  $\delta \in \mathbb{R}^m$ . The Lagrangean that corresponds to the constrained maximization problem in 3 is

$$\mathcal{L}(Var(X), \delta, \lambda) = \delta' Var(X) \delta - \lambda (\delta' \delta - 1),$$

where  $\lambda \in \mathbb{R}^m$ 

Taking derivatives we obtain the first order condition:

$$\frac{\partial \mathcal{L}}{\partial \delta} \stackrel{!}{=} 0$$
$$2Var(X)\delta - 2\lambda_k \delta \stackrel{!}{=} 0$$
$$Var(X)\delta = \lambda_k \delta$$

Which is now reduced to a common Eigenvalue problem as posed in (1).

$$Y = \Gamma'(X - \mu) \tag{4}$$

where  $Y \in \mathbb{R}^{n \times m}$  is the matrix of rotations,  $\Gamma \in \mathbb{R}^{m \times m}$  is the matrix of eigenvectors,  $\mu \in \mathbb{R}^m$  is the vector of sample means. [Härdle and Simar, 2015]

In section two we lay out the mathematical foundations for the operations we are about to perform. In particular, we will try to reformulate any complicated eigenvalue problem into a straightforward one by diagonalizing the matrix in question, without altering the eigenvalues we would like to compute. We follow these justifications by proposing two main algorithms for computing eigenvalues, first the Jacobi-Method for symmetric matrices, then the QR-Method for arbitrary square matrices in section 3. Additionally, for the QR-Method we define two extensions which try to increase the initial QR-algorithm's speed. For all algorithms we provide implementations in the Python-programming-language [van Rossum, 1995, Hunter, 2007, McKinney, 2010]. In section 4 we will analyse the implemented routines by critically reflecting upon the accuracy of the obtained results as well as their efficiency. In the final section we provide a final algorithm which combines the strengths

of the defined procedures by chosing the algorithm that is most fit for the underlying problem.

## 2 Similarity Transformations

In general we want to reformulate the eigenvalue problem of a complicated matrix into an eigenvalue problem of a simple matrix, which yields the same eigenvalues. Simple matrices in our case will be diagonal matrices, since with them it is possible to identify their eigenvalues simply as entries on the main diagonal. Such a transformation that conserves the eigenvalues of a matrix is called a *similarity transformation*.

Two  $n \times n$  matrices A and B are called *similar* if there exists an invertible matrix P such that

$$A = P^{-1}BP. (5)$$

It is obvious that the similarity relationship is commutative as well as transitive. If A and B are similar, it holds that

$$B - \lambda I = P^{-1}BP - \lambda P^{-1}IP$$
$$= A - \lambda I.$$

Hence A and B have the same eigenvalues. This fact also follows immediately from the transitivity of the similarity relationship and the fact that a matrix is similar to the diagonal matrix formed from its eigenvalues, as stated in the spectral-decomposition. Important types of similarity transformations are based around orthogonal matrices. If Q is orthogonal and

$$A = Q'BQ$$
,

A and B are called orthogonally similar [Gentle, 1998]. We will use orthogonal similarity transformations to diagonalize matrices we wish to know the eigenvalues of.

# 3 Algorithms

#### 3.1 Jacobi Method

The Jacobi-Method for computing the eigenvalues of a symmetric matrix  $A \in \mathbb{R}^{n \times n}$  deploys a sequence of orthogonal similarity transformations that eventually results in

$$A = P\Lambda P^{-1} \Leftrightarrow \Lambda = P^{-1}AP$$

where  $\Lambda$  is diagonal and P consists of a sequence of matrix multiplications  $P = \prod_{k=1}^{K} V_{p_k,q_k}(\theta_k)$  and  $V_{p_k,q_k}(\theta_k)$  is of the form proposed in (18). More specific the *Jacobi iteration* is

$$A^{(k)} = V'_{p_k,q_k}(\theta_k)A^{(k-1)}V_{p_k,q_k}(\theta_k),$$
(6)

where  $p_k, q_k$  and  $\theta_k$  are chosen such that  $A^k$  resembles more a diagonal matrix than  $A^{k-1}$ . Specifically they will be chosen as to reduce the sum of squares of the off-diagonal elements. As we saw in (18) it is easy to chose an angle  $\theta_k$  in order to introduce a zero in a single Givens rotation. Here we use the rotations in the context of a similarity transformation, so it is a little more complicated.

We require that  $a_{pq}^{(k)} = 0$ , this implies

$$a_{pq}^{(k-1)}(\cos^2\theta - \sin^2\theta) + \left(a_{pp}^{(k-1)} - a_{qq}^{(k-1)}\right)\cos\theta\sin\theta = 0.$$
 (7)

Jacobi Method **ALGORITHMS** 3.1

Demonstrate jacobi on a 5x5 matrix Iteration: 10 Iteration: 75

Figure 1: Progress Jacobi-Method •

We can use the trigonometric identities

$$\cos(2\theta) = \cos^2\theta \sin^2\theta$$

$$\sin(2\theta) = 2\cos\theta\sin\theta,$$

in (7) we have

$$\tan(2\theta) = \frac{2a_{pq}^{(k-1)}}{a_{pp}^{(k-1)} - a_{qq}^{(k-1)}}.$$

From this we can retrieve the angle and obtain the rotation matrix in each iteration [Gentle, 1998].

The algorithm converges if the off-diagonal elements are sufficiently small. The best index pair at a given iteration is the pair (p,q) that satisfies

$$|a_{pq}^{(k-1)}| = \max_{i < j} |a_{ij}^{(k-1)}|.$$

If this choice is made, the Jacobi Method can be shown to converge [Gentle, 1998].

