

Calcul Scientifique

Projet de Calcul Scientifique

Groupe EF06

Élèves :

THEVENET Louis
SABLAYROLLES Guillaume

1.

1.1.

Matrix dimension	Matrix type	Exec. time for eig (s)	Exec. time for power_v11, (s)
200×200	Type 1	9.000e-02	1.510e+00
400×400	Type 1	4.000e-02	1.831e + 01
600×600	Type 1	6.000e-02	6.021e+01
200×200	Type 2	3.000e-02	3.000e-02
400×400	Type 2	4.000e-02	4.000e-02
600×600	Type 2	7.000e-02	1.700e-01
200×200	Type 3	1.000e-02	5.000e-02
400×400	Type 3	3.000e-02	5.200e-01
600×600	Type 3	7.000e-02	1.270e+00
200×200	Type 4	2.000e-02	1.670e + 00
400×400	Type 4	3.000e-02	2.094e+01
600×600	Type 4	6.000e-02	5.456e + 01

Table 1: Execution time for different sizes and types of matrices

We can see that the power_v11 algorithm is generally slower than the eigen function especially for the type 2 and 4 matrices.

1.2.

```
nb_it = 1;
norme = norm(beta*v - z, 2)/norm(beta,2);

while(norme > eps && nb_it < maxit)
    beta_old = beta;
    v = z/norm(z, 2);
    z = A*v;
    beta = (v'*z)/(v'*v);
    norme = abs(beta-beta_old)/abs(beta_old);
    nb_it = nb_it + 1;
end</pre>
```

Listing 1: Inner loop of the new algorithm

Matrix dimension	Matrix type	Exec. time for $power_v11$, (s)	Exec. time for $power_v12$, (s)
200×200	Type 1	1.960e+00	3.200e-01
400×400	Type 1	1.888e+01	2.660e+00
600×600	Type 1	5.031e+01	7.070e+00
200×200	Type 2	1.000e-02	1.000e-02
400×400	Type 2	7.000e-02	1.000e-02
600×600	Type 2	1.800e-01	4.000e-02
200×200	Type 3	3.000e-02	1.000e-02
400×400	Type 3	6.100e-01	1.100e-01
600×600	Type 3	1.270e+00	2.600e-01

200×200	Type 4	1.530e+00	2.900e-01
400×400	Type 4	2.113e+01	3.060e+00
600×600	Type 4	5.914 + e01	6.480e+00

We can see that the power_v12 algorithm is globally faster than the power_v11.

1.3.

The main drawback of the deflated power method is the numerous matrix-vector products required to compute the eigenvectors as well as the fact that each iteration compute only one eigenvalue which can be slow if a lot of eigenvalues are desired.

1.4.

If we apply Algorithm 1 to m vectors, there is no reason for the columns of V to converge to a base. Each vector will converge toward a different projection of the dominant eigenvalue.

1.5.

In Algorithm 2, the matrix H is a smaller matrix, with dimension $n \times m$, therefore, even for larger matrices A, computing the spectral decomposition of H will not be computionally expensive.

1.6.

1.7.

- 1: function Subspace Iter v1 (Raleigh-Ritz Projection)
- 2: **Input**: $A \in \mathbb{R}^{n \times n}$, ε , MaxIter, PercentTrace
- 3: Output : $n_{\rm ev}$ dominant eigenvectors $V_{\rm out}$ and the corresponding eigenvalues $\Lambda_{\rm out}$
- 4: Generate an initial set of m orthonormal vectors $V \in \mathbb{R}^{n \times m}$; k = 0; PercentReached = 0
- 5: repeat until (line 52) PercentReached > PercentTrace $\forall n_{\text{ev}} = m \lor k > \text{MaxIter}$
- 6: $k \leftarrow k + 1$
- 7: Compute Y such that $Y = A \cdot V$
- 8: $V \leftarrow$ orthonormalisation of the columns of Y
- 9: Rayleigh-Ritz projection (line 60) applied on matrix A and orthonormal vectors V
- 10: Convergence analysis step (line 70): save eigenpairs that converged and update PercentReached

1.8.

There is $A \in \mathbb{R}^{n \times n}$, a square matrix, for $A \times A$ there are $\mathbb{O}(n^3)$ Flops. Since repeating p times the operation, there are $\mathbb{O}(p \times n^3)$ Flops to compute A^p . After, there is $V \in \mathbb{R}^{n \times m}$ which leads to $\mathbb{O}(p \times n^3 \times m)$ Flops to compute $A^p \times V$.

1.9.

1.10.

Matrix	dimen-	Matrix type	Iterations f	or	Iterations	for	Iterations	for	$p(A^p)$
sic	n		subspace_ite	r0	subspace_it	er1	subspace_i	ter2	
200 ×	200	Type 1	2309		263		132		2
200 ×	200	Type 1	2309		263		88		3
200 ×	200	Type 1	2309		263		53		5
200 ×	< 200	Type 1	2309		263		27		10

Table 2: After computing

When increasing the value of p to compute A^p in $subspace_iter2$, the number of flops to compute the results is: Iterations(iter2) $\simeq \frac{\text{Iterations(iter2)}}{p}$. This is due to the eigenvalues of A^p raised to the p power also. Increasing the conditionment of the matrix leading to a faster convergence.

1.11.

The accuracy differs because eigenpairs are computed from the columns of new matrix V. However, the firsts columns are recalculated at each step, that will lead to have different approximate size for the new eigenpairs computed. By recomputing them, the quality reduces and diverge from the approximate size of the first one.

