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Trilogy on Computing Maximal Eigenpair

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Abstract The eigenpair here means the twins consist of eigenvalue and its eigenvector. This paper introduces the three steps of our study on computing the maximal eigenpair. In the first two steps, we construct efficient initials for a known but dangerous algorithm, first for tridiagonal matrices and then for irreducible matrices, having nonnegative off-diagonal elements. In the third step, we present two global algorithms which are still efficient and work well for a quite large class of matrices, even complex for instance.

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1 Introduction

This paper is a continuation of [4]. For the reader's convenience, we review shortly the first part of [4], especially the story of the proportion of 1000 and 2 of iterations for two different algorithms.

The most famous result on the maximal eigenpair should be the Perron-Frobenius theorem. For nonnegative (pointwise) and irreducible A, if Trace (A) > 0, then the theorem says there exists uniquely a maximal eigenvalue $\rho(A) > 0$ with positive left-eigenvector u and positive right-eigenvector g such that

$$uA = \lambda u, \qquad Ag = \lambda g, \qquad \lambda = \rho(A).$$

These eigenvectors are also unique up to a constant. Before going to the main body of the paper, let us make two remarks.

- 1) We need to study the right-eigenvector g only. Otherwise, use the transpose A^* instead of A.
- 2) The matrix A is required to be irreducible with nonnegative off-diagonal elements, its diagonal elements can be arbitrary. Otherwise, use a shift A+mI for large m:

$$(A+mI)g = \lambda g \iff Ag = (\lambda - m)g, \tag{1}$$

their eigenvector remains the same but the maximal eigenvalues are shifted to each other.

Consider the following matrix:

$$Q = \begin{pmatrix} -1^2 & 1^2 & 0 & 0 & \cdots \\ 1^2 & -1^2 - 2^2 & 2^2 & 0 & \cdots \\ 0 & 2^2 & -2^2 - 3^2 & 3^2 & \cdots \\ \vdots & \vdots & \ddots & \ddots & \ddots \\ 0 & 0 & 0 & N^2 & -N^2 - (N+1)^2 \end{pmatrix}.$$
(2)

The main character of the matrix is the sequence $\{k^2\}$. The sum of each row equals zero except the last row. Actually, this matrix is truncated from the corresponding infinite one, in which case we have known that the maximal eigenvalue is -1/4 (refer to [2; Example 3.6]).

Example 1 Let N=7. Then the maximal eigenvalue is -0.525268 with eigenvector:

 $g \approx (55.878, 26.5271, 15.7059, 9.97983, 6.43129, 4.0251, 2.2954, 1)^*,$

where the vector v^* = the transpose of v.

We now want to practice the standard algorithms in matrix eigenvalue computation. The first method in computing the maximal eigenpair is the *Power Iteration*, introduced in 1929. Starting from a vector v_0 having a nonzero component in the direction of g, normalized with respect to a norm $\|\cdot\|$. At the kth step, iterate v_k by the formula

$$v_k = \frac{Av_{k-1}}{\|Av_{k-1}\|}, \quad z_k = \|Av_k\|, \qquad k \geqslant 1.$$
 (3)

Then we have the convergence: $v_k \to g$ (first pointwise and then uniformly) and $z_k \to \rho(Q)$ as $k \to \infty$. If we rewrite v_k as

$$v_k = \frac{A^k v_0}{\|A^k v_0\|},$$

one sees where the name "power" comes from. For our example, to use the Power Iteration, we adopt the ℓ^1 -norm and choose $v_0 = \tilde{v}_0/\|\tilde{v}_0\|$, where

 $\tilde{v}_0 = (1, 0.587624, 0.426178, 0.329975, 0.260701, 0.204394, 0.153593, 0.101142)^*$

This initial comes from a formula to be given in the next section. In Figure 1 below, the upper curve is g, the lower one is modified from \tilde{v}_0 , renormalized so that its last component becomes one. Clearly, these two functions are quite different, one may worry about the effectiveness of the choice of v_0 . Anyhow, having the experience of computing its eigensystem, I expect to finish the computation in a few of seconds. Unexpectly, I got a difficult time to compute

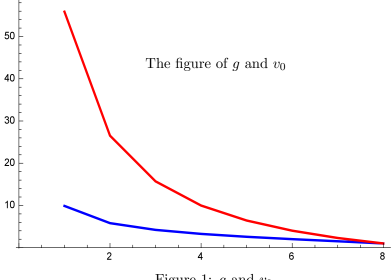
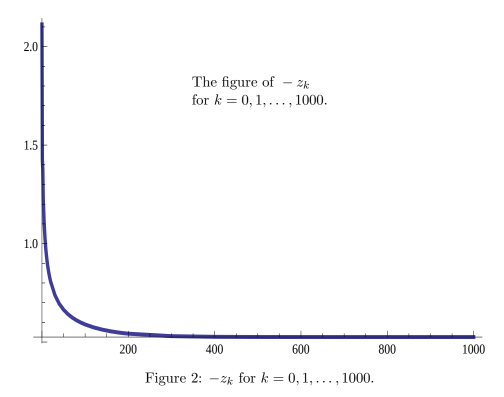


Figure 1: g and v_0 .

the maximal eigenpair for this simple example. Altogether, I computed it for 180 times, not in one day, using 1000 iterations. The printed pdf-file of the outputs has 64 pages. Figure 2 gives us the outputs.