Figure 1 visualizes the progress of the *Jacobi*-method on a symmetric  $5 \times 5$ matrix. As we can see, in the first iteration the element  $a_43$  is eliminated. In the subsequent operations the Jacobi-method continues to eliminate any non-zero entries on the off-diagonal until the algorithm convergences after 10 iterations.

#### Algorithm 1 jacobi Q

**Require:** symmetric matrix A

Ensure: 0 < precision < 1

initialize:  $L \leftarrow A; U \leftarrow I; L_{max} \leftarrow 1$ 

1: while  $L_{max} > precision$  do

Find indices i, j of largest value in lower triangle of abs(L)

3:  $L_{max} \leftarrow L_{i,j}$ 

 $\alpha \leftarrow \frac{1}{2} \cdot \arctan(\frac{2A_{i,j}}{A_{i,i} - A_{j,j}})$ 

5:

 $V_{i,i}, V_{j,j} \leftarrow \cos \alpha; V_{i,j}, V_{j,i} \leftarrow -\sin \alpha, \sin \alpha$   $A \leftarrow V'AV; U \leftarrow UV$ 6:

7:

8: end while

9: **return** diag(A), U

#### QR-Method 3.2

The most widely used algorithm to extract eigenvalues is the so called *QR*-method. The most important advantage of the QR-method over the Jacobi-method is that it can be applied to non-symmetric matrices. Note however, that it is simpler for symmetric matrices, since the eigenvalues are real-valued.

The QR-method to extract the eigenvalues of a square matrix  $A \in \mathbb{R}^{n \times n}$  is performed by first computing the titular QR decomposition of A.

$$A = QR, (8)$$

where Q is an orthogonal and R is an upper triangular matrix. Then define the QR iteration as

$$A^{k} = Q'_{k-1}A_{k-1}Q_{k-1} = R_{k-1}Q_{k-1}$$
(9)

Note hereby that all matrices in the sequence  $\{A_k\}$  share the same eigenvalues, since this procedure is a similarity transformation due to Q's orthogonality. Additionally, in an implementation it is usually preferable to compute the QR-iteration in the way shown at the rightmost part of equation (9) [Börm and Mehl, 2012]. Although, mathematically, each statement is exactly identical there is a practical difference in due to computational imperfections and limited machine precision.

Figure 2: Progress basic QR-Method Demonstrate qrm1 on a 5x5 matrix Iteration: 0 Iteration: 1

Iteration: 10 Iteration: 75

Iteration: 75

Iteration: 75

The reason being that the computation of  $Q'_{k-1}A_{k-1}Q_{k-1}$  obviously requires two matrix multiplications whereas the result of  $R_{k-1}Q_{k-1}$  can be readily obtained by one. When combining multiple steps over a long sequence of QR iterations the additional computations lead to additional rounding errors, which can have an influence on the accuracy of the obtained results. Additionally, less computations lead of course to a faster procedure in general.

Figure 3.2 visualizes the progress of the basic QR-method on the same  $5 \times 5$  matrix as in Figure 1. Compared to the Jacobi-method it does not explicitly pick a single element that will be eliminated per iteration. Instead, the QR-method extracts the eigenvalues by a process that is called "chasing". By that we mean that alternating steps are being performed, which create non-zero eintries in positions (i+2,i), (i+3,i) and (i+3,i+1) and restore them to zero, as the nonzero entries are moved farther down the matrix [Gentle, 1998]. We can also see that compared to the Jacobi-Method, so far, the QR-algorithm lacks in speed. Where the Jacobi-method was almost done diagonalizing the matrix in iteration 10, the basic QR-algorithm still had multiple non-zero entries left. Thus we would like to make minor improvements on the algorithm's efficiency.

```
Algorithm 2 QRM1 \bigcirc

Require: square matrix A
  initialize: conv \leftarrow False

1: while not conv do

2: Q, R \leftarrow QR-Factorization of A

3: A \leftarrow RQ

4: if A is diagonal then

5: conv \leftarrow True

6: end if

7: end while

8: return diag(A), Q
```

#### 3.2.1 Hessenberg Variant

In order to speed up the QR-method it is advisable to transform the matrix to its upper Hessenberg form. A matrix A is of upper Hessenberg form if it is upper triangular except for the first subdiagonal, which may be non-zero. In particular  $a_{ij} = 0 \ \forall i > j+1$ :

$$\begin{bmatrix} X & X & X & \dots & X & X \\ X & X & X & \dots & X & X \\ 0 & X & X & \dots & X & X \\ 0 & 0 & X & \dots & X & X \\ \vdots & \vdots & & \ddots & \vdots & \vdots \\ 0 & 0 & 0 & \dots & X & X \end{bmatrix}$$

A matrix can be reduced to Hessenberg form in a finite number of similarity transformations using Householder transformations or Givens rotations. For symmetric matrices the transformation into a Hessenberg-form results in a tridiagonal matrix. But even for non-symmetric matrices, the Hessenberg-form allows a large saving in subsequent computations. After the transformation we can deploy the previously defined QR-method [Gentle, 1998].

