EIGENEDGE package for MATLAB

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1 Introduction

The EigenEdge MATLAB package contains open source implementations of methods for working with eigenvalue distributions of large random matrices. In particular, it contains the Atomic method to compute the limit empirical spectrum of sample covariance matrices (proposed in Dobriban, 2015).

• Version: 0.0.1

• Requirements: Tested on MATLAB R2014b.

• Author: Edgar Dobriban

• License: GPL-3

In addition, this package contains the code to reproduce all simulation results from the paper Dobriban (2015). These are contained in the \Experiments\Atomic folder.

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2 Installation

Extract the archive in any folder, say to <path> . The main functions are in the Code directory, which needs to be on the Matlab path, along with all of its subfolders. This can be accomplished in at least three ways. First, you can add the following lines to your Matlab startup:

```
addpath('<path>/pvalue_weighting_matlab/Code')
addpath('<path>/pvalue_weighting_matlab/Code/Basic')
addpath('<path>/pvalue_weighting_matlab/Code/External Helper Code')
```

The second option is to add those line to scripts that call functions in this package. The third option is to execute the setpaths.m script every time you start a new session with the EIGENEDGE package. This will include the current folder in the Matlab search path.

An example computation is in the \Experiments\Examples\example.m file. This is described in Section 3.

This file is the main documentation for the package. To start, look at the example (Section 3) or at the methods implemented (Sections 4).

3 Example

Here we give a simple example of using the Atomic function for computing limit spectra of sample covariance matrices. We refer to the paper Dobriban (2015) for a full description of the method.

3.1 Background

We begin with the problem statement (borrowed from our paper). Consider a large $n \times p$ matrix **X**, whose rows x^i are independent random vectors. Suppose that x^i are mean zero, and their covariance matrix is the $p \times p$ matrix $\Sigma = \mathbb{E} x^i x^{i\top}$. To estimate Σ , we form the sample covariance matrix

$$\widehat{\Sigma} = \frac{1}{n} \mathbf{X}^{\top} \mathbf{X}.$$

If n and p are of comparable size, then $\widehat{\Sigma}$ deviates substantially from the true covariance. The asymptotic theory of random matrices describes the behavior of the eigenvalues of $\widehat{\Sigma}$ as n,p grow large proportionally (e.g. Bai and Silverstein, 2009). It is well known that, if the distribution of the eigenvalues of Σ tends to a limit population spectral distribution (SD) H, as $n,p\to\infty$ and the aspect ratio $p/n\to\gamma$, then the random eigenvalue distribution of $\widehat{\Sigma}$ also tends to a deterministic limit empirical spectrum F (Marchenko and Pastur, 1967; Silverstein, 1995). This limit differs from the true spectrum.

3.2 Quick example

The Atomic method is designed to compute the limit sample spectrum F from the limit population spectrum H. It is easy to use: in its simplest form, we need to provide a vector of population eigenvalues t (in this case H is a uniform mixture of point masses at t_i), and an aspect ratio gamma, and call atomic(t,gamma). Atomic will return a grid and pointwise density estimates of the limit spectrum F.

The code below (excerpted from example.m) shows an example, in which the eigenvalues are placed at t = [1; 5], and the aspect ratio gamma = 1/2:

```
t = [1; 5]; %location of population eigenvalues: H = 1/n\sum delta_{t_i}
gamma = 1/2; %aspect ratio gamma = p/n
[grid, density] = atomic(t, gamma); %compute limit spectrum
```

The density can be plotted with

```
figure, plot(grid, density,'r','LineWidth',4) %plot
xlabel('Eigenvalue')
ylabel('Density');
```

The results are shown below:

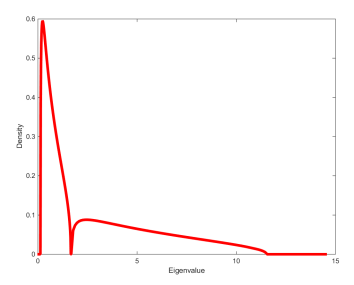


Figure 1: Density of the limit spectrum

3.3 Longer example

4 Computational methods

4.1 Atomic: computing the limit spectrum

The Atomic method computes the limit sample spectrum of covariance matrices from the limit population spectrum. The main user interface for this computation is atomic(). This function takes as input the population spectral distribution H, aspect ratio γ , and other parameters, and computes the density of the limit spectrum on a grid.

The basic syntax of this function is:

In the basic setup, it requires only two inputs:

- t: a vector of positive entries, the locations of the population eigenvalues (length J)
- gamma: aspect ratio $\gamma = p/n$

In the basic case the population spectrum is defined as an equal mixture of point masses at t_i : $H = J^{-1} \sum_{i=1}^{J} \delta_{t_i}$.

The outputs are:

- grid: a positive grid where the density is computed
- density: pointwise numerical approximations to the density on the grid

The complete syntax of the function is:

[grid, density,m, v, mass_at_0, K_hat, l_hat, u_hat, x, f_hat] = atomic(t,gamma,varargin)

Here, in addition to the above two arguments t,gamma, a much larger set of input arguments can be specified within varargin. A complete call, including all parameters, looks like:

The extra arguments w, r, w_int, epsilon, M are optional. You can specify any subset of them, however, you must always use them in this order. If you wish to specify some later arguments, and leave some earlier ones default, then place an empty array [] in their place. For instance, to specify epsilon = 1e-6 without specifying w etc, call atomic(t,gamma,[],[],[],1e-6).

The inputs specify the population spectrum H as a mixture of point masses and uniform distributions:

$$H = \sum_{i=1}^{J} w_i \delta_{t_i} + \sum_{t=1}^{T} w_t^* U(a_t, b_t).$$

Here δ_t is the point mass at t and U(a,b) is the uniform distribution on [a,b]. The parameters of the mixture can be set using the parameters of atomic(), which are, in order:

- w: mixture weights of the components of t in the input spectrum. This must be a positive vector of size J. The default is a uniform vector of length J: w = (1, 1, ..., 1)/J. If a continuous part of the component is also specified with \mathbf{r} , then this parameter must not be empty. The reason is that these weights and \mathbf{w}_{int} implictly depend on each other, as they must sum to 1.
- r: intervals where the continuous part of the spectrum is supported. A real matrix of size $J \times 2$, the t-th row encodes the endpoints $[a_t, b_t]$ of the i-th continuous component of the spectrum, a uniform distribution on $[a_t, b_t]$.
- w_int: weights of the continuous part of the spectrum. This must be a non-negative vector. The entry w_int[t] specifies the weights of the *i*-th continuous component of the spectrum, a uniform distribution on $[a_t, b_t]$. If this is empty, then the defualt value is a uniform distribution subject to the constraint sum(w)sum(w_int) =1+.
- epsilon: accuracy parameter. As the accuracy parameter $\varepsilon \to 0$, the density approximation becomes more accurate. However, only the convergence of the procedure is guaranteed, not the actual accuracy.
- M: number of grid points in each interval of the support. This is an additional parameter which allows the user to specify how many grid points should be used in each interval in the support of the sample spectrum. This is available for additional flexibility.
- alg: The algorithm used to compute the density. The available choices are {'atomic', 'fp', 'Newton'};. These stand for: 'atomic' is the Atomic algorithm proposed in Dobriban (2015), 'fp' is the fixed point method described in Couillet et al. (2011), and 'Newton' is Newton's method, which the author learned from Jack W. Silverstein, and which is described in Dobriban (2015). Please see that paper for more information on each method.

- 4.2 Moments and Quantiles of the ESD
- 4.3 Utilities
- 4.3.1 Arithmetic and geometric models for eigenvalues

References

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