The figure shows that the convergence of z_k goes quickly at the beginning of the iterations. This means that our initial v_0 is good enough. Then the

convergence goes very slow which means that the Power Iteration Algorithm converges very slowly.

Let us have a look at the convergence of the power iteration. Suppose that the eigenvalues are all different for simplicity. Denote by (λ_j, g_j) the eigenpairs with maximal one (λ_0, g_0) . Write $v_0 = \sum_{j=0}^{N} c_j g_j$ for some constants (c_j) . Then $c_0 \neq 0$ by assumption and

$$A^{k}v_{0} = \sum_{j=0}^{N} c_{j}\lambda_{j}^{k}g_{j} = c_{0}\lambda_{0}^{k} \left[g_{0} + \sum_{j=1}^{N} \frac{c_{j}}{c_{0}} \left(\frac{\lambda_{j}}{\lambda_{0}}\right)^{k}g_{j}\right].$$

Since $|\lambda_j/\lambda_0| < 1$ for each $j \ge 1$ and $||g_0|| = 1$, we have

$$\frac{A^k v_0}{\|A^k v_0\|} = \frac{c_0}{|c_0|} g_0 + O\left(\left|\frac{\lambda_1}{\lambda_0}\right|^k\right) \quad \text{as } k \to \infty,$$

where $|\lambda_1| := \max\{|\lambda_j| : j > 0\}$. Since $|\lambda_1/\lambda_0|$ can be very closed to 1, this explains the reason why the convergence of the method can be very slow.

Before moving further, let us mention that the power method can be also used to compute the minimal eigenvalue $\lambda_{\min}(A)$, simply replace A by A^{-1} . That is the *Inverse Iteration* introduced in 1944:

$$v_k = \frac{A^{-1}v_{k-1}}{\|A^{-1}v_{k-1}\|} \Longleftrightarrow v_k = \frac{A^{-k}v_0}{\|A^{-k}v_0\|}.$$
 (4)

It is interesting to note that the equivalent assertion on the right-hand side is exactly the the input-output method in economy.

To come back to compute the maximal $\rho(A)$ rather than $\lambda_{\min}(A)$, we add a shift z to A: replacing A by A-zI. Actually, it is even better to replace the last one by zI-A since we will often use $z>\rho(A)$ rather than $z<\rho(A)$, the details will be explained at the beginning of Section 4 below. When z is close enough to $\rho(A)$, the leading eigenvalue of $(zI-A)^{-1}$ becomes $(z-\rho(A))^{-1}$. Furthermore, we can even use a variant shift $z_{k-1}I$ to accelerate the convergence speed. Throughout this paper, we use varying shifts rather than a fixed one only. Thus, we have arrived at the second algorithm in computing the maximal eigenpair, the Rayleigh Quotient Iteration (RQI), a variant of the Inverse Iteration. From now on, unless otherwise stated, we often use the ℓ^2 -norm. Starting from an approximating pair (z_0, v_0) of the maximal one $(\rho(A), g)$ with $v_0^*v_0 = 1$, use the following iteration.

$$v_k = \frac{(z_{k-1}I - A)^{-1}v_{k-1}}{\|(z_{k-1}I - A)^{-1}v_{k-1}\|}, \qquad z_k = v_k^* A v_k, \qquad k \geqslant 1.$$
 (5)

If (z_0, v_0) is close enough to $(\rho(A), g)$, then

$$v_k \to g$$
 and $z_k \to \rho(A)$ as $k \to \infty$.

Since for each $k \ge 1$, $v_k^* v_k = 1$, we have $z_k = v_k^* A v_k / (v_k^* v_k)$. That is where the name "Rayleigh Quotient" comes from. Unless otherwise stated, z_0 is setting to be $v_0^* A v_0$.

Having the hard time spent in the first algorithm, I wondered how many iterations are required using this algorithm. Of course, I can no longer bear 1000 iterations. To be honest, I hope to finish the computation within 100 iterations. What happens now?

Example 2 For the same matrix Q and \tilde{v}_0 as in Example 1, by RQI, we need two iterations only:

$$z_1 \approx -0.528215$$
, $z_2 \approx -0.525268$.

The result came to me, not enough to say surprisingly, I was shocked indeed. This shows not only the power of the second method but also the effectiveness of my initial v_0 . From the examples above, we have seen the story of the proportion of 1000 and 2.

For simplicity, from now on, we often write $\lambda_j := \lambda_j(-Q)$. In particular $\lambda_0 = -\rho(Q) > 0$. Instead of our previous v_0 , we adopt the uniform distribution:

$$v_0 = (1, 1, 1, 1, 1, 1, 1, 1)^* / \sqrt{8}.$$

This is somehow fair since we usually have no knowledge about g in advance.