Figure 3: Progress Hessenberg-QR-Method

Demonstrate qrm2 on a 5x5 matrix Iteration: 0

Iteration: 1

Iteration: 10

Iteration: 75

Iteration: 75

Iteration: 75

Figure 3 visualizes the progress of the Hessenberg variant of the QR-method. In order to make it comparable to the previous algorithms the same  $5 \times 5$ -dimensional matrix is being evaluated. We can readily see, that the transformation to Hessenberg-form results in a tridiagonal matrix. This facilitates computations and explains the vastly improved resulting matrix after 10 iterations compared to the basic QR-method. However, it still does not match the progress of the Jacobi-method after the same number of iterations.

#### Algorithm 3 QRM2 Q

Require: square matrix A1:  $A \leftarrow \text{hessenberg}(A)$ 2: continue with: QRM1(A)

#### 3.2.2 Accelerated Variant

We could already improve the QR-method and cut down on computational cost. However, we still cannot match the results of the Jacobi method. To this effect we present the final adjustment on the QR-method to improve convergence speed. The general idea is, that we deliberately create an additional zero entry on the main diagonal by subtracting a scalar on each element, perform the QR-iteration and finally undo the subtraction. In particular, we define

$$T^{m} = \begin{bmatrix} \alpha_{1}^{m} & \beta_{1}^{m} & 0 & 0 & \dots & 0 \\ \beta_{1}^{m} & \alpha_{2}^{m} & \beta_{2}^{m} & & & & \\ 0 & \beta_{2}^{m} & \alpha_{3}^{m} & \beta_{3}^{m} & & \vdots & & \\ & & \ddots & \ddots & \ddots & & \\ & & & \beta_{n-2}^{m} & \alpha_{n-1}^{m} & \beta_{n-1}^{m} \\ 0 & & & \beta_{n-1}^{m} & \alpha_{n}^{m} \end{bmatrix}$$

$$(10)$$

$$T^{m} = T - t_{n,n}I$$
$$T^{m} = QR$$
$$T^{m+1} = T^{m} + t_{n,n}I$$

After this we can define the accelerated iteration step as

$$R_m = Q_m' \left( T_m - \alpha_n^m I \right) \tag{11}$$

$$T_{m+1} = Q_m' \left( T_m - \alpha_n^m I \right) Q_m + \alpha_n^m I \tag{12}$$

$$=Q_m'T_mQ_m\tag{13}$$

Again  $T_{m+1}$  is similar to  $T_m$ .

Figure 4: Progress Accelerated QR-Method

Demonstrate qrm3 on a 5x5 matrix
Iteration: 1

Iteration: 1

Iteration: 75

Iteration: 75

Figure 4 visualizes the progress of the accelerated QR-method. For comparability, the matrix used is the same  $5 \times 5$  matrix as before. Most notably is, that the accelerated method still performs worse than the Jacobi method. So far results seem similar compared to the Hessenberg variant of the QR-method. The case can be made that the accelerated method performs considerably better than the basic QR-method and slightly better than the Hessenberg variant after 10 iterations than the. Howbeit, this claim warrants further analysis.

#### Algorithm 4 QRM3 Q

```
Require: square matrix A \in \mathbb{R}^{p \times p}

1: T \leftarrow \text{hessenberg}(A), conv \leftarrow False

2: while not conv do

3: Q, R \leftarrow \text{QR-Factorization of } T - t_{p-1,p-1}I

4: T \leftarrow RQ + t_{p-1,p-1}I

5: if T is diagonal then

6: conv \leftarrow True

7: end if

8: end while

9: return diag(T), Q
```

# 4 Analysis

In order to test the quality of our implemented algorithms we will follow a two pronged approach. First, we will scrutinize the quality of the obtained results. Once we can be sure that the implemented routines are reliable or, to be more precise, under which conditions our routines are reliable we can analyze their convergence behavior and compare the efficiency accross algorithms in terms of computational cost.

## 4.1 Accuracy

If we wish to investigate the accuracy of our algorithms, we first need to define what we mean by that. We set up an environment of unit tests that compare the absolute difference between the vector of true eigenvalues of a matrix and the vector 4.1 Accuracy 4 ANALYSIS

Table 1: Failed unit tests accross matrix-sizes Q									
Algorithm	Maximum number								
	of iterations	3	4	5	6	7			
Jacobi	-	0	0	0	0	0			
QRM1	10	785	952	994	999	1000			
	100	110	224	349	474	565			
	1000	7	16	33	38	45			
	10000	1	1	1	1	3			
	100000	0	0	0	0	0			
QRM2	10	826	975	997	1000	1000			
	100	111	191	360	467	602			
	1000	9	16	22	35	58			
	10000	0	0	3	2	0			
	100000	0	0	0	0	0			
QRM3	10	251	616	896	983	997			
	100	29	52	116	222	311			
	1000	6	2	8	19	21			
	10000	0	0	1	1	3			
	100000	0	0	0	0	0			

of computed eigenvalues by our custom algorithms. If any element of this  $L_1$ -norm is larger than a threshold of  $10^{-5}$  we consider the test as failed. In order to obtain matrices with known eigenvalues we will reverse the spectral decomposition, which states that every square, symmetric matrix  $A \in \mathbb{R}^{m \times m}$  can be written as

$$A = \Gamma \Lambda \Gamma' = \sum_{j=1}^{m} \lambda_j \gamma_j \gamma_j', \tag{14}$$

where  $\Lambda = diag(\lambda_1, \ldots, \lambda_m)$  is a diagonal matrix with the eigenvalues of A on its main diagonal and where  $\Gamma = (\gamma_1, \ldots, \gamma_m)$  is the matrix containing the associated eigenvectors [Härdle and Simar, 2015]. This means we can draw a random vector of eigenvalues  $\lambda_{true}$  and a random orthogonal matrix to construct a test matrix  $A_{test}$ . Once the matrix  $A_{test}$  is initialized, we can plug it into the eigenvalue-routine we wish to test.

