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Bayesian Wavelet Estimation of Long Memory Parameter

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A Bayesian wavelet estimation method for estimating parameters of a stationary I(d) process is represented as an useful alternative to the existing frequentist wavelet estimation methods. The effectiveness of the proposed method is demonstrated through Monte Carlo simulations. The sampling from the posterior distribution is through the Markov Chain Monte Carlo (MCMC) easily implemented in the WinBUGS software package.

Key words: Bayesian method, wavelet, discrete wavelet transform (DWT), I(d) process, long memory

Introduction

Stationary processes exhibiting long range dependence have been widely studied now since the works of Granger and Joyeux (1980) and Hosking (1981). The long range dependence has found applications in many areas, including economics, finance, geosciences, hydrology, and statistics. The estimation of the long-memory parameter of the stationary long-memory process is one of the important tasks in studying this process.

There exist parametric, non-parametric and semi-parametric methods of estimation for the long-memory parameter in literature. In the parametric method, the long-memory parameter is one of the several parameters that determine the parametric model; hence the usual classical methods such as the maximum likelihood estimation can be applied. The non-parametric method, not assuming restricted parametric form of the model, usually uses regression methods by regressing the logarithm of some sampling statistics for estimation.

The widely and often used Geweke and Poter-Hudak (1983) estimation method belongs to non-parametric methods. The semi-parametric method makes intermediate assumptions by not specifying the covariance structure at short ranges. The article by Bardet et al. (2003) surveyed some semi-parametric estimation methods and compared their finite sample performance by Monte-Carlo simulation.

Wavelet has now been widely used in statistics, especially in time series, as a powerful multiresolution analysis tool since 1990's. See Vidakovic (1999) for reference from the statistical perspective. The wavelet's strength rests in its ability to localize a process in both time and frequency scale simultaneously.

This article presents a Bayesian Wavelet estimation method of the long-memory parameter d and variance σ^2 of a stationary long-memory I(d) process implemented in the MATLAB computing environment and the WinBUGS software package.

Methodology

A time series $\{X_t\}$ is a fractionally integrated process, $I(d)$, if it follows:

$$(1-L)^d X_t = \varepsilon_t,$$

where $\varepsilon_t \sim i.i.d. N(0, \sigma_\varepsilon^2)$ and L is the lag operator defined by $LX_t = X_{t-1}$. The parameter d is not necessarily an integer so that fractional differencing is allowed. The process $\{X_t\}$ is

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stationary if $|d| < 0.5$. The fractionally differencing operator $(1-L)^d$ is defined by the general binomial expansion:

$$(1-L)^d = \sum_{k=0}^{\infty} \binom{d}{k} (-L)^k,$$

where

$$\binom{d}{k} = \frac{\Gamma(d+1)}{\Gamma(k+1)\Gamma(d-k+1)}$$

and $\Gamma(\cdot)$ is the usual Gamma function.

Denote the autocovariance function of $\{X_t\}$ as $\gamma(k)$, that is $\gamma(k) = E(X_t X_{t-s})$ where $k=|t-s|$. The formula for $\gamma(k)$ of a stationary $I(d)$ process is well-known (Beran 1994, pp. 63):

$$\begin{aligned} \gamma(0) &= \sigma_\varepsilon^2 \Gamma(1-2d) / \Gamma^2(1-d), \\ \gamma(k+1) &= \gamma(k)(k+d)/(k+1-d), \\ &k = 0, 1, 2, \dots \end{aligned}$$

When $0 < d < 0.5$, the $\gamma(k)$ has a slow hyperbolic decay, hence the process $\{X_t\}$ is a long-memory process.

The fractional difference parameter d and the nuisance parameter σ^2 are usually unknown in an $I(d)$ process. They need to be estimated from the observed time series X_t , $t=1, \dots, N$.

Assume $N=2^J$ for some positive integer J in order to apply the fast algorithm of the Discrete Wavelet Transform (DWT) on $X = (X_t)_{t=1}^N$. Let $\omega = WX$ denote the DWT of X , where $\omega = (c_{j_0}^T, d_{j_0}^T, d_{j_0+1}^T, d_{j_0-1}^T)^T$. The j_0 is the lowest resolution level for which we use $j_0=0$ in this article. The smoothed wavelet coefficient vector $c_{j_0} = (c_{j_0,0}, c_{j_0,1}, \dots, c_{j_0,2^{j_0}-1})^T$. At the resolution level j , the detailed wavelet coefficient vector $d_j = (d_{j,0}, d_{j,1}, \dots, d_{j,2^j-1})^T$ for $j=0, 1, \dots, J-1$.

