L_Bijman_S_Gijsbers_HW4

November 27, 2023

1 Homework set 4

Before you turn this problem in, make sure everything runs as expected (in the menubar, select Kernel \rightarrow Restart Kernel and Run All Cells...).

Please submit this Jupyter notebook through Canvas no later than Mon Nov. 27, 9:00. Submit the notebook file with your answers (as .ipynb file) and a pdf printout. The pdf version can be used by the teachers to provide feedback. A pdf version can be made using the save and export option in the Jupyter Lab file menu.

Homework is in **groups of two**, and you are expected to hand in original work. Work that is copied from another group will not be accepted.

2 Exercise 0

Write down the names + student ID of the people in your group.

Loes Bijman 15211312

Sacha Gijsbers 12798525

3 About imports

Please import the needed packages by yourself.

4 Sparse matrices

A sparse matrix or sparse array is a matrix in which most of the elements are zero. There is no strict definition how many elements need to be zero for a matrix to be considered sparse. In many examples, the number of nonzeros per row or column is a small fraction, a few percent or less, of the total number of elements of the row or column. By contrast, if most of the elements are nonzero, then the matrix is considered dense.

In the context of software for scientific computing, a sparse matrix typically refers to a storage format, in which elements which are known to be zero are not stored. In Python, the library scipy.sparse defines several sparse matrix classes, such as scipy.sparse.csr_array. To construct such an object, one passes for each nonzero element the value, and the row and column coordinates. In some cases, one can also just pass the nonzero (off-)diagonals, see scipy.sparse.diags.

Functions for dense matrices do not always work with sparse matrices. For example for the product of a sparse matrix with a (dense) vector, there is the member function scipy.sparse.csr_array.dot, and for solving linear equations involving a sparse matrix, there is the function scipy.sparse.linalg.spsolve.

```
[]: # Import some basic packages
     import numpy as np
     import matplotlib.pyplot as plt
     import math
[]: from scipy.sparse import csr_array
     # This is how to create a sparse matrix from a given list of (row, column,
     ⇔value) tuples.
     row = [0, 3,
                       1,
                            0]
     col = [0, 3,
                     1.
     data = [4.0, 5.0, 7.0, 9.0]
     M = csr_array((data, (row, col)), shape=(4, 4))
     print("When printing a sparse matrix, it shows its nonzero entries:")
     print(M)
     print("If you want to see its `dense` matrix form, you have to use `mat.
     print(M.toarray())
     # This is how to perform matrix-vector products.
     x = np.array([1, 2, 3, 4])
     print("For x=\{\}, Mx = \{\}".format(x, M.dot(x)))
    When printing a sparse matrix, it shows its nonzero entries:
                    4.0
      (0, 0)
      (0, 2)
                    9.0
      (1, 1)
                    7.0
      (3, 3)
                    5.0
    If you want to see its `dense` matrix form, you have to use `mat.toarray()`:
    [[4. 0. 9. 0.]
     [0. 7. 0. 0.]
     [0. 0. 0. 0.]
     [0. 0. 0. 5.]]
    For x=[1 \ 2 \ 3 \ 4], Mx = [31. \ 14. \ 0. \ 20.]
[]: from scipy.sparse import diags, SparseEfficiencyWarning
     from scipy.sparse.linalg import spsolve
```

warnings.simplefilter('ignore', SparseEfficiencyWarning) # Suppress confusing

import warnings

 \hookrightarrow warning

```
# This is how to create a sparse matrix from a given list of subdiagonals.
diagonals = [[1, 2, 3, 4], [1, 2, 3], [1, 2]]
M = diags(diagonals, [0, 1, 2])
print("This matrix has values on its diagonal and on offdiagonals 1 and 2 rows⊔
 →ABOVE it.")
print(M.toarray())
M = diags(diagonals, [0, -1, -2])
print("This matrix has values on its diagonal and on offdiagonals 1 and 2 rows⊔
 ⇔BELOW it.")
print(M.toarray())
print("If you want to visualize the matrix for yourself, use `plt.imshow`:")
plt.imshow(M.toarray())
plt.colorbar()
plt.show()
# This is how to solve sparse systems.
b = np.array([1, 2, 3, 4])
x = spsolve(M, b)
print("For b={}, the solution x to Mx=b is {}".format(b, x))
print("And indeed, Mx - b = {}".format(M.dot(x) - b))
This matrix has values on its diagonal and on offdiagonals 1 and 2 rows ABOVE
it.
[[1. 1. 1. 0.]
[0. 2. 2. 2.]
[0. 0. 3. 3.]
 [0. 0. 0. 4.]]
```

This matrix has values on its diagonal and on offdiagonals 1 and 2 rows BELOW it.