The number of failures out of 1000 unit tests on matrices of different dimensions are catalogued in Table 1. Most notably, we see that the Jacobi-method outperforms any of the QR-methods in terms of accuracy. This is mostly due to its convergence

4.2 Efficiency 5 CONCLUSION

criterion being defined differently than the convergence criterion of the QR-methods. Simply put, the Jacobi-method will always converge in this setup. Also keep in mind, that since the Jacobi-method is only allowed for symmetric matrices, it essentially eliminates two non-zero off-diagonal elements at once. Comparing only the QR-methods amongst each other one can clearly see, that the quality of the obtained results increases, the more willing an analyst is to wait. Convergence is slower for the QR-methods and a premature stop leads to poor eigenvalues. One the other hand, if the QR-methods converge, they unanimously pass all tests. It would therefore be possible to always achieve an arbitrary grade of precision by setting the maximum number of iterations to infinity.

## 4.2 Efficiency

Now that we can be confident that the routines offer accurate results

#### 5 Conclusion

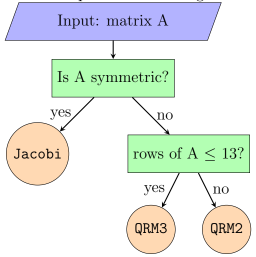


Figure 6: Decision process of final eigenvalue routine

# 6 Appendix

#### 6.1 Householder-Reflections

Our goal is still to diagonalize a matrix in order to programmatically extract it's eigenvalues. So far we have seen that there exist such transformations that conserve the eigenvalues of a given matrix. However, we require transformations that, additionally, eliminate non-zero entries on the off-diagonal elements of said matrix. A greedy technique, that eliminates all but the first elements of a vector is proposed in the form of Householder-Reflections.

Let u and v be orthonormal vectors and let x be a vector in the space spanned by u and v, such that

$$x = c_1 u + c_2 + v$$

for some scalars  $c_1$  and  $c_2$ . The vector

$$\tilde{x} = -c_1 u + c_2 v$$

is a reflection of x through the line difined by the vector u. Now consider the matrix

$$P = I - 2uu'. (15)$$

Note that

$$Px = c_1 u + c_2 v - 2c_1 u u u' - 2c_2 v u u'$$

$$= c_1 u + c_2 v - 2c_1 u' u u - 2c_2 u' v u$$

$$= -c_1 u + c_2 v$$

$$= \hat{x}.$$

The matrix P is called a reflector. The usefulness of Householder-Reflections stems from the fact that it is easy to transform a vector of the form

$$x = (x_1, x_2, \dots, x_n)$$

into a vector

$$\hat{x} = (\hat{x}_1, 0, \dots, 0).$$

If  $Qx = \hat{x}$ , then  $||x||_2 = ||\hat{x}||_2$  and thus  $\hat{x}_1 = \pm ||x||_2$ , since it is the only non-zero entry. To construct the reflector let

$$v = (x_1 + sign(x_1)||x||_2, x_2, \dots, x_n)$$
(16)

and  $u = \frac{v}{||v||_2}$  [Gentle, 1998]. We use the sign-function, which simply returns the sign of its argument in order to avoid the numerical problem known as catastrophic cancellation. It can occur when adding two very close, but different, floating point numbers of differing signs. In some unfortunate cases both of these numbers get represented by the same computer number and, because of their opposing signs cancel each other out. In our case this would mean, that we reflect the vector onto the origin. Fortunately, by making use of the sign function we can make sure that both summands will share the same sign, thus mitigating any concerns about catastrophic cancellation.

We use reflectors to compute the so called QR factorization of an aribitrary square matrix  $A \in \mathbb{R}^{n \times n}$ .

$$A = QR \tag{17}$$

where Q is orthogonal and R is upper triangular. We use Householder transformations to reflect the  $i^{th}$  column and produce zeros below the (i, i) element. The QR-factorization of a matrix  $A \in \mathbb{R}^5$  would therefore consist of five Householder-reflections with  $Q = P_5 P_4 P_3 P_2 P_1$ . The number of computations for the QR factorization in this fashion is  $2n^3/3$  multiplications and  $2n^3/3$  additions [Gentle, 1998].

#### 6.2 Givens-Rotations

Another way of forming the QR-factorization is by using orthogonal transformations which rotate a vector in a way such that a specific element becomes 0 and only one other element in the vector being changed. These transformations are called Givens transformations, Givens Rotations or Jacobi transformations

Figure 7: Rotation of  $x \supseteq x_2$  (4,3) (5,0)A Givens rotation in  $\mathbb{R}^2$ .

Using orthogonal transformations we can also rotate a vector in such a way that a specified element becomes 0 and only one other element in the vector is changed. The basic idea can be seen in a two-dimensional space. We wish to rotate the vector  $x = (x_1, x_2)$  to  $\tilde{x} = (\tilde{x_1}, 0)$  as with a reflector.