Example 3 Let Q be the same as above. Use the uniform distribution v_0 and set $z_0 = v_0^*(-Q)v_0$. Then

$$(z_1, z_2, z_3, \mathbf{z_4}) \approx (4.78557, 5.67061, 5.91766, \mathbf{5.91867}).$$

 $(\lambda_0, \lambda_1, \lambda_2) \approx (0.525268, 2.00758, \mathbf{5.91867}).$

The computation becomes stable at the 4th iteration. Unfortunately, it is not what we want λ_0 but λ_2 . In other words, the algorithm converges to a pitfall. Very often, there are n-1 pitfalls for a matrix having n eigenvalues. This shows once again our initial \tilde{v}_0 is efficient and the RQI is quite dangerous.

Hopefully, everyone here has heard the name *Google's PageRank*. In other words, the Google's search is based on the maximal left-eigenvector. On this topic, the book [8] was published 11 years ago. In this book, the Power Iteration is included but not the RQI. It should be clear that for PageRank, we need to consider not only large system, but also fast algorithm.

It may be the correct position to mention a part of the motivations for the present study.

- Google's search—PageRank.
- Input—output method in economy. In this and the previous cases, the computation of the maximal eigenvector is required.
- Stability speed of stochastic systems. Here, for the stationary distribution of a Markov chain, we need to compute the eigenvector; and for the stability rate, we need to study the maximal (or the fist nontrivial) eigenvalue.

 Principal component analysis for BigData. One choice is to study the socalled five-diagonal matrices. The second approach is using the maximal eigenvector to analysis the role played by the components, somehow similar to the PageRank.

- For image recognition, one often uses Poisson or Toeplitz matrices, which are more or less the same as the Quasi-birth-death matrices studied in queueing theory. The discrete difference equations of elliptic partial differential equations are included in this class: the block-tridiagonal matrices.
- The effectiveness of random algorithm, say Markov Chain Monte Carlo for instance, is described by the convergence speed. This is also related to the algorithms for machine learning.
- As in the last item, a mathematical tool to describe the phase transitions is the first nontrivial eigenvalue (the next eigenpair in general). This is the original place where the author was attracted to the topic.

Since the wide range of the applications of the topic, there is a large number of publications. The author is unable to present a carefully chosen list of references here, what instead are two random selected references: [8] and [11].

Up to now, we have discussed only a small size $8 \times 8 \, (N=7)$ matrix. How about large N? In computational mathematics, one often expects the number of iterations grows in a polynomial way N^{α} for α greater or equal to 1. In our efficient case, since $2=8^{1/3}$, we expect to have $10000^{1/3}\approx 22$ iterations for $N+1=10^4$. The next table subverts completely my imagination.

N+1	z_0	z_1	$z_2 = \lambda_0$	upper/lower
8	0.523309	0.525268	0.525268	$1+10^{-11}$
100	0.387333	0.376393	0.376383	$1+10^{-8}$
500	0.349147	0.338342	0.338329	$1+10^{-7}$
1000	0.338027	0.327254	0.32724	$1+10^{-7}$
5000	0.319895	0.30855	0.308529	$1+10^{-7}$
7500	0.316529	0.304942	0.304918	$1+10^{-7}$
10^{4}	0.31437	0.302586	0.302561	$1+10^{-7}$

Table 1 Comparison of RQI for different N

Here z_0 is defined by

$$z_0 = 7/(8\delta_1) + v_0^*(-Q)v_0/8,$$

where v_0 and δ_1 are computed by our general formulas to be defined in the next section. We compute the matrices of order $8, 100, \ldots, 10^4$ by using MatLab in a notebook, in no more than 30 seconds, the iterations finish at the second step. This means that the outputs starting from z_2 are the same and coincide with λ_0 . See the first row for instance, which becomes stable at the first step indeed. We do not believe such a result for some days, so we checked it in

different ways. First, since $\lambda_0 = 1/4$ when $N = \infty$, the answers of λ_0 given in the fourth column are reasonable. More essentially, by using the output v_2 , we can deduce upper and lower bounds of λ_0 (using [2; Theorem 2.4(3)]), and then the ratio upper/ lower is presented in the last column. In each case, the algorithm is significant up to 6 digits. For the large scale matrices here and in 4, the computations are completed by Yue-Shuang Li.

2 Efficient initials: tridiagonal case

It is the position to write down the formulas of v_0 and δ_1 . Then our initial z_0 used in Table 1 is a little modification of δ_1^{-1} : a convex combination of δ_1^{-1} and $v_0^*(-Q)v_0$.

Let us consider the tridiagonal matrix (cf. [3; §3] and [6; §4.4]). Fix $N \ge 1$, denote by $E = \{0, 1, ..., N\}$ the set of indices. By a shift if necessary, we may reduce A to Q with negative diagonals: $Q^c = A - mI$, $m := \max_{i \in E} \sum_{j \in E} a_{ij}$,

$$Q^{c} = \begin{pmatrix} -b_{0} - c_{0} & b_{0} & 0 & 0 & \cdots \\ a_{1} & -a_{1} - b_{1} - c_{1} & b_{1} & 0 & \cdots \\ 0 & a_{2} & -a_{2} - b_{2} - c_{2} & b_{2} & \cdots \\ \vdots & \vdots & \ddots & \ddots & \ddots \\ 0 & 0 & 0 & a_{N} & -a_{N} - c_{N} \end{pmatrix}.$$

Thus, we have three sequences $\{a_i > 0\}$, $\{b_i > 0\}$, and $\{c_i \ge 0\}$. Our main assumption here is that the first two sequences are positive and $c_i \not\equiv 0$. In order to define our initials, we need three new sequences, $\{h_k\}$, $\{\mu_k\}$, and $\{\varphi_k\}$.