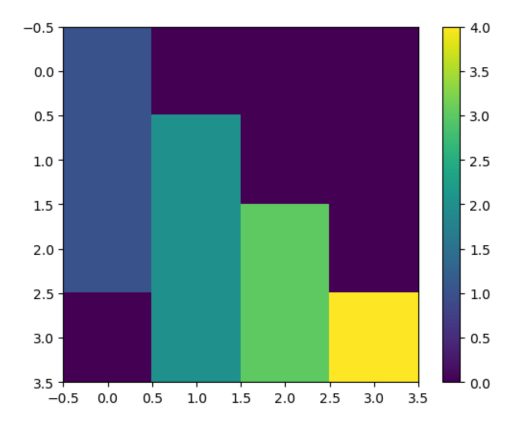
[[1. 0. 0. 0.]

[1. 2. 0. 0.]

[1. 2. 3. 0.]

[0. 2. 3. 4.]]

If you want to visualize the matrix for yourself, use `plt.imshow`:



For
$$b=[1\ 2\ 3\ 4]$$
, the solution x to Mx=b is [1. 0.5 0.33333333 0.5] And indeed, Mx - b = [0. 0. 0. 0.]

5 Exercise 1

Consider the following boundary value problem involving a nonlinear ordinary differential equation:

$$y''(x) + \exp(y(x)) = 0, \quad 0 < x < 1, \quad y(0) = y(1) = 0.$$
 (1)

The purpose of this exercise is to approximate the solution to this boundary value problem, by discretizing the problem and then solving the resulting system of nonlinear equations.

Problem (1) will be discretized using finite differences. Suppose we use n+2 discretization points for x, denoted $x_k = kh$ for $k \in \{0, \dots, n+1\}$ and h = 1/(n+1). The approximate solution is denoted $y_k = y(x_k)$.

We will use a second-order central finite difference approximation for the second derivative:

$$y''(x_k) \approx \frac{y_{k-1} - 2y_k + y_{k+1}}{h^2}. (2)$$

The term $\exp(y(x_k))$ can simply be approximated by $\exp(y_k)$. Thus for $x=x_k$, equation (1) becomes

$$\frac{y_{k-1} - 2y_k + y_{k+1}}{h^2} + \exp y_k = 0, \quad k = 1, \dots, n.$$
 (3)

The boundary conditions (the conditions y(0) = y(1) = 1), lead to the requirement that $y_0 = y_{n+1} = 0$. To find the remaining values y_k , k = 1, ..., n, equation (3) will be used for k = 1, ..., n. In this way, one obtains n equations for n unknowns, to which, in principle, a rootfinding method can be applied.

We will write $\vec{y} = \begin{bmatrix} y_1 \\ \vdots \\ y_n \end{bmatrix}$ for the vector of values to be determined.

5.1 (a) (2 pts)

As a first step, finish the function SecondDerMatrix that returns a matrix \mathbf{M} that maps the vector \vec{y} to the vector of the approximate values $y''(x_k)$, $k=1,\ldots,n$ given in (2). To get full points for this part of the exercise you must create output in the form of a sparse matrix.

```
[]: from scipy.sparse import csr_array
     def SecondDerMatrix(n):
         111
         Creates matrix M that maps any vector y to the vector of approximate values \Box
      ⇔for the second derivative,
         using the finite differences approximation. This matrix assumes that the \Box
      ⇒boundary conditions are
         y(0) = y(1) = 0.
         Input: n(int), size of the matrix
         Output: sparse matrix M
         111
         h = 1/(n+1)
         diagonals = [(n)*[-2], (n-1)*[1], (n-1)*[1]]
         M = diags(diagonals, [0, 1, -1])
         M /= h**2
         return M
     print(SecondDerMatrix(8).toarray())
```

```
ΓΓ-162.
           81.
                   0.
                         0.
                                0.
                                       0.
                                              0.
                                                     0.1
    81. -162.
                  81.
                         0.
                                0.
                                       0.
                                              0.
                                                     0.]
     0.
           81. -162.
                        81.
                                0.
                                       0.
                                              0.
                                                     0.]
 Γ
     0.
            0.
                  81. -162.
                               81.
                                       0.
                                              0.
                                                     0.]
                        81. -162.
0.
            0.
                   0.
                                      81.
                                                     0.]
0.
            0.
                   0.
                         0.
                               81. -162.
                                             81.
                                                     0.]
81. -162.
     0.
            0.
                         0.
                                0.
                                                    81.]
                   0.
            0.
                                0.
                                       0.
                                             81. -162.]]
     0.
                   0.
                         0.
```