It is easy to see that the orthogonal matrix

$$Q = \begin{bmatrix} \cos \theta & \sin \theta \\ -\sin \theta & \cos \theta \end{bmatrix}$$

performs the desired rotation, if  $\cos \theta = \frac{x_1}{||x||_2}$  and  $\sin \theta = \frac{x_2}{||x||_2}$ 

In general, we can construct an orthogonal  $matrixV_{pq}$ , that will transform the vector

$$x = (x_1, \dots, x_p, \dots x_q, \dots, x_n)$$

to

$$\tilde{x} = (x_1, \dots, \tilde{x}_p, \dots 0, \dots, x_n)$$

. The matrix that does this is

$$V_{pq}(\theta) = \begin{bmatrix} 1 & & & & & \\ & \ddots & & & \\ & & \cos \theta & & \sin \theta & \\ & & & \ddots & \\ & & -\sin \theta & & \cos \theta & \\ & & & \ddots & \\ & & & & 1 \end{bmatrix}$$
 (18)

where  $\cos \theta = \frac{x_p}{||x||}$  and  $\sin \theta = \frac{x_q}{||x||}$ .

A rotation matrix is therefore the same as an identity matrix, in which we change four elements [Börm and Mehl, 2012]. We will use Givens rotations primarily in the Jacobi-Method.

#### 6.3 Eigenvalue Routines

```
import numpy as np
import copy

def hreflect1D(x):
    """
    Calculate Householder reflection: Q = I - 2*uu'.

Parameters:
    X: numpy array.

Returns:
    Qx: reflected vector.
    Q: Reflector (matrix).
```

```
n n n
15
       # Construct v:
16
      v = copy.deepcopy(x)
17
      v[0] += np.linalg.norm(x)
18
19
       # Construct u: normalize v.
20
      vnorm = np.linalg.norm(v)
21
       if vnorm:
22
           u = v / np.linalg.norm(v)
23
       else:
24
           u = v
26
       # Construct Q:
27
       Q = np.eye(len(x)) - 2 * np.outer(u, u)
28
       Qx = np.dot(Q, x)
29
30
       return Qx, Q
31
32
33
  def qr_factorize(X, offset=0):
34
       n n n
35
       Compute QR factorization of X s.t. QR = X.
36
37
      Parameters:
38
           - X: square numpy ndarray.
39
           - offset: (int) either 0 or 1. If offset is unity: compute
40
              Hessenberg -
                      matrix.
41
42
       Returns:
43
           Q: square numpy ndarray, same shape as X. Rotation matrix.
44
           R: square numpy ndarray, same shape as X. Upper triangular
45
              matrix if
              offset is 0, Hessenberg-matrix if offset is 1.
46
       n n n
47
```

```
assert offset in [0, 1]
48
      assert type(X) == np.ndarray
49
      assert X.shape[0] == X.shape[1]
50
51
      R = copy.deepcopy(X)
52
      Q = np.eye(X.shape[0])
53
      for i in range(X.shape[0]-offset):
           Pi = np.eye(R.shape[0])
           _, Qi = hreflect1D(R[i+offset:, i])
57
           Pi[i+offset:, i+offset:] = Qi
58
59
           Q = Pi.dot(Q)
60
           R = Pi.dot(R)
61
62
      return Q.T, R
63
```

```
11 11 11
  Algorithms for solving eigenvalue problems.
3
  1. Compute diagonalization of 2x2 matrices via jacobi iteration.
  2. Generalize Jacobi iteration for symmetric matrices.
  n n n
  import numpy as np
8 import copy
  import warnings
10 from scipy import linalg as lin
  from algorithms import helpers
12
13
  def jacobi2x2(A):
       n n n
      Diagonalize a 2x2 matrix through jacobi step.
16
17
      Solve: U' A U = E s.t. E is a diagonal matrix.
18
19
```

```
Parameters:
20
           A - 2x2 numpy array.
21
      Returns:
22
           A - 2x2 diagonal numpy array
23
       11 11 11
24
      assert type(A) == np.ndarray
25
      assert A.shape == (2, 2)
      assert A[1, 0] == A[0, 1]
27
28
      alpha = 0.5 * np.arctan(2*A[0, 1]/(A[1, 1] - A[0, 0]))
29
      U = np.array([[np.cos(alpha), np.sin(alpha)],
30
                      [-np.sin(alpha), np.cos(alpha)]])
31
      E = np.matmul(U.T, np.matmul(A, U))
32
      return E
33
34
35
  def jacobi(X, precision=1e-6, debug=False):
36
37
      Compute Eigenvalues and Eigenvectors for symmetric matrices.
38
39
      Parameters:
40
           X - 2D numpy ndarray which represents a symmetric matrix
41
           precision - float in (0, 1). Convergence criterion.
42
43
      Returns:
44
           A - 1D numpy array with eigenvalues sorted by absolute
45
            value
           U - 2D numpy array with associated eigenvectors (column).
       n n n
47
      assert 0 < precision < 1.
48
      assert type(X) == np.ndarray
49
      n, m = X.shape
50
      assert n == m
51
      assert all(np.isclose(X - X.T, np.zeros(n)).flatten())
52
      A = copy.deepcopy(X)
53