First, we define the sequence $\{h_k\}$:

$$h_0 = 1, \ h_n = h_{n-1}r_{n-1}, \qquad 1 \leqslant n \leqslant N;$$
 (6)

here we need another sequence $\{r_k\}$:

$$r_0 = 1 + \frac{c_0}{b_0}, \quad r_n = 1 + \frac{a_n + c_n}{b_n} - \frac{a_n}{b_n r_{n-1}}, \qquad 1 \leqslant n < N.$$

Here and in what follows, our iterations are often of one-step. Note that if $c_k = 0$ for every k < N, then we do not need the sequence $\{h_k\}$, simply set $h_k \equiv 1$. An easier way to remember this (h_i) is as follows. It is nearly harmonic of Q^c except at the last point N:

$$Q^{c \text{ the last row}} h = 0,$$
 (7)

where $B^{\text{the last row}}$ means the matrix modified from B by removing its last low.

We now use H-transform, it is designed to remove the sequence (c_i) :

$$\widetilde{Q} = \operatorname{Diag}(h_i)^{-1} Q^c \operatorname{Diag}(h_i).$$

Then

$$\widetilde{Q} = \begin{pmatrix} -b_0 & b_0 & 0 & 0 & \cdots \\ a_1 & -a_1 - b_1 & b_1 & 0 & \cdots \\ 0 & a_2 & -a_2 - b_2 & b_2 & \cdots \\ \vdots & \vdots & \ddots & \ddots & \ddots \\ 0 & 0 & 0 & a_N & -a_N - c_N \end{pmatrix}$$

for some modified $\{a_i > 0\}$, $\{b_i > 0\}$, and $c_N > 0$. Of course, Q^c and \widetilde{Q} have the same spectrum. In particular, under the H-transform,

$$(\lambda_{\min}(-Q^c), g) \to (\lambda_{\min}(-\widetilde{Q}) = \lambda_{\min}(-Q^c), \operatorname{Diag}(h_i)^{-1}g).$$

From now on, for simplicity, we denote by Q the matrix replacing c_N by b_N in \widetilde{Q} .

Next, we define the second sequence $\{\mu_k\}$:

$$\mu_0 = 1, \quad \mu_n = \mu_{n-1} \frac{b_{n-1}}{a_n}, \qquad 1 \leqslant n \leqslant N.$$
 (8)

And then define the third one $\{\varphi_k\}$ as follows:

$$\varphi_n = \sum_{k=n}^N \frac{1}{\mu_k b_k}, \qquad 0 \leqslant n \leqslant N. \tag{9}$$

We are now ready to define v_0 and δ_1 (or z_0) using the sequences (μ_i) and (φ_i) .

$$\tilde{v}_0(i) = \sqrt{\varphi_i}, \ i \leqslant N; \qquad v_0 = \tilde{v}_0 / \|\tilde{v}_0\|; \quad \|\cdot\| := \|\cdot\|_{L^2(\mu)}$$
 (10)

$$\delta_1 = \max_{0 \le n \le N} \left[\sqrt{\varphi_n} \sum_{k=0}^n \mu_k \sqrt{\varphi_k} + \frac{1}{\sqrt{\varphi_n}} \sum_{n+1 \le j \le N} \mu_j \varphi_j^{3/2} \right] =: z_0^{-1}$$
 (11)

with a convention $\sum_{\emptyset} = 0$.

Finally, having constructed the initials (v_0, z_0) , the RQI goes as follows. Solve w_k :

$$(-Q - z_{k-1}I)w_k = v_{k-1}, \qquad k \geqslant 1;$$
 (12)

and define

$$v_k = w_k / ||w_k||, \qquad z_k = (v_k, -Q v_k)_{L^2(\mu)}.$$

Then

$$v_k \to g$$
 and $z_k \to \lambda_0$ as $k \to \infty$.

Before moving further, let us mention that there is an explicit representation of the solution (w_i) to equation (12). Assume that we are given $v := v_{k-1}$ and $z := z_{k-1}$. Set

$$M_{sj} = \mu_j \sum_{k=j}^{s} \frac{1}{\mu_k b_k}, \qquad 0 \leqslant j \leqslant s \leqslant N.$$
 (13)

Define two independent sequences $\{A(s)\}\$ and $\{B(s)\}\$, recurrently:

$$\begin{cases}
A(s) = -\sum_{0 \le j \le s-1} M_{s-1,j} (v(j) + zA(j)), \\
B(s) = 1 - z \sum_{0 \le j \le s-1} M_{s-1,j} B(j), & 0 \le s \le N.
\end{cases}$$
(14)

Set

$$x = \frac{\sum_{j=0}^{N} \mu_j (v(j) + zA(j)) - \mu_N b_N A(N)}{\mu_N b_N B(N) - z \sum_{j=0}^{N} \mu_j B(j)}.$$
 (15)

Then the required solution $w_k := \{w(s) : s \in E\}$ can be expressed as w(s) = A(s) + xB(s) $(s \in E)$.