5.2 (b) (1 pt)

Second-order central finite differences are exact for quadratic functions. In order to test your implementation, choose n = 10 and apply the second derivative matrix from part (a) to a quadratic function y(x) with y(0) = y(1) = 0 for which you know the second derivative y''(x).

```
[]: def QuadraticFunction(x):
         return 5*x**2 - 5*x
     n = 10
     h = 1/(n+1)
     y = []
     for k in range(1,n+1):
         x = k*h
         y.append(QuadraticFunction(x))
     M = SecondDerMatrix(n)
     print(M.toarray())
     print(y)
     print(M.dot(y))
    [[-242.
              121.
                      0.
                             0.
                                   0.
                                          0.
                                                0.
                                                       0.
                                                             0.
                                                                    0.]
```

```
[ 121. -242.
                121.
                         0.
                                0.
                                       0.
                                              0.
                                                    0.
                                                                  0.]
                                                           0.
          121. -242.
                       121.
                                0.
                                       0.
                                              0.
                                                    0.
                                                           0.
                                                                  0.]
                121. -242.
 121.
                                       0.
                                                                  0.]
 Γ
                       121. -242.
                                                                  0.]
     0.
            0.
                   0.
                                     121.
                                              0.
                                                    0.
                                                           0.
 121. -242.
                                                                  0.]
     0.
            0.
                         0.
                                            121.
                                                    0.
                                                           0.
 Γ
                                     121. -242.
                                                                  0.]
     0.
            0.
                                                  121.
 0.
            0.
                         0.
                                            121. -242.
                                                         121.
                                                                  0.]
                   0.
                                0.
                                       0.
 Γ
     0.
            0.
                   0.
                         0.
                                0.
                                       0.
                                              0.
                                                  121. -242.
                                                                121.7
 Γ
     0.
            0.
                   0.
                         0.
                                0.
                                       0.
                                              0.
                                                    0.
                                                         121. -242.]]
[-0.41322314049586784, -0.7438016528925621, -0.9917355371900827,
-1.15702479338843, -1.2396694214876034, -1.2396694214876034,
-1.1570247933884295, -0.9917355371900829, -0.7438016528925613,
-0.4132231404958677]
[10. 10. 10. 10. 10. 10. 10. 10. 10. 10.]
```

5.3 (c) (2 pts)

Defining
$$\vec{y} = \begin{bmatrix} y_1 \\ \vdots \\ y_n \end{bmatrix}$$
 and $E(\vec{y}) = \begin{bmatrix} \exp(y_1) \\ \vdots \\ \exp(y_n) \end{bmatrix}$, the equations (3) can be written in the form

$$F(\vec{y}) := \mathbf{M} \cdot \vec{y} + E(\vec{y}) = \vec{0}.$$

Finish the function F that defines $F(\vec{y}) = \mathbf{M} \cdot \vec{y} + E(\vec{y})$. Finish the function JacobianF that computes the Jacobian $\mathbf{J}_F(\vec{y})$ of $F(\vec{y})$. To get full points for this part of the exercise, the Jacobian must be computed in the form of a sparse matrix.

```
[]: def F(y):
         M = SecondDerMatrix(len(y))
         Ey = np.exp(np.array(y))
         return M.dot(y) + Ey
     def JacobianF(y):
         n = len(y)
         h = 1/(n+1)
         diag_list = []
         for yi in y:
             diag_list.append((1/h**2)*(-2) + np.exp(yi))
         diagonals = [diag_list, (n-1)*[1/h**2], (n-1)*[1/h**2]]
         J = diags(diagonals, [0, 1, -1])
         return J
     y = [1,2,3,2,1]
     print(F(y))
     print(JacobianF(y).toarray())
```

```
7.3890561 -51.91446308
                                            7.3890561
[ 2.71828183
                                                          2.71828183]
[[-69.28171817
                36.
                               0.
                                             0.
                                                           0.
                                                                     ]
[ 36.
               -64.6109439
                                                                     ]
                              36.
                                             0.
                                                           0.
                                                           0.
                                                                     ]
[ 0.
                36.
                             -51.91446308
                                           36.
[ 0.
                 0.
                              36.
                                           -64.6109439
                                                          36.
                                                                     1
-69.28171817]]
   0.
                 0.
                               0.
                                            36.
```

5.4 (d) (3 pts)

- 1. Write down the first order Taylor expansion $T_F(\vec{y}, \vec{s})$ for $F(\vec{y} + \vec{s})$.
- 2. In order to check your implementation of the Jacobian matrix, compute and print both $F(\vec{y} + \vec{s})$ and its first order Taylor approximation $T_F(\vec{y}, \vec{s})$ for a choice \vec{y} and \vec{s} .
- 3. Verify numerically that the error $||F(\vec{y}+\vec{s})-T_F(\vec{y},\vec{s})||_2$ is $\mathcal{O}(||\vec{s}||_2^2)$. Hint: take vectors \vec{s} with $||\vec{s}||_2 = \mathcal{O}(h)$ for multiple values for h, e.g. $h = 10^{-k}$ for a range of k.