```

```
U = np.eye(A.shape[0])
54
      L = np.array([1])
55
      iterations = 0
56
57
      while L.max() > precision:
58
           L = np.abs(np.tril(A, k=0) - np.diag(A.diagonal()))
59
           i, j = np.unravel_index(L.argmax(), L.shape)
           alpha = 0.5 * np.arctan(2*A[i, j] / (A[i, i]-A[j, j]))
61
62
           V = np.eye(A.shape[0])
63
           V[i, i], V[j, j] = np.cos(alpha), np.cos(alpha)
64
           V[i, j], V[j, i] = -np.sin(alpha), np.sin(alpha)
65
66
           A = np.dot(V.T, A.dot(V))
67
           U = U.dot(V)
68
           iterations += 1
69
70
      # Sort by eigenvalue (descending order) and flatten A
71
      A = np.diag(A)
72
      order = np.abs(A).argsort()[::-1]
73
      if debug:
74
           return iterations
75
76
      return A[order], U[:, order]
77
78
79
  def qrm(X, maxiter=15000, debug=False):
80
81
      Compute Eigenvalues and Eigenvectors using the QR-Method.
82
83
      Parameters:
84
           - X: square numpy ndarray.
85
      Returns:
86
           - Eigenvalues of A.
87
           - Eigenvectors of A.
88
```

```
n n n
89
       n, m = X.shape
90
       assert n == m
91
92
       # First stage: transform to upper Hessenberg-matrix.
93
       A = copy.deepcopy(X)
94
       conv = False
       k = 0
97
       # Second stage: perform QR-transformations.
98
       while (not conv) and (k < maxiter):
           k += 1
100
           Q, R = helpers.qr_factorize(A)
101
           A = R.dot(Q)
102
103
           conv = np.alltrue(np.isclose(np.tril(A, k=-1), np.zeros((n))
104
             , n))))
105
       if not conv:
106
           warnings.warn("Convergence was not reached. Consider
107
             raising maxiter.")
       if debug:
108
           return k
109
       Evals = A.diagonal()
110
       order = np.abs(Evals).argsort()[::-1]
111
       return Evals[order], Q[order, :]
112
113
114
  def qrm2(X, maxiter=15000, debug=False):
       n n n
116
       First compute similar matrix in Hessenberg form, then compute
117
         the
       Eigenvalues and Eigenvectors using the QR-Method.
118
119
       Parameters:
120
```

```
- X: square numpy ndarray.
121
       Returns:
122
           - Eigenvalues of A.
123
           - Eigenvectors of A.
124
       n n n
125
       n, m = X.shape
126
       assert n == m
127
128
       # First stage: transform to upper Hessenberg-matrix.
129
       A = lin.hessenberg(X)
130
       conv = False
131
       k = 0
132
133
       \# Second stage: perform QR-transformations.
134
       while (not conv) and (k < maxiter):
135
           k += 1
136
           Q, R = helpers.qr_factorize(A)
137
           A = R.dot(Q)
138
139
           conv = np.alltrue(np.isclose(np.tril(A, k=-1), np.zeros((n
140
             , n))))
141
       if not conv:
142
           warnings.warn("Convergence was not reached. Consider
143
             raising maxiter.")
       if debug:
144
           return k
145
       Evals = A.diagonal()
       order = np.abs(Evals).argsort()[::-1]
       return Evals[order], Q[order, :]
148
149
150
  def qrm3(X, maxiter=15000, debug=False):
151
152
       First compute similar matrix in Hessenberg form, then compute
153
```

```
the
       Eigenvalues and Eigenvectors using the QR-Method.
154
155
       Parameters:
156
            - X: square numpy ndarray.
157
       Returns:
158
            - Eigenvalues of A.
159
           - Eigenvectors of A.
160
       H/H/H
161
       n, m = X.shape
162
       assert n == m
163
164
       \hbox{\it\# First stage: transform to upper Hessenberg-matrix.}
165
       T = lin.hessenberg(X)
166
167
       conv = False
168
       k = 0
169
170
       \# Second stage: perform QR-transformations.
171
       while (not conv) and (k < maxiter):
172
           k += 1
173
           Q, R = helpers.qr_factorize(T - T[n-1, n-1] * np.eye(n))
174
           T = R.dot(Q) + T[n-1, n-1] * np.eye(n)
175
176
            conv = np.alltrue(np.isclose(np.tril(T, k=-1), np.zeros((n
177
              , n))))
178
       if not conv:
179
            warnings.warn("Convergence was not reached. Consider
180
             raising maxiter.")
       if debug:
181
            return k
182
       Evals = T.diagonal()
183
       order = np.abs(Evals).argsort()[::-1]
184
       return Evals[order], Q[order, :]
185
```