To finish the algorithm, we return to the estimates of $(\lambda_{\min}(-Q^c), g(Q^c))$ $(g(Q^c) = g(-Q^c))$ or further $(\rho(A), g(A))$ if necessary, where g(A), for instance, denotes the maximal eigenvector of A. Suppose that the iterations are stopped at $k = k_0$ and set $(\bar{z}, \bar{v}) = (z_{k_0}, v_{k_0})$ for simplicity. Then, we have

$$(\lambda_{\min}(-Q^c), \operatorname{Diag}(h_i)^{-1}g(Q^c)) = (\lambda_{\min}(-\widetilde{Q}), g(\widetilde{Q})) \approx (\overline{z}, \overline{v}),$$

and so

$$(\lambda_{\min}(-Q^c), g(Q^c)) \approx (\bar{z}, \operatorname{Diag}(h_i)\bar{v}).$$
 (16)

Because $\lambda_{\min}(-Q^c) = m - \rho(A)$, we obtain

$$(\rho(A), g(A)) \approx (m - \bar{z}, \operatorname{Diag}(h_i) \bar{v}).$$
 (17)

Now, the question is the possibility from the tridiagonal case to the general one.

3 Efficient initials: the general case ([3; $\S 4.2$] and [6; $\S 4.5$])

When we first look at the question just mentioned, it seems quite a long distance to go from the special tridiagonal case to the general one. However, in the eigenvalue computation theory, there is the so-called Lanczos tridiagonalization procedure to handle the job, as discussed in [3; Appendix of §3]. Nevertheless, what we adopted in [3; §4] is a completely different approach. Here is our main idea. Note that the initials v_0 and δ_1 constructed in the last section are explicitly expressed by the new sequences. In other words, we have used three new sequences $\{h_k\}$, $\{\mu_k\}$, and $\{\varphi_k\}$ instead of the original three $\{a_i\}$, $\{b_i\}$, and $\{c_i\}$ to describe our initials. Very fortunately, the former three sequences do have clearly the probabilistic meaning, which then leads us a way to go to the general setup. Shortly, we construct these sequences by solving three linear equations (usually, we do not have explicit solution in such a general setup). Then use them to construct the initials and further apply the RQI-algorithm.

Let $A = (a_{ij} : i, j \in E)$ be the same as given at the beginning of the paper. Set $A_i = \sum_{j \in E} a_{ij}$ and define

$$Q^c = A - \Big(\max_{i \in E} A_i\Big)I.$$

We can now state the probabilistic/analytic meaning of the required three sequences (h_i) , (μ_i) , and (φ_i) .

- (h_i) is the harmonic function of Q^c except at the right endpoint N, as mentioned in the last section.
- (μ_i) is the invariant measure (stationary distribution) of the matrix Q^c removing the sequence (c_i) .
- (φ_i) is the tail related to the transiency series, refer to [3; Lemma 24 and its proof].

We now begin with our construction. Let $h=(h_0,h_1,\ldots,h_N)^*$ (with $h_0=1$) solve the equation

$$Q^{c \text{ the last row}} h = 0$$

and define

$$\widetilde{Q} = \operatorname{Diag}(h_i)^{-1} Q^c \operatorname{Diag}(h_i).$$

Then for which we have

$$c_0 = \ldots = c_{N-1} = 0, \quad c_N =: q_{N,N+1} > 0.$$

This is very much similar to the tridiagonal case.

Next, set Q = Q. Let $\varphi = (\varphi_0, \varphi_1, \dots, \varphi_N)^*$ (with $\varphi_0 = 1$) solve the equation

$$\varphi^{\text{the first row}} = P^{\text{the first row}} \, \varphi.$$

where

$$P = \operatorname{Diag}((-q_{ii})^{-1})Q + I.$$

Thirdly, assume that $\mu := (\mu_0, \mu_1, \dots, \mu_N)$ with $\mu_0 = 1$ solves the equation

$$Q^{*}$$
 the last row $\mu^* = 0$.

Having these sequences at hand, we can define the initials

$$\tilde{v}_0(i) = \sqrt{\varphi_i}, \ i \leqslant N; \quad v_0 = \tilde{v}_0 / \|\tilde{v}_0\|_{\mu}; \quad z_0 = (v_0, -Qv_0)_{\mu}.$$

Then, go to the RQI as usual. For $k \ge 1$, let w_k solve the equation

$$(-Q - z_{k-1}I)w_k = v_{k-1}$$

and set

$$v_k = w_k / ||w_k||_{\mu}, \qquad z_k = (v_k, -Qv_k)_{\mu}.$$

Then we often have $(z_k, v_k) \to (\lambda_0, g)$ as $k \to \infty$.

We remark that there is an alternative choice (more safe) of z_0 :

$$z_0^{-1} = \frac{1}{1 - \varphi_1} \max_{0 \leqslant n \leqslant N} \left[\sqrt{\varphi_n} \sum_{k=0}^n \mu_k \sqrt{\varphi_k} + \frac{1}{\sqrt{\varphi_n}} \sum_{n+1 \leqslant j \leqslant N} \mu_j \varphi_j^{3/2} \right]$$

which is almost a copy of the one used in the last section.