McCoy and Walden (1996) argued heuristically that the DWT coefficients of X has the following distribution:

$$d_{j,k} \sim N(0, \sigma_j^2),$$

where $j = 0, 1, \dots, J-1$; $k = 0, 1, \dots, 2^j - 1$,

$c_{0,0} \sim N(0, \sigma_{-1}^2)$, and the $d_{j,k}$'s and $c_{0,0}$ are approximately uncorrelated due to the whitening property of the DWT. The σ_j^2 , $j=-1, 0, 1, \dots, J-1$ depend on d and σ_ε^2 as

$$\sigma_j^2 = 2^{J-j} \times 2 \times 4^{-d} \sigma_\varepsilon^2 \int_{2^{-(J-j+1)}}^{2^{-(J-j)}} \sin^{-2d}(\pi f) df.$$

When $J-j \geq 2$, $f < 2^{-2}$, then $\sin(\pi f) \approx \pi f$, so that σ_j^2 can be simplified as (see equation (2.10) of McCoy and Walden 1996)

$$\sigma_j^2 = (2\pi)^{-2d} \sigma_\varepsilon^2 2^{2(J-j)d} (2-2^{2d}) / (1-2d) \quad (1)$$

where $j = -1, 0, \dots, J-2$.

McCoy and Walden (1996) used these facts to estimate d and σ_ε^2 by the Maximum Likelihood Method. They demonstrated through simulation that d could be estimated as well, or better by wavelet methods than the best Fourier-based method.

Jensen (1999) derived the similar result about the distribution of the wavelet coefficients, and by the fact that $Var(d_{j,k}) \propto 2^{-2jd}$, he used the Ordinary Least Squares method to estimate d . That is, by regressing \log of the sample variance of the wavelet coefficients at resolution level j , against $\log(2^{-2j})$ for $j=2, 3, \dots, J-2$, he obtained the OLS estimate of d . The sample variance of the wavelet coefficients at resolution level j is estimated by the sample second moment of the observed wavelet coefficients at resolution level j .

Vannucci and Corradi (1999) section 5 proposed a Bayesian approach. They used independent priors and assumed Inverse Gamma distribution for σ_ε^2 and a Beta distribution for $2d$. They did not use formula (1), instead, they used a recursive algorithm to compute the variances of wavelet coefficients. The posterior inference is done through Markov chain Monte Carlo (MCMC) sampling procedure. They did

not give details of the implementation in the paper.

McCoy and Walden (1996) did not give the variance of their estimates. Jensen (1999) only estimated d using the OLS method, it is not clear how σ_ϵ^2 is estimated. In both cases, the estimated d can not be guaranteed in the range $(-0.5, 0.5)$.

Here, we propose a Bayesian approach to estimate d and σ_ϵ^2 in the same spirit of Vannucci and Corradi (1999) section 5. The distinction of this article from Vannucci and Corradi (1999) is that firstly, we use the explicit formula (1) for the variances of wavelet coefficients at resolution level j instead the recursive algorithm to compute these variances; secondly, the MCMC is implemented in the WinBUGS software package.

Denoting $\theta = (d, \sigma_\epsilon^2)$, the parameters of the models for the data ω . If a prior distribution of $\pi(\cdot)$ of θ is chosen, i.e., $\theta \sim \pi(\theta)$, then by Bayesian formula, the posterior distribution of θ is

$$\pi(\theta | \omega) \propto f(\omega | \theta)\pi(\theta)$$

where $f(\omega | \theta)$ is the likelihood of the data ω given the parameters θ , which is the density of the multivariate normal distribution $N(0, \Sigma)$ with

$$\Sigma = \text{diag}(\sigma_{-1}^2, \Sigma_0, \Sigma_1, \dots, \Sigma_{J-1})$$

and $\Sigma = \text{diag}(\sigma_j^2, \dots, \sigma_j^2)$

for $j = 0, 1, \dots, J - 1$ is a $2^j \times 2^j$ diagonal matrix.

The inference of θ is based on the posterior distribution $\pi(\theta | \omega)$. The MCMC methods are popular to draw repeated samples from the intractable $\pi(\theta | \omega)$. We focus on the implementation of the Gibbs sampling for estimating d and σ_ϵ^2 in the WinBUGS software. The easy programming in the WinBUGS software provides practitioners an useful and

convenient tool to carry out Bayesian computation for long memory time series data analysis.

The following priors will be used. The first prior is the Jefferys' noninformative prior subject to the constraints of the range of model parameters:

$$\pi(\theta) \propto [J(\theta)]^{1/2} I_{(0,+\infty)}(\sigma_\epsilon^2) I_{(-0.5,0.5)}(d),$$

where $I(\cdot)$ is an indicator function for the subscripted set and $J(\theta)$ is the Fisher information for θ :

$$J(\theta) = -E \left[\frac{\partial^2 \ln f(\omega | \theta)}{\partial \theta \theta^T} \right].$$

Simple calculation shows that $J(\theta) \propto 1/\sigma_\epsilon^2$. The second prior is the other independent priors on d and σ_ϵ^2 , i.e.,

$$\pi(\theta) = \pi(d)\pi(\sigma_\epsilon^2).$$

The prior for $d+0.5$ is $Beta(\alpha, \beta)$ where $\alpha > 0, \beta > 0$ are the hyperparameters. This prior restricts $|d| < 0.5$, thus imposing stationarity for the time series. When $\alpha = \beta = 1$, the prior is the noninformative uniform prior. When historical information or expert opinion is available, α and β can be selected to reflect this extra information, thus obtaining an informative prior. Hyper priors can also be used on α and β to reflect uncertainties on them, thus forming a hierarchical Bayesian model.