#### 6.4 Analysis: Figures

```
import os
2 import copy
3 import pandas as pd
  import numpy as np
5 import seaborn as sns
6 from scipy import linalg as lin
7 from scipy.stats import ortho_group
  from matplotlib import pyplot as plt
  datadir = os.path.join("analysis", "benchmarks.csv")
  outpath = os.path.join("media", "plots")
  trials = pd.read_csv(datadir, index_col=0)
  trials.groupby(["algorithm", "dimension"]).iterations.describe()
16 # Boxplot iteration:
fig = plt.figure(figsize=(10, 5))
18 sns.boxplot(x="dimension", y="iterations", hue="algorithm", data=
   trials)
plt.yscale("log")
  plt.title("Iterations needed before Convergence")
21 plt.savefig(os.path.join(outpath, "iterations_boxplot.png"))
 plt.show()
 plt.close()
24
25 # Boxplot elapsed time:
fig = plt.figure(figsize=(10, 5))
27 sns.boxplot(x="dimension", y="time", hue="algorithm", data=trials)
plt.title("Time needed before Convergence")
plt.ylabel("time (sec)")
plt.yscale('log')
plt.savefig(os.path.join(outpath, "time_boxplot.png"))
plt.show()
plt.close()
```

```
34
  # Visualize Algorithm-Progress:
35
np.random.seed(42)
37 size = 5
  Lambda = np.diag(np.random.randint(low=0, high=10, size=size))
G = ortho_group.rvs(dim=size)
  X = np.dot(G, Lambda.dot(G.T))
41
42
  def plot_factory(func):
43
      def plotter(savepath, **fig_kw):
           def algorithm_generator(*args, **kwargs):
45
               return func(*args, **kwargs)
46
47
           fig, ax = plt.subplots(nrows=2, ncols=2, **fig_kw)
48
           algorithm_iterator = algorithm_generator()
49
           j = -1
50
51
           for i, A in enumerate(algorithm_iterator):
52
               if i in (0, 1, 10, 75):
53
                   j += 1
54
55
                   hm = ax[j // 2, j \% 2].imshow(A,
56
                                                    cmap=plt.get_cmap('
57
                                                      seismic'),
                                                    vmin=-X.max(),
58
                                                    vmax=X.max())
59
                   ax[j // 2, j % 2].set_yticks([])
                   ax[j // 2, j % 2].set_xticks([])
61
                   ax[j // 2, j % 2].set_title("Iteration: " + str(i)
62
                     )
63
                   if i > 75:
64
                        break
65
66
```

```
fig.subplots_adjust(right=0.8)
67
           cbar_ax = fig.add_axes([0.85, 0.15, 0.05, 0.7])
68
           fig.colorbar(hm, cax=cbar_ax)
69
70
           sup_title = "Demonstrate {} on a {}x{} matrix".format(
71
                func.__name__,
72
                *X.shape)
73
74
           fig.suptitle(sup_title)
75
           fig.savefig(savepath)
76
77
           return fig, ax
78
79
       return plotter
80
81
82
  @plot_factory
83
  def jacobi():
84
       11 11 11
85
       Compute Eigenvalues and Eigenvectors for symmetric matrices
86
        using the
       jacobi method.
87
88
       Yields:
89
            st A - 2D numpy array of current iteration step.
90
       11 11 11
91
       A = copy.deepcopy(X)
92
       U = np.eye(A.shape[0])
93
       L = np.array([1])
94
       iterations = 0
95
96
       while iterations < 5000:
97
           L = np.abs(np.tril(A, k=0) - np.diag(A.diagonal()))
98
           i, j = np.unravel_index(L.argmax(), L.shape)
99
           alpha = 0.5 * np.arctan(2*A[i, j] / (A[i, i]-A[j, j]))
100
```

```
101
            V = np.eye(A.shape[0])
102
            V[i, i], V[j, j] = np.cos(alpha), np.cos(alpha)
103
            V[i, j], V[j, i] = -np.sin(alpha), np.sin(alpha)
104
105
            A = np.dot(V.T, A.dot(V))
106
            U = U.dot(V)
107
            iterations += 1
108
            yield A
109
   @plot_factory
112
   def qrm1():
113
        n n n
114
       Create generator for transformed matrices after applying the
115
         QR-Method.
116
        Yields:
117
            - T: 2D-numpy array. Similar matrix to X.
118
        11 11 11
119
       # First stage: transform to upper Hessenberg-matrix.
120
       T = copy.deepcopy(X)
121
122
       k = 0
123
       \# Second stage: perform QR-transformations.
124
       while k < 5000:
125
            k += 1
126
            Q, R = np.linalg.qr(T)
127
            T = R.dot(Q)
128
            yield T
129
130
   @plot_factory
132
133
   def qrm2():
        n n n
134
```

```
Create generator for transformed matrices after applying the
135
         QR-Method.
136
       Yields:
137
           - T: 2D-numpy array. Similar matrix to X.
138
       n n n
139
       # First stage: transform to upper Hessenberg-matrix.
140
       T = lin.hessenberg(X)
141
142
       k = 0
143
       # Second stage: perform QR-transformations.
       while k < 5000:
145
           if k == 0:
146
                yield X
147
           k += 1
148
           Q, R = np.linalg.qr(T)
149
           T = R.dot(Q)
150
           yield T
151
152
153
  @plot_factory
154
  def qrm3():
155
       n n n
156
       First compute similar matrix in Hessenberg form, then compute
157
         the
       Eigenvalues and Eigenvectors using the accelerated QR-Method.
158
159
       Yields:
160
            * T - 2D numpy array of current iteration step.
161
       n n n
162
       # First stage: transform to upper Hessenberg-matrix.
163
       T = lin.hessenberg(X)
164
       k = 0
165
       n, _= X.shape
166
167
```

```
\# Second stage: perform QR-transformations.
168
       while k < 5000:
169
           if k == 0:
170
                yield X
171
           k += 1
172
           Q, R = np.linalg.qr(T - T[n-1, n-1] * np.eye(n))
173
           T = R.dot(Q) + T[n-1, n-1] * np.eye(n)
174
175
           yield T
176
177
  jacobi(os.path.join(outpath, "jacobi.png"))
179
  qrm1(os.path.join(outpath, "qrm1.png"))
180
   qrm2(os.path.join(outpath, "qrm2.png"))
181
  qrm3(os.path.join(outpath, "qrm3.png"))
182
183
  plt.show()
184
  plt.close()
185
```