The procedure for returning to the estimates of $(\lambda_{\min}(-Q^c), g(Q^c))$ or further $(\rho(A), g(A))$ is very much the same as in the last section.

To conclude this section, we introduce two examples to illustrate the efficiency of the extended initials for tridiagonally dominant matrices. The next two examples were computed by Xu Zhu, a master student in Shanghai.

Example 4 (Block-tridiagonal matrix) Consider the matrix

$$Q = \begin{pmatrix} A_0 & B_0 & 0 & 0 & \cdots \\ C_1 & A_1 & B_1 & 0 & \cdots \\ 0 & C_2 & A_2 & B_2 & \cdots \\ \vdots & \vdots & \ddots & \ddots & \ddots \\ 0 & 0 & 0 & C_N & A_N \end{pmatrix},$$

where A_k, B_k, C_k are 40×40 -matrices, B's and C's are identity matrices, and A's are tridiagonal matrices. For this model, two iterations are enough to arrive at the required results (Table 2).

 Table 2
 Outputs for Poisson matrix

N+1	z_0	z_1	$z_2 = \lambda_0$
1600	7.985026	7.988219	7.988263
3600	7.993232	7.994676	7.994696
6400	7.996161	7.988256	7.987972

Example 5 (Toeplitz matrix) Consider the matrix

$$A = \begin{pmatrix} 1 & 2 & 3 & \cdots & n-1 & n \\ 2 & 1 & 2 & \cdots & n-2 & n-1 \\ \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ n-1 & n-2 & n-3 & \cdots & 1 & 2 \\ n & n-1 & n-2 & \cdots & 2 & 1 \end{pmatrix}.$$

For this model, three iterations are enough to arrive at the required results (Table 3).

 Table 3
 Outputs for Toeplitz matrix

N+1	$z_0 \times 10^6$	$z_1 \times 10^6$	$z_2 \times 10^6$	$z_3 = \lambda_0$
1600	0.156992	0.451326	0.390252	0.389890
3600	0.157398	2.30731	1.97816	1.97591
6400	0.157450	7.32791	6.25506	6.24718

As mentioned before, the extended algorithm should be powerful for the tridiagonally dominant matrices. How about more general case? Two questions are often asked to me by specialists in computational mathematics: do you allow more negative off-diagonal elements? How about complex matrices? My answer is: they are too far away from me, since those matrices can not be a generator of a Markov chain, I do not have a tool to handle them. Alternatively, I have studied some more general matrices than the tridiagonal ones: the block-tridiagonal matrices, the lower triangular plus upper-diagonal, the upper triangular plus lower-diagonal, and so on. Certainly, we can do a lot case by case, but this seems still a long way to achieve a global algorithm. So we do need a different idea.

4 Global algorithms

Several months ago, AlphaGo came to my attention. From which I learnt the subject of machine learning. After some days, I suddenly thought, since we are doing the computational mathematics, why can not let the computer help us to find a high efficiency initial value? Why can not we leave this hard task to the computer? If so, then we can start from a relatively simple and common initial value, let the computer help us to gradually improve it.

The first step is easy, simply choose the uniform distribution as our initial v_0 :

$$v_0 = (1, 1, \cdots, 1)^* / \sqrt{N+1}.$$

As mentioned before, this initial vector is fair and universal. One may feel strange at the first look at "global" in the title of this section. However, with this universal v_0 , the power iteration is already a global algorithm. Unfortunately, the convergence of this method is too slow, and hence is often not practical. To quicken the speed, we should add a shift which now has a very heavy duty for our algorithm. The main trouble is that the usual Rayleigh quotient $v_0^*Av_0/(v_0^*v_0)$ can not be used as z_0 , otherwise, it will often lead to a pitfall, as illustrated by Example 3. The main reason is that our v_0 is too rough and so z_0 deduced from it is also too rough. Now, how to choose z_0 and further z_n ?

Clearly, for avoiding the pitfalls, we have to choose z_0 from the outside of the spectrum of A (denoted by $\mathrm{Sp}(A)$), and as close to $\rho(A)$ as possible to quicken the convergence speed. For nonnegative A, $\mathrm{Sp}(A)$ is located in a circle with radius $\rho(A)$ in the complex plane. Thus, the safe region should be on the outside of $\mathrm{Sp}(A)$. Since $\rho(A)$ is located at the boundary on the right-hand side of the circle, the effective area should be on the real axis on the right-hand side of, but a little away from, $\rho(A)$.

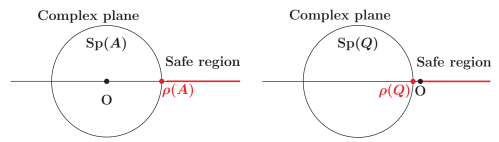


Figure 3: Safe region in complex plane.

For the matrix Q used in this paper, since $\rho(Q) < 0$, its spectrum Sp(Q) is located on the left-hand side of the origin. Then, one can simply choose $z_0 = 0$ as an initial. See Figure 3.

Having these idea in mind, we can now state two of our global algorithms. Each of them uses the same initials:

$$v_0 = \text{uniform distribution}, \qquad z_0 = \max_{0 \le i \le N} \frac{Av_0}{v_0}(i),$$

where for two vectors f and g, $(f/g)(i) = f_i/g_i$.