A $Gamma(\alpha_1, \alpha_2)$ prior is chosen for the precision $\tau^2 = 1/\sigma_\epsilon^2$, where $\alpha_1 > 0, \alpha_2 > 0$ are the hyperparameters. When α_1 and α_2 are close to zero, the prior for σ_ϵ^2 is practically equivalent to $\pi(\sigma_\epsilon^2) \propto 1/\sigma_\epsilon^2$, an improper prior. The non-informative prior $\pi(\sigma_\epsilon^2) \propto I_{(0,+\infty)}(\sigma_\epsilon^2)$ can also be chosen.

Simulation

The MCMC sampling is carried out in the WinBUGS software package. WinBUGS is the current windows-based version of the BUGS (Bayesian inference Using Gibbs Sampling), a newly developed, user-friendly and free software package for general-purpose Bayesian computation, Lunn et al. (2000). It is developed by the MRC, Biostatistics Group, Institute of Public Health (www.mrc-bsu.cam.ac.uk/bugs), Cambridge.

In WinBUGS programming, user only needs to specify the full proper data distribution and prior distributions, WinBUGs will then use certain sophisticated sampling methods to sample the posterior distribution.

In this Monte Carlo experiment, we compare the proposed Bayesian approach with the approach in McCoy and Walden (1996) and Jensen (1999). Different values of d , N and different prior distributions $\pi(\theta)$ are used to determine the effectiveness of the estimation procedure. Also used were two different wavelet bases to compare the effect of this choice.

The Davis and Harte (1987) algorithm was used to generate an $I(d)$ process because of its efficiency compared to other computationally intensive methods (McLeod & Hipel (1978)). This algorithm generates a Gaussian time series with the specified autocovariances by discrete Fourier transform and discrete inverse Fourier transform. It is well known that Fast Fourier Transform (FFT) can be carried out in $O(N \log N)$ operations, so the computation is fast.

The generation of the $I(d)$ process using the Davis and Harte algorithm and the DWT of the generated $I(d)$ process are carried out in the MATLAB 6.5 on a Pentium III running Windows 2000. The DWT tool used is the *WAVELAB802* developed by the team from the Statistics Department of Stanford University (<http://www-stat.stanford.edu/~wavelab>).

The following two different wavelet basis for comparison were chosen: (a) Harr wavelet; (b) LA(8): Daubechies least asymmetric compactly supported wavelet basis with four vanish moments, see p.198 of Daubechies (1992).

The periodic boundary handling is used. The data of the discrete wavelet transformed $I(d)$

process is first saved in a file in R data file format. Then WinBUGS1.4 is activated under MATLAB to run a script file that implements the proposed Bayesian estimation procedure. The estimation results from WinBUGS1.4 are then converted to the MATLAB variables for further uses.

The model parameters are estimated under the following independent priors on d and σ_ε^2

$$(a) \quad \begin{aligned} d &\sim Unif(-0.5, 0.5), \\ &\sim Gamma(0.01, 0.01); \end{aligned}$$

$$(b) \quad d \sim Unif(-0.5, 0.5), \sigma_\varepsilon^2 \sim Unif(0, 1000).$$

The prior (a) is practically equivalent to Jefferys' noninformative prior:

$$\pi(d, \sigma_\varepsilon^2) \propto \frac{1}{\sigma_\varepsilon^2} I_{(0, +\infty)}(\sigma_\varepsilon^2) I_{(-0.5, 0.5)}(d).$$

BUGS only allow the use of proper prior specification, so the non-informative or improper prior distribution can be regarded as the limit of a corresponding proper prior.

The estimation results using the proposed Bayesian approach for the simulated $I(d)$ process and the method by Jensen (1999) and McCoy and Walden (1996) are found in Table 1 for Haar wavelets and Table 2 for LA(8) wavelets. For the chosen prior, it reports the estimated posterior mean, posterior standard deviation (SD). In addition, it also tabulated in the parenthesis below the value of Mean and SD the 95% credible intervals of the parameters using the 2.5% and 97.5% quantiles of the random samples.

In all cases, two independent chains of 10500 iterations each were run, keeping every tenth one, after burn-in 500, with random initial values. The posterior inference is based on the actual random samples of 2000. For the case of $N=256$, $d=0.1$, $\sigma_\varepsilon^2=1.0$ and prior (b), Figure 1 shows the trace of the random samples and the kernel estimates of the posterior densities of the parameters.