Subquestion 1.

The first order Taylor expansion for $F(\vec{y} + \vec{s})$ is given by: $F(\vec{y} + \vec{s}) \approx F(\vec{y}) + J_F(\vec{y}) \cdot \vec{s}$. Therefore, we have

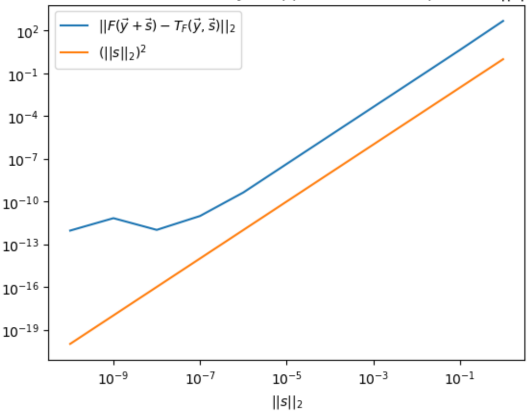
$$\begin{split} T_F(\vec{y}, \vec{s}) &= F(\vec{y}) + J_F(\vec{y}) \cdot \vec{s} \\ &= \mathbf{M} \cdot \vec{y} + E(\vec{y}) + J_F(\vec{y}) \cdot \vec{s} \end{split}$$

```
[]: # Subquestions 2 and 3.

# 2
def T_F(y,s):
```

```
return F(y) + JacobianF(y).dot(s)
# Veryify with a choice of y and s
y = np.array([1,2,3,2,1])
s = np.array([0.01, 0.01, 0.01, 0.01, 0.01])
print(f"T_F(y,s) = \{T_F(y,s)\}")
print(f"F(y+s) = \{F(y+s)\}")
# 3
n = 10 \# length of s and y
y = np.array(range(0,n))
s = np.ones(n)
s_norm = np.linalg.norm(s)
error_list = []
norm_s_list = []
h_list = []
for k in range(0,11):
    h = 10**(-k)
    # create s with the right norm
    new_s = h/s_norm * s
    h list.append(h)
    norm_s_list.append(np.linalg.norm(new_s)**2)
    error_list.append(np.linalg.norm(F(y+new_s) - T_F(y,new_s)))
print(norm_s_list)
print(error_list)
plt.loglog(h_list, error_list, label = '$|| F(\\vec y +\\vec s) - T_F(\\vec_\)
 \rightarrowy,\\vec s)||_2$')
plt.loglog(h_list, norm_s_list, label = '$(||s||_2)^2$')
plt.xlabel('$||s|| 2$')
plt.title('Error of the first order Taylor approximation compared to $||s||_2$')
plt.legend()
plt.show()
                                                                    2.38546465]
T_F(y,s) = \begin{bmatrix} 2.38546465 & 7.46294666 & -51.71360771 & 7.46294666 \end{bmatrix}
F(y+s) = [ 2.38560102 7.46331735 -51.71260007 7.46331735 ]
                                                                  2.38560102]
[0.9999999999999, 0.00999999999999, 0.0001, 1e-06, 1e-08,
1.0000000000000002e-10, 1e-12, 9.999999999998e-15, 1.000000000000001e-16,
1e-18, 1.000000000000001e-20]
[485.50991502569156, 4.403382566259102, 0.043616858204482134,
0.00043575487691715163, 4.357135051294295e-06, 4.3563044426600625e-08,
4.3330658480417245e-10, 9.6611339270968e-12, 1.0180810756198736e-12,
6.77605788862772e-12, 9.10599483307802e-13]
```

Error of the first order Taylor approximation compared to $||s||_2$



5.5 (e) (2 pts)

- 1. Finish the function NewtonSolve below to solve the system of equations.
- 2. Take n=40, and experiment with your function. Try to find a choice of y0 such that the method doesn't converge, as well as a choice of y0 such that the method converges. In your answer, list the types of convergence behavior you found. Show a convergent example (if you found any) and a nonconvergent example (if you found any). Show the solutions you found for each example.