Algorithm 1 (Specific Rayleigh quotient iteration) At step $k \ge 1$, for given $v := v_{k-1}$ and $z := z_{k-1}$, let w solve the equation

$$(zI - A)w = v.$$

Set $v_k = w/\|w\|$ and let $z_k = v_k^* A v_k$.

This algorithm goes back to [3; §4.1 with Choice I].

Algorithm 2 (Shifted inverse iteration) Everything is the same as in Algorithm 1, except redefine z_k as follows:

$$z_k = \max_{0 \leqslant i \leqslant N} \frac{Av_k}{v_k}(i)$$

for $k \geqslant 1$ (or equivalently, $k \geqslant 0$).

The comparison of these algorithms is the following: with unknown small probability, Algorithm 1 is less safe than Algorithm 2, but the former one has a faster convergence speed than the latter one with possibility 1/5 for instance. A refined combination of the above two algorithms is presented in [6; §2], say Algorithm 4_2 for instance.

With the worrying on the safety and convergence speed in mind, we examine two examples which are non-symmetric.

The first example below is a lower triangular plus the upper-diagonal. It is far away from the tridiagonal one, we want to see what can be happened.

Example 6 ([6; Example 7]) Let

$$Q = \begin{pmatrix} -1 & 1 & 0 & 0 & \cdots & 0 & 0 \\ a_1 & -a_1 - 2 & 2 & 0 & \cdots & 0 & 0 \\ a_2 & 0 & -a_2 - 3 & 3 & \cdots & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \vdots & \cdots & N-1 & 0 \\ a_{N-1} & 0 & 0 & 0 & \cdots & -a_{N-1} - N & N \\ a_N & 0 & 0 & 0 & \cdots & 0 & -a_N - N-1 \end{pmatrix}.$$
(18)

For this matrix, we have computed several cases:

$$a_k = 1/(k+1), \ a_k \equiv 1, \ a_k = k, \ a_k = k^2.$$

Among them, the first one is the hardest and is hence presented below. For different N, the outputs of our algorithm are given in Table 4.

N+1	$egin{array}{c} ext{Table 4.} \ extbf{\emph{z}}_1 \end{array}$	$oldsymbol{z_1}$ ne outpu $oldsymbol{z_2}$	z_3	ent N by or $oldsymbol{z_4}$	ar algorithm z ₅	z_6
8	0.276727	0.427307	0.451902	0.452339		
16	0.222132	0.367827	0.399959	0.400910		
32	0.187826	0.329646	0.370364	0.372308	0.372311	
50	0.171657	0.311197	0.357814	0.360776	0.360784	
100	0.152106	0.287996	0.343847	0.349166	0.349197	
500	0.121403	0.247450	0.321751	0.336811	0.337186	
1000	0.111879	0.233257	0.313274	0.334155	0.335009	0.335010
5000	0.0947429	0.205212	0.293025	0.328961	0.332609	0.332635
10^{4}	0.0888963	0.194859	0.284064	0.326285	0.332113	0.332188

The next example is upper triangular plus lower-diagonal. It is motivated from the classical branching process. Denote by $(p_k : k \ge 0)$ a given probability measure with $p_1 = 0$. Let

$$Q = \begin{pmatrix} -1 & p_2 & p_3 & p_4 & \cdots & p_{N-1} & \sum_{k \geqslant N} p_k \\ 2p_0 & -2 & 2p_2 & 2p_3 & \cdots & 2p_{N-2} & 2\sum_{k \geqslant N-1} p_k \\ 0 & 3p_0 & -3 & 3p_2 & \cdots & 3p_{N-3} & 3\sum_{k \geqslant N-2} p_k \\ \vdots & \vdots & \vdots & \ddots & \ddots & \ddots & \ddots \\ \vdots & \vdots & \vdots & \ddots & \ddots & -(N-1) & (N-1)\sum_{k \geqslant 2} p_k \\ 0 & 0 & 0 & 0 & \cdots & Np_0 & -Np_0 \end{pmatrix}.$$

The matrix is defined on $E := \{1, 2, ..., N\}$. Set $M_1 = \sum_{k \in E} k p_k$. When $N = \infty$, it is subcritical iff $M_1 < 1$, to which the maximal eigenvalue should be positive. Otherwise, the convergence rate should be zero.

Now, we fix

$$p_0 = \alpha/2, \ p_1 = 0, \ p_2 = (2 - \alpha)/2^2, \ \dots p_n = (2 - \alpha)/2^n, \dots, \qquad \alpha \in (0, 2).$$

Then $M_1=3(2-\alpha)/2$ and hence we are in the subcritical case iff $\alpha\in(4/3,2).$

Example 7 ([6; Example 9]) Set $\alpha=7/4$. We want to know how fast the local $(N<\infty)$ maximal eigenvalue becomes stable (i.e., close enough to the converge rate at $N=\infty$). Up to $N=10^4$, the steps of the iterations we need are no more than 6. To quicken the convergence, we adopt an improved algorithm. Then the outputs of the approximation of the minimal eigenvalue of -Q for different N are given in Table 5.