The autocorrelation function of the random samples shows very little autocorrelations for the drawn series of the random samples. The two parallel chains mix well after small steps of the initial stage. All other diagnostics for convergence indicate a good convergence behavior.

In most cases, the Bayesian wavelet estimates of d and σ_ε^2 are quite good. They are very close to the truth. The 95% credible interval

given by the Bayesian wavelet approach is well centered around the true parameter and is also very tight.

The estimation results using the two different priors (a) and (b) are very similar. The estimates by Jensen's method differ most from those by the other methods. It seems that LA(8) generally gives better estimates than Haar. This is in agreement with the results of McCoy and Walden (1996) section 5.2.

Figure 1: Trace and Kernel Density Plot for d and σ_ε^2 .

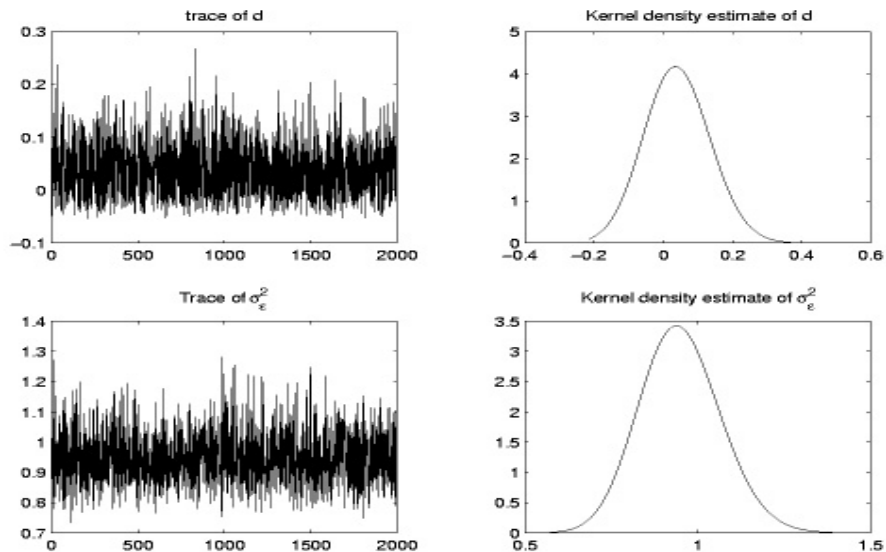


Table 1: Estimation of the simulated $I(d)$ process when $N=256$ Using Haar Basis.

Parameter	Jensen	MW	Prior (a)		Prior (b)	
			Mean	SD	Mean	SD
d=0.1	0.1620	0.1629	0.1686	0.0499	0.1692	0.0499
			(0.0739,	0.2711)	(0.0768,	0.2674)
$\sigma_\varepsilon^2 = 1.0$		1.0226	1.0452	0.0931	1.0485	0.0977
			(0.8801,	1.2460)	(0.8791,	1.2540)
d=0.25	0.1431	0.1858	0.1887	0.0465	0.1880	0.0462
			(0.1049,	0.2827)	0.1008,	0.2854)
$\sigma_\varepsilon^2 = 1.0$		1.0789	1.1000	0.0972	1.1068	0.1021
			(0.9331,	1.3150)	(0.9289,	1.3220)
d=0.4	0.4121	0.4384	0.4301	0.0351	0.4284	0.0364
			(0.3567,	0.4902)	(0.3489,	0.4901)
$\sigma_\varepsilon^2 = 1.0$		1.0189	1.0445	0.0934	1.0571	0.0975
			(0.8775,	1.2395)	(0.8822,	1.2640)
d=0.1	0.1227	0.0681	0.0709	0.0470	0.0719	0.0472
			(-0.0176,	0.1663)	(-0.0172,	0.1679)
$\sigma_\varepsilon^2 = 2.0$		2.1482	2.1787	0.1918	2.1943	0.1948
			(1.8455,	2.5745)	(1.8570,	2.5975)
d=0.25	0.2468	0.1855	0.1858	0.0477	0.1847	0.0462
			(0.0995,	0.2858)	(0.0938,	0.2785)
$\sigma_\varepsilon^2 = 2.0$		1.9369	1.9674	0.1729	1.9791	0.1770
			(1.6570,	2.3275)	(1.6715,	2.3675)
d=0.4	0.2154	0.3096	0.3127	0.0467	0.3105	0.0476
			(0.2238,	0.4069)	(0.2165,	0.4079)
$\sigma_\varepsilon^2 = 2.0$		1.7783	1.8130	0.1540	1.8305	0.1619
			(1.5435,	2.1385)	(1.5300,	2.1665)

Table 2: Estimation of the simulated $I(d)$ process when $N=256$ Using LA(8) Basis.