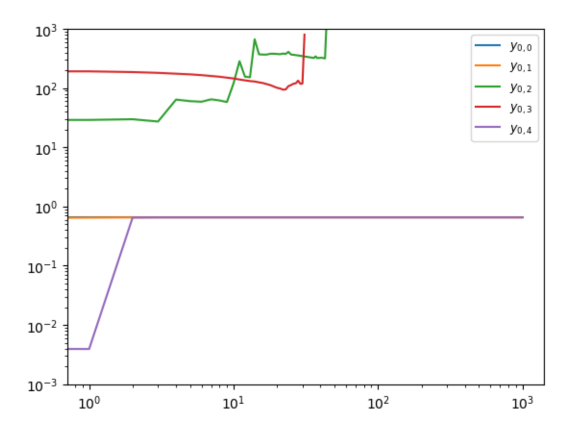
```
[]: # Subquestion 1.
def NewtonSolve(y0, K):
    """ Use Newton's method to solve F(y) = 0 with initial guess y0 and K
    iterations.
    Input: y0 (initial guess), K 9number of iterations
    Output: approximate solution to the root problem """
    y = y0
    newton_iterations = []
    for _ in range(K):
        sk = spsolve(JacobianF(y), -F(y))
```

```
y += sk
             newton_iterations.append(np.linalg.norm(y))
         return y,newton_iterations
     y0 = [1,2,1]
     K = 10
     print(NewtonSolve(y0,K))
    (array([0.10544867, 0.14144676, 0.10544867]), [1.3105006689808303,
    0.15731114883711553, 0.20544229052823915, 0.20553838885786144,
    0.205538389248722, 0.20553838924872203, 0.205538389248722, 0.20553838924872203,
    0.205538389248722, 0.20553838924872203])
[]: # Subquestion 2, code part
     y0 0 = np.zeros(40)
     y0_1 = np.ones(40)
     y0_2 = np.full(40,4, dtype = float)
     y0_3 = np.concatenate((np.full(9,4),np.full(31,1)), dtype=float)
     y0_4 = np.concatenate((np.full(8,4),np.full(32,1)), dtype=float)
     K = 1000
     newton_iterations = []
     y0,newton_iterations = NewtonSolve(y0_0, K)
     y1, newton_iterations_1 = NewtonSolve(y0_1, K)
     y2, newton_iterations_2 = NewtonSolve(y0_2, K)
     y3, newton_iterations_3 = NewtonSolve(y0_3, K)
     y4, newton_iterations_4 = NewtonSolve(y0_4, K)
     plt.loglog(range(K), newton_iterations, label = '$y_{0,0}$')
     plt.loglog(range(K), newton_iterations_1, label = '$y_{0,1}$')
     plt.loglog(range(K), newton_iterations_2, label = '$y_{0,2}$')
     plt.loglog(range(K), newton_iterations_3, label = '$y {0,3}$')
     plt.loglog(range(K), newton_iterations_4, label = '$y_{0,4}$')
     plt.legend(fontsize=9)
    plt.ylim(10**(-3),10**3)
    /var/folders/wr/_f23fm512gg3qhxthyg301vc0000gn/T/ipykernel_1070/3664664084.py:11
    : RuntimeWarning: overflow encountered in exp
      diag_list.append((1/h**2)*(-2) + np.exp(yi))
    /var/folders/wr/_f23fm512gg3qhxthyg30lvc0000gn/T/ipykernel_1070/3664664084.py:3:
    RuntimeWarning: overflow encountered in exp
      Ey = np.exp(np.array(y))
    /Users/loesbijman/anaconda3/envs/my-env/lib/python3.12/site-
```

packages/scipy/sparse/linalg/_dsolve/linsolve.py:293: MatrixRankWarning: Matrix
is exactly singular

warn("Matrix is exactly singular", MatrixRankWarning)

[]: (0.001, 1000)



We have tested the following initial conditions:

$$y_{0,0} = [0,...,0]^T;$$

$$y_{0,1} = [1, ..., 1]^T;$$

$$y_{0,2} = [4, ..., 4]^T;$$

 $y_{0,3} = [4, ..., 4, 1, ..., 1]^T$ with 9 fours in the beginning and 31 ones;

 $y_{0,4} = [4,...,4,1,...,1]^T$ with 8 fours in the beginning and 32 ones.

In the figure we see that the norm of the solution diverges for $y_{0,3}$ and $y_{0,4}$. The other solutions converge to the same norm. We thus see that small changes in y_0 can have large impacts on the divergence of Newton's method.