Table 5. The outputs in the subcritical case						
$N^{-\alpha}$	z_1	z_2	z_3	z_4		
8	0.637800	0.638153				
16	0.621430	0.625490	0.625539			
50	0.609976	0.624052	0.624997	0.625000		
100	0.606948	0.623377	0.624991	0.625000		
500	0.604409	0.622116	0.624962	0.625000		
1000	0.604082	0.621688	0.624944	0.625000		
5000	0.603817	0.620838	0.62489	0.625000		
10^{4}	0.603784	0.620511	0.624861	0.625000		

The computation in each case costs no more than one minute. Besides, starting from N=50, the final outputs are all the same: 0.625, which then can be regarded as a very good approximation of $\lambda_{\min}(-Q)$ at infinity $N=\infty$.

It is the position to compare our global algorithm with that given in the last section. At the first look, here in the two examples above, we need about 6 iterations, double of the ones given in the last section. Note that for the initials of the algorithm in the last section, we need solve three additional linear equations, which are more or less the same as three additional iterations. Hence the efficiency of these two algorithms are very close to each other. Actually, the computation time used for the algorithm in the last section is much more than the new one here.

It is quite surprising that our new algorithms work for a much general class of matrices, out of the scope of [3]. Here we consider the maximal eigenpair only.

The example below allows partially negative off-diagonal elements.

Example 8 ([9; Example (7)], [6; Example 12]) Let

$$A = \begin{pmatrix} -1 & 8 & -1 \\ 8 & 8 & 8 \\ -1 & 8 & 8 \end{pmatrix}.$$

Then The eigenvalues of A are as follows.

$$17.5124, -7.4675, 4.95513.$$

The corresponding maximal eigenvector is

$$(0.486078, 1.24981, 1)^*$$

which is positive.

Started at $z_0 = 24$, the outputs of our algorithms are given in Table 6.

\boldsymbol{n}	z_n : Algorithm 1	z_n : Algorithm 2
1	17.3772	18.5316
2	17.5124	17.5416
3		17.5124

Table 6. The outputs for a matrix with more negative elements

Furthermore, we can even consider some complex matrices.

Example 9 ([10; Example 2.1], [6; Example 15]) Let

$$A = \begin{pmatrix} 0.75 - 1.125 \, i & 0.5882 - 0.1471 \, i & 1.0735 + 1.4191 \, i \\ -0.5 - i & 2.1765 + 0.7059 \, i & 2.1471 - 0.4118 \, i \\ 2.75 - 0.125 \, i & 0.5882 - 0.1471 \, i & -0.9265 + 0.4191 \, i \end{pmatrix},$$

where the coefficients are all accurate, to four decimal digits. Then ${\cal A}$ has eigenvalues

$$3, -2-i, 1+i$$

with maximal eigenvector

$$(0.408237, 0.816507, 0.408237)^*$$
.

The outputs (y_n) (but not (z_n)) of [6; Algorithm 14], a variant of Algorithm 2, are as follows.

y_1^{7} . The outp	uts for a complex matrix y_2	y_3
3.03949 - 0.0451599 i	3.00471 - 0.0015769 i	3

We mention that a simple sufficient condition for the use of our algorithms is the following:

$$\operatorname{Re}(A^n) > 0$$
 for large enough n , up to a shift mI . (19)

Then we have the Perron–Frobenius property: there exists the maximal eigenvalue $\rho(A) > 0$ having simple left- and right-eigenvectors.

Hopefully, the reader would now be accept the use of "global" here for our new algorithms. They are very much efficient indeed. One may ask about the convergence speed of the algorithms. Even though we do not have a universal estimate for each model in such a general setup, it is known however that the shifted inverse algorithm is a fast cubic one, and hence should be fast enough in practice. This explains the reason why our algorithms are fast enough in the general setup. Certainly, in the tridiagonal dominate case, one can use the algorithms presented in the previous sections. Especially, in the tridiagonal situation, we have analytically basic estimates which guarantee the efficiency of the algorithms. See [4] for a long way to reach the present level.

When talking about the eigenvalues, the first reaction for many people (at least for me, 30 years ago) is that well, we have known a great deal about the

subject. However, it is not the trues. One may ask himself that for eigenvalues, how large matrix have you computed by hand? As far as I know, 2×2 only in analytic computation by hand. It is not so easy to compute them for a 3×3 matrix, except using computer. Even I have worked on the topic for about 30 years, I have not been brave enough to compute the maximal eigenvector, we use its mimic only to estimate the maximal eigenvalue (or more generally the first nontrivial eigenvalue). The first paper I wrote on the numerical computation is [3]. It is known that the most algorithms in computational mathematics are local, the Newton algorithm (which is a quadratic algorithm) for instance. Hence, our global algorithms are somehow unusual.

About three years ago, I heard a lecture that dealt with a circuit board optimization problem. The author uses the Newton method. I said it was too dangerous and could fall into the trap. The speaker answered me that yes, it is dangerous, but no one in the world can solve this problem. Can we try annealing algorithm? I asked. He replied that it was too slow. We all know that in the global optimization, a big problem (not yet cracked) is how to escape from the local traps. The story we are talking about today seems to have opened a small hole for algorithms and optimization problems, and perhaps you will be here to create a new field.

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