Parameter	Jensen	MW	Prior (a)		Prior (b)	
			Mean	SD	Mean	SD
d=0.1	0.0759	0.1701	0.1757	0.0466	0.1755	0.0446
			(0.0894,	0.2734)	(0.0936,	0.2707)
$\sigma_\varepsilon^2 = 1.0$		1.0037	1.0222	0.0935	1.0270	0.0899
			(0.8529,	1.2295)	(0.8626,	1.2190)
d=0.25	0.0904	0.2611	0.2651	0.0508	0.2661	0.0502
			(0.1681,	0.3680)	(0.1705,	0.3741)
$\sigma_\varepsilon^2 = 1.0$		1.0154	1.0398	0.0916	1.0412	0.0888
			(0.8791,	1.2295)	(0.8824,	1.2255)
d=0.4	0.4906	0.4369	0.4304	0.0359	0.4295	0.0362
			(0.3548,	0.4905)	(0.3536,	0.4895)
$\sigma_\varepsilon^2 = 1.0$		1.0148	1.0413	0.0953	1.0502	0.0932
			(0.8669,	1.2370)	(0.8826,	1.2450)
d=0.1	0.0542	0.1110	0.1175	0.0529	0.1151	0.0535
			(0.0166,	0.2278)	(0.0183,	0.2298)
$\sigma_\varepsilon^2 = 2.0$		2.1233	2.1594	0.1926	2.1694	0.1894
			(1.8185,	2.5765)	(1.8235,	2.5650)
d=0.25	0.1977	0.2609	0.2608	0.0535	0.2637	0.0540
			(0.1556,	0.3697)	(0.1630,	0.3761)
$\sigma_\varepsilon^2 = 2.0$		1.8372	1.8745	0.1609	1.8849	0.1709
			(1.5870,	2.2165)	(1.5795,	2.2420)
d=0.4	0.2632	0.3111	0.3130	0.0454	0.3117	0.0463
			(0.2257,	0.4045)	(0.2236,	0.4040)
$\sigma_\varepsilon^2 = 2.0$		1.7469	1.7942	0.1635	1.7995	0.1595
			(1.5080,	2.1510)	(1.5145,	2.1225)

Figure 2: Box plots of the estimates for $N=128$.

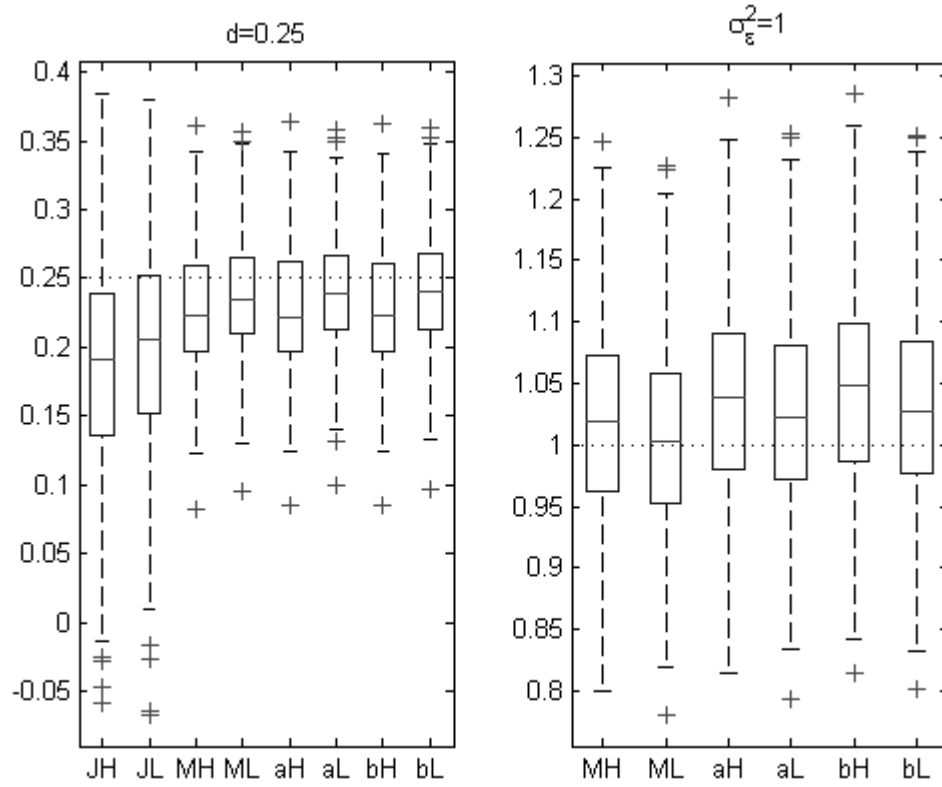
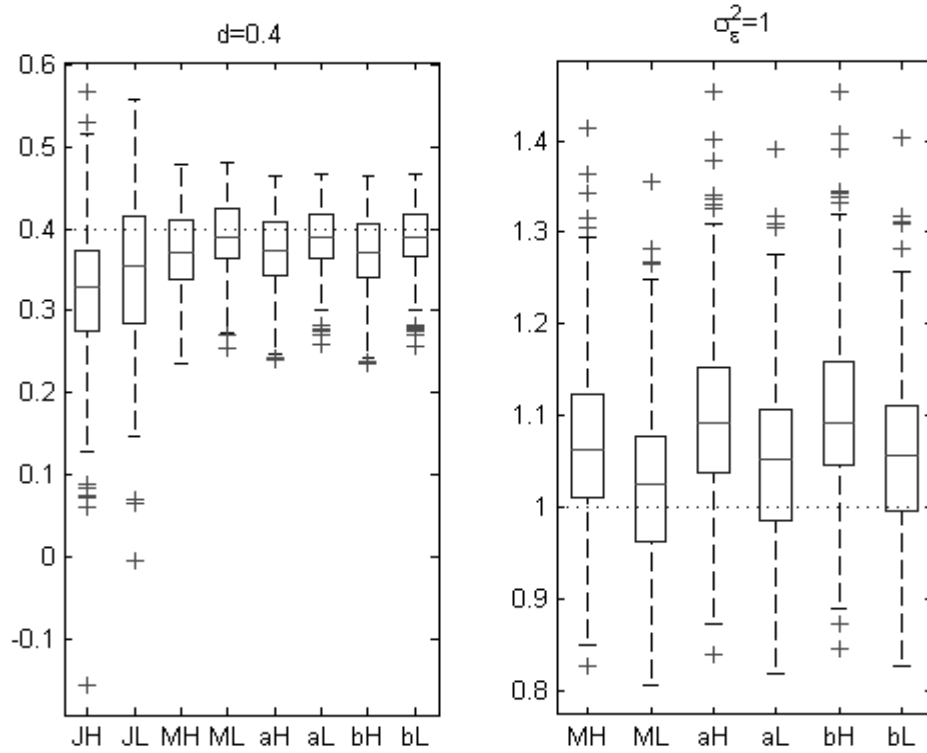