1.12.

By freezing the converged columns, the algorithm will not have to recalculate them everytime. Which means that the accuracy for the eigenpairs will be more equal. The first and last will have the same approximate size.

2.

2.1.

 Σ_k is of size (k, k)

 U_k is of size (q, k)

 V_k is of size (p,k)

2.2.

eps	10^{-8}
maxit	10000
search_space	400
percentage	0.995
puiss	1

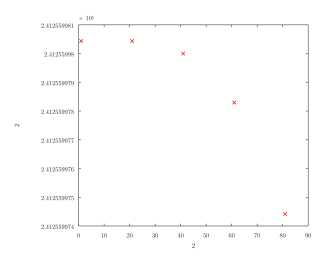
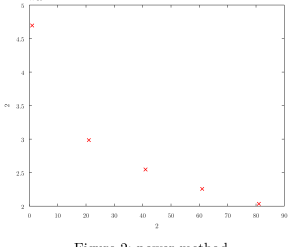


Figure 1: eig method



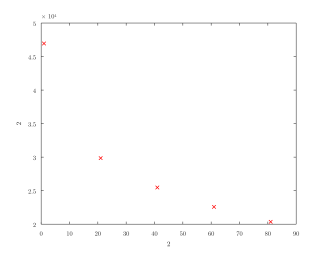
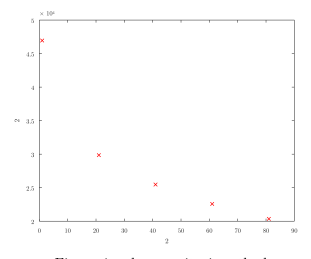
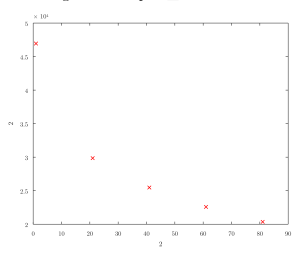


Figure 2: power method

Figure 3: subspace_iter0 method





 $Figure \ 4: subspace_iter1 \ method$

Figure 5: subspace_iter2 method

eps	10^{-8}
maxit	7500
search_space	600
percentage	0.995
puiss	2

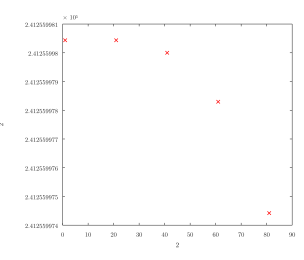
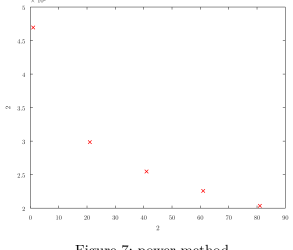


Figure 6: eig method



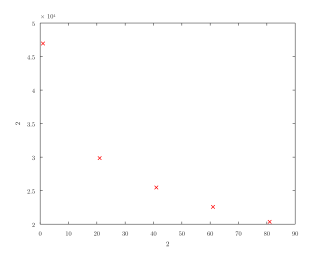
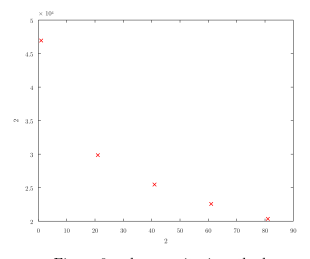
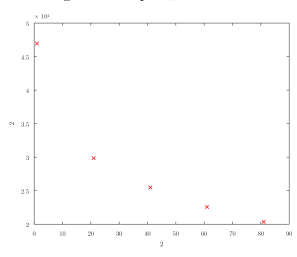


Figure 7: power method

Figure 8: subspace_iter0 method





 $Figure \ 9: subspace_iter1 \ method$

Figure 10: $subspace_iter2 method$

eps	10^{-8}
maxit	3000
search_space	500
percentage	0.995
puiss	1

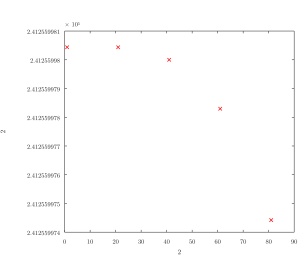


Figure 11: eig method

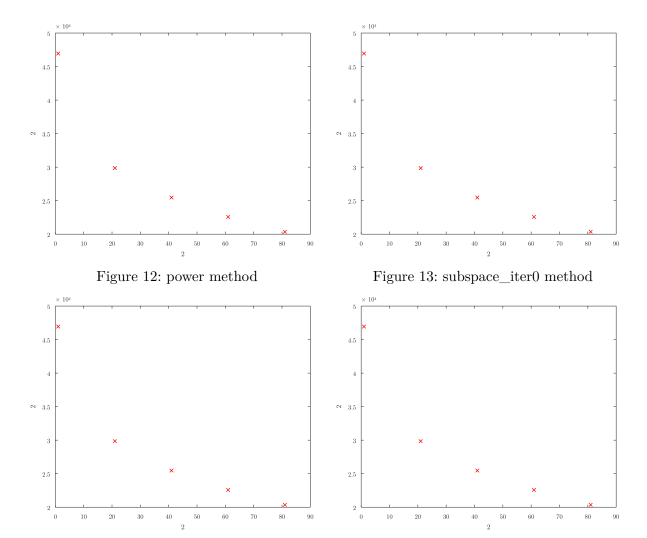


Figure 14: $subspace_iter1$ method

Figure 15: $subspace_iter2 method$