#### 6.5 Analysis: Unit tests

```
11 11 11
  Automated tests for different algorithms.
  n n n
  import os
  import sys
  import numpy as np
7 from threading import Thread
  import pandas as pd
9 from algorithms import eigen
  from scipy.stats import ortho_group
  from tqdm import tqdm
  from functools import wraps
13
  data_out = os.path.join("data", "accuracy_tests.csv")
15
17
  class AlgoTest(object):
18
      tests = {"algorithm": [],
19
                "dimension": [],
20
                "maxiter": [],
21
                "failed": []}
22
23
      def __init__(self, algo, dim, filepath, n_tests=1000, jobs=1,
24
                    *args, **kwargs):
25
           assert n_tests % jobs == 0
26
           self.algorithm = self.__get_test_algorithm(algo, *args, **
27
            kwargs)
           self.dim = dim
28
           self.n = n_tests // jobs
29
           self.failed = []
30
           self.jobs = jobs
31
           self.result = None
32
           self.path = filepath
33
```

```
self.maxiter = kwargs.get("maxiter", None)
34
35
          if not os.path.exists(self.path):
36
               self.save(header=["algorithm", "dimension", "maxiter",
37
                  "failed"])
38
      def __get_test_algorithm(self, algorithm, *args, **kwargs):
39
           @wraps(algorithm)
           def algo(X):
41
               return algorithm(X, *args, **kwargs)
42
           return algo
44
      def __get_test_matrix(self):
45
           """Return matrix with assosiated Eigenvalues."""
46
           eigenvalues = np.random.uniform(size=self.dim)
47
           eigenvectors = ortho_group.rvs(dim=self.dim)
48
          Lambda = np.diag(eigenvalues)
49
50
          matrix = np.dot(eigenvectors, Lambda).dot(eigenvectors.T)
51
52
           order = np.abs(eigenvalues).argsort()[::-1]
53
           return matrix, eigenvalues[order]
54
55
      def __run_test(self):
56
           """Run singular test."""
57
          mat, true_eig = self.__get_test_matrix()
58
           test_eig, _ = self.algorithm(mat)
59
           test_res = np.alltrue(np.isclose(true_eig, test_eig))
           self.failed.append(not test_res)
61
62
      def __run_tests(self):
63
           """Run multiple tests."""
64
           for _ in range(self.n):
65
               try:
66
                   self.__run_test()
67
```

```
except (KeyboardInterrupt, SystemExit):
68
                    self.__save()
69
                    sys.exit(0)
70
71
       def run(self):
72
           """Distribute tests accross threads."""
73
           threadlist = [None] * self.jobs
75
           for i in range(self.jobs):
77
               threadlist[i] = Thread(target=self.__run_tests, daemon
78
                 =True)
               threadlist[i].start()
79
80
           for thread in threadlist:
81
               thread.join()
82
83
           self.result = sum(self.failed)
84
           self.tests["algorithm"].append(self.algorithm.__name__)
85
           self.tests["dimension"].append(self.dim)
86
           self.tests["failed"].append(self.result)
87
           if self.algorithm.__name__ == "jacobi":
88
               self.tests["maxiter"].append(None)
89
           else:
90
               self.tests["maxiter"].append(self.maxiter)
91
92
       def save(self, header=False):
93
           if os.path.exists(self.path):
                    print("Saving results.")
           df = pd.DataFrame(self.tests)
96
           with open(self.path, 'a') as f:
97
               df.to_csv(f, header=header, index=False,
98
                          columns = ["algorithm", "dimension", "maxiter"
99
                             , "failed"])
100
```

```
101
  # Unit tests
102
  if __name__ == "__main__":
103
       # Define Flags.
104
       JOBS = 20
105
       MAXITER = (10, 100, 1000, 10000, 100000)
106
       DIMS = range(3, 8)
107
       ALGOS = {'jacobi': eigen.jacobi,
108
                 'qrm': eigen.qrm,
109
                 'qrm2': eigen.qrm2,
110
                 'qrm3': eigen.qrm3}
112
       # Define parameters of all runs.
113
       parameters = []
114
       for algo in ALGOS.values():
115
            for maxiter in MAXITER:
116
                for dim in range(3, 8):
117
                     param = (algo,
118
                               maxiter if algo.__name__ != "jacobi" else
119
                                  None,
                               dim)
120
                    parameters.append(param)
121
122
       # Check progress of previous runs.
123
       if os.path.exists(data_out):
124
           print("Check progress:")
125
           required = [(algo.__name__, m, dim) for (algo, m, dim) in
126
             parameters]
           required = set(required)
127
128
           progress = pd.read_csv(data_out)
129
            done = []
131
            for (a, d, m, _) in progress.values:
132
                if np.isnan(m):
133
```

```
param = (a, None, d)
134
                else:
135
                     param = (a, int(m), d)
136
                done.append(param)
137
            done = set(done)
138
139
            to_do = required.difference(done)
140
            parameters = [(ALGOS[a], m, d) for a, m, d in to_do]
141
142
       for algo, maxiter, dim in tqdm(parameters):
143
            if algo is eigen.jacobi:
                algo_test = AlgoTest(filepath=data_out,
145
                                        algo=algo,
146
                                        dim=dim,
147
                                        jobs=JOBS)
148
            else:
149
                algo_test = AlgoTest(filepath=data_out,
150
                                        algo=algo,
151
                                        dim=dim,
152
                                        maxiter=maxiter,
153
                                        jobs=JOBS)
154
            algo_test.run()
155
156
       algo_test.save()
157
158
       results = pd.read_csv(data_out)
159
       results.head(10)
160
       results.loc[results.algorithm == 'jacobi', 'maxiter'] = '-'
161
162
       crosstab = pd.crosstab(index=[results.algorithm, results.
163
         maxiter],
                                 columns=results.dimension,
164
                                 values=results.failed,
165
                                 aggfunc=np.mean,
166
                                 dropna=True)
167
```

print(crosstab.to\_latex())

# 7 References

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