Figure 3: Box plots of the estimates for $N=128$.



Frequentist Comparison

Also compared were the estimates of the three methods in repeatedly simulated I(d) process. Figure 2 is the box plots of the estimates for d and σ_ε^2 respectively of 200 replicates with $N=128$, $d=0.25$ and $\sigma_\varepsilon^2=1.0$. Figure 3 is the box plots of the estimates for 200 replicates with $N=128$, $d=0.40$ and $\sigma_\varepsilon^2=1.0$. The x -axis labels in the box plot read as follows: 'JH' denotes the case by the Jensen method using Haar; 'JL' denotes the case by the Jensen method using LA(8); and so forth. Because of the long computation time associated with the Gibbs sampling for the large number of simulated I(d) processes, we limit the burn-in to 100 iterations and the number of random samples to 500. Because only the posterior mean was calculated using the generated random samples, not much information was lost even when the slightly short chain was used.

For the estimates of d , the mean square errors of the McCoy and Walden and The Bayesian method using these two priors are very similar, and they are all smaller than the one by Jensen's OLS. LA(8) gives less biased estimates than Haar. The mean estimates for d given by the Bayesian method using LA(8) is similar to those by McCoy and Walden. In all methods, it seems the estimates for d and σ_ε^2 are a little biased in that \hat{d} tends to underestimate d and $\hat{\sigma}_\varepsilon^2$ tends to overestimate σ_ε^2 .

Conclusion

Bayesian wavelet estimation method for the stationary I(d) process provides an alternative to the existing frequentist wavelet estimation methods. Its effectiveness is demonstrated through Monte Carlo simulations implemented in the WinBUGS computing package.

A future effort is to extend the Bayesian wavelet method to more general fractional process such as $ARFIMA(p,d,q)$. The hypothesis testing problem for the I(d) process can also be explored via the Bayesian wavelet approach.

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Appendix:

This appendix includes the MATLAB code for first simulating the $I(d)$ process, then transforming it by DWT, and the WinBUGS program for the MCMC computation. In the WinBUGS programming, the symbol “~” is for the stochastic node which has the specified distribution denoted on the right side, the symbol “←” is for the deterministic node which has the specified expression denoted on the right side. All the likelihood function, the prior distributions and initial values of the nodes without parents must be specified in the programs.

The MATLAB code:

```
function x=Generatex(J, d, sig2eps)
%Generate the I(d) process
%input:
%J: where N=2^J sample size
%d: long memory parameter of the I(d) process, abs(d)<0.5
%sig2eps:  $\sigma^2 \epsilon^2$ 

%output:
%x: the time series

N=2^J;
c=[];
% generate the autocovariance function by the formular of covariance
% function for LRD
c(1)=sig2eps*gamma(1-2*d)/((gamma(1-d))^2);
%for i=1:N-1 c(i+1)=c(1)*gamma(i+d)*gamma(1-d)/(gamma(d)*gamma(i+1-d)); end;
for i=1:N-1 c(i+1)=c(i)*(i+d-1)/(i-d); end;

x=GlrddDH(c);

function x=GlrddDH(c);
%GlrddDH.m Generating the stationary gaussian time series with specified
% autocovariance series c
% using Davis and Harte's method, Appendix of 'Tests for Hurst Effect',
% Biometrika, V74, No. 1 (Mar., 1987), 95-101

%c: autocovariance series

[temp, N]=size(c); %c is a row vector

cCirculant=[];
for i=1:N-2 cCirculant(i)=c(N-i); end;

cFull=[];
cFull=[c cCirculant];
```

```

g=[];
g=fft(cFull);
%Fast Fourier Transform of cFull

Z=[];
Z=complex(normrnd(0,1,1,N), normrnd(0,1, 1,N));
Z(1)=normrnd(0,sqrt(2)); %Be careful to specify sqrt(2), if you want variance of Z(1) to be 2
Z(N)=normrnd(0,sqrt(2));

ZCirculant=[];
for i=1:N-2 ZCirculant(i)=conj(Z(N-i)); end;

ZFull=[];
ZFull=[Z ZCirculant];

X=[];
X=ifft(ZFull.*sqrt(g)).*sqrt(N-1);

x=[];
x=real(X(1:N));

function [dJensen, dMW, sigMW, dBS, sigBS]=GetdHatSig2Hat(x, j0, filter)
%Wavelet estimation of Long Range Dependence parameters
%
%input:
%x: the observed I(d) process
%j0: lowest resolution level of the DWT
%filter: wavelet filter

%output:
%dJensen: estimate of d by Jensen 1999
%dMW: estimate of d by McCoy & Walden 1996
%sigMW: estimate of  $\sigma_\epsilon^2$  by McCoy & Walden 1996
%dBS: estimate of d by Bayesian Wavelet Method for prior (a), (b)
%dBS.a, dBS.b
%sigBS: estimate of  $\sigma_\epsilon^2$  by Bayesian Wavelet Method for prior (a), (b)
%sigBS.a, sigBS.b

N=length(x);
J=log2(N);

w=[];

w = FWT_PO(x,j0,filter)'; %w is a coulumn vector

resolution=[]; % data used in WinBUGS14
resolution(1:2^j0,1)=j0-1;
for j = j0:(J-1)
    resolution(2^j+1 : 2^(j+1),1)=j;
end;

vwj=[];
for j=j0+1:(J-1)
    vwj(j, :)=j, mean(w(dyad(j)).^2)];

```

```

end;

tempd=[];
tempd=-[ones(J-2,1), log(2.^(2*vwj(2:J-1,1)))]\log(vwj(2:J-1,2));
dJensen=tempd(2);

OPTIONS=optimset(@fminbnd);
dMW=fminbnd(@NcllhMW, -0.5, 0.5, OPTIONS, j0, w, J);
sigMW=findSig2epsHat(dMW, j0, w, J);

n=N-2^(J-1);    %the first n data of w, approximation of variance

%function mat2bugs() converts matlab variable to BUGS data file
mat2bugs('c:\WorkDir\LRD_data.txt', 'w',w,'twopowl', 2^j0, 'n', n, 'N', N,
        'resolution', resolution, 'J', J, 'pi', pi,'K',500);

%set the current directory at MATLAB to 'C:\Program Files\WinBUGS14\'
cd 'C:\Program Files\WinBUGS14\';

%prior (a)
dos('WinBUGS14 /par BWIdSt_a.odc');
Sa=bugs2mat('C:\WorkDir\bugsIndex.txt', 'C:\WorkDir\bugs1.txt');

dBS.a=mean(Sa.d);    %the posterior mean as the estimate of d
sigBS.a=mean(Sa.sig2eps);    %the posterior mean as the estimate of sig2eps

%prior (b)
dos('WinBUGS14 /par BWIdSt_b.odc');
Sb=bugs2mat('C:\WorkDir\bugsIndex.txt', 'C:\WorkDir\bugs1.txt');

dBS.b=mean(Sb.d);    %the posterior mean as the estimate of d
sigBS.b=mean(Sb.sig2eps);    %the posterior mean as the estimate of sig2eps

cd 'C:\WorkDir';

function y=NcllhMW(d, j0, w, J);
%NcllhMW.m --- Negative Concentrated log likelihood of McCoy & Walden
%
%input:
%d: the long memory parameter, a value in (0,0.5)
%j0: Lowest Resolution Level
%w: w=Wx, x is the observed time series
%J: N=2^J sample size
%
%output:
%y: Negative Concentrated log likelihood for the given data w

```

```

m=J-j;
bmP(j+1)=2*4^(-d)*quad(@sinf,2^(-m-1),2^(-m),[],[],d);
%by McCoy & Walden's formula, P37, (2.9)
smP(j+1)=2^m*bmP(j+1);
end;

bpp1P=gamma(1-2*d)/((gamma(1-d))^2)-sum(bmP);
%B_{p+1} in McCoy & Walden's notation, p=J here
spp1P=2^J*bpp1P*(bpp1P>0);
%S_{p+1} in McCoy & Walden's notation, it should be nonnegative

if spp1P>0
    sig2epsHat=w(1)^2/spp1P;
else
    sig2epsHat=0;
end;
sumlogsmP=0;
for j = j0:(J-1)
    sig2epsHat=sig2epsHat+sum(w(2^j+1 : 2^(j+1)).^2)/smP(j+1);
    sumlogsmP=sumlogsmP+2^j*log(smP(j+1));
end;
sig2epsHat=sig2epsHat/N;
%McCoy & Walden, Page 49, formular (5.1)

y=N*log(sig2epsHat)+log(spp1P)+sumlogsmP;
%McCoy & Walden, Page 49

function sig2epsHat=findSig2epsHat(d, j0, w, J);
%find Sig2epsHat by McCoy & Walden Page 49, formular (5.1)
%
%input:
%d: the long memory parameter, a value calculated by function NcllhMW();
%j0: Lowest Resolution Level
%J: N=2^J sample size

N=2^J;

bmP=[];
smP=[];
for j=j0:(J-1) %j is the resolution level
    m=J-j;
    bmP(j+1)=2*4^(-d)*quad(@sinf,2^(-m-1),2^(-m),[],[],d);
    %by McCoy & Walden's formula, P37, (2.9)
    smP(j+1)=2^m*bmP(j+1);
end;

bpp1P=gamma(1-2*d)/((gamma(1-d))^2)-sum(bmP);
%B_{p+1} in McCoy & Walden's notation, p=J here
spp1P=2^J*bpp1P*(bpp1P>0);
%S_{p+1} in McCoy & Walden's notation, it should be nonnegative

```

```

N=2^J;
bmP=[];
smP=[];
for j=j0:(J-1) %j is the resolution level

if spp1P>0
  sig2epsHat=w(1)^2/spp1P;
else
  sig2epsHat=0;
end;

for j = j0:(J-1)
  sig2epsHat=sig2epsHat+sum(w(2^j+1 : 2^(j+1)).^2)/smP(j+1);
end;

sig2epsHat=sig2epsHat/N;
\end{verbatim}

```

The WinBUGS script file: BWIdSt_a.odc

```

check('C:/MyDir/LRD_model_a.odc')
data('C:/MyDir/LRD_data.txt')
compile(1)
gen.inits()
update(100)
set(d)
set(sig2eps)
update(500)
coda(*, 'C:/Documents and Settings/MyDir/bugs')
#save('C:/Documents and Settings/MyDirlog.txt')
quit()

```

The WinBUGS model file: LRD_model_a.odc

```

model {
# This takes care of the father wavelet coefficients from level L+1 to J-1
# which are detailed wavelet coefficients, $D$
for (i in twopowl+1:n) {
  tau[i]<-1/(pow(2*pi, -2*d)*sig2eps*pow(2, 2*d*(J-resolution[i]))*(2-pow(2,2*d))/(1-2*d))
  w[i] ~ dnorm (0, tau[i])
}
}

```

```

#The following takes care the wavelet coefficients at the resolution level J-1.
#It uses the exact formula instead of the approximation.
for (i in 1:K) { sinf[i]<-pow(sin(pi*(0.25+i/(4*K))),-2*d)}
integration<-sum(sinf[])/(4*K)
B1<-2*pow(4,-d)*sig2eps*integration

tau1<-1/(2*B1)      #S_1=2*B_1 in McCoy & Walden 1996's notation

for (i in (n+1): N) {
  w[i] ~ dnorm(0, tau1)
}

# This takes care of the scaling coefficients on the lowest level $j_0=L$
# which are mother wavelet coefficients, $C$

# twopow1 <- pow(2, L)

for (jp1 in 1:J-1) {      #jp1=j+1, m=J-j
  b[jp1]<-(2*pow(2*pi, -2*d)*sig2eps*pow(2, -(J-jp1+1)*(1-2*d)) *(1-pow(2,2*d-1))/(1-2*d))
}
  bpp1<-sig2eps*exp(loggam(1-2*d))/pow(exp(loggam(1-d)),2)-sum(b[])-B1;
  #B_{p+1} in McCoy & Walden's notation
  spp1<-pow(2,J)*bpp1*step(bpp1)+1.0E-6;
  #S_{p+1} in McCoy & Walden's notation, this should be positive

  tau0 <- 1/spp1
  for (i in 1: twopow1) {
    w[i] ~ dnorm(0, tau0)
  }

#note: m=J-resolution[i] in McCoy & Walden's 1996 paper

# prior (a)
d~dunif(-0.5, 0.5)
sig2eps<-1/ tau2
tau2~dgamma(1.0E-2,1.0E-2)

#prior (b)
# d~dunif(-0.5, 0.5)
# sig2eps~dunif(0,1000)
}

```