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Risk Management, Signal Processing and Econometrics: A New Tool for Forecasting the Risk of Disease Outbreaks

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Abstract

This paper takes a novel approach for forecasting the risk of disease emergence by combining risk management, signal processing and econometrics to develop a new forecasting approach. We propose quantifying risk using the Value at Risk criterion and then propose a two staged model based on Multivariate Singular Spectrum Analysis and Quantile Regression (MSSA-QR model). The proposed risk measure (PLVaR) and forecasting model (MASS-QR) is used to forecast the worst cases of waterborne disease outbreaks in 22 European and North American countries based on socio-economic and environmental indicators. The results show that the proposed method perfectly forecasts the worst case scenario for less common waterborne diseases whilst the forecasting of more common diseases requires more socio-economic and environmental indicators.

Keywords: Value at Risk; Disease; Outbreaks; Forecasting; Quantile Regression; Multivariate Singular Spectrum Analysis.

1. Introduction

2 The accurate forecasting of disease outbreaks continue to challenge re-
3 searchers, governments and policy makers (Graham et al. , 2018; Metcalf
4 and Lessler , 2018). The task itself is challenging as an outbreak is a result

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5 of interactions between pathogens/parasites, hosts and other environmental
6 variables (Alizon et al. , 2013; Griffiths et al. , 2011).

7 Accordingly, in the recent past, researchers have adopted a variety of tools
8 from different parts of science to forecast disease outbreaks. For instance,
9 Lowe et al. (2017) used precipitation, minimum temperature, and El Niño
10 index forecasts to predict the dengue incidence in Ecuador. Their results
11 show that using climatological forecasts could improve the accuracy of dengue
12 outbreak forecast. Han and Drake (2016) proposed using statistical machine
13 learning methods to forecast the outbreaks of a disease. They argued that
14 applying machine learning methods to existing big data on environmental,
15 epidemiological and molecular systems could help public health authorities
16 to predict the flow or risks of disease emergence (including outbreak risks).
17 Liao et al. (2017) used a Bayesian Belief Network (BBN) to predict the risk
18 of further outbreaks. They suggest that the BBN technique can be used for
19 early warnings of infectious diseases.

20 Although many of the methods considered in disease outbreak risk fore-
21 casting proved to be accurate and effective, most of the research forecasts the
22 number of cases/incidence, ratios or the probability of occurrence as outbreak
23 risks. On the other hand, in risk management, one is usually interested in
24 worst case scenarios. For instance, in financial risk analysis, instead of fore-
25 casting the average value of an asset, it is common to forecast the value which
26 is the lowest with 95% confidence. Such values are referred to as Value at
27 Risk (Davino et al., 2014) and shows the value of the asset in in extremely
28 negative conditions (the probability of extreme events taking place is 5%).

29 In this paper, we are concerned with forecasting the worst case scenarios
30 for disease outbreaks. Relying on financial risk analysis, a new risk measure
31 is proposed to present the worst case scenario. More specifically, a model
32 based on the Multivariate Singular Spectrum Analysis (Sanei and Hassani,
33 2015) and the Quantile Regression (Koenker , 2005) is developed to forecast
34 the disease outbreak worst case scenario. The proposed method is used to
35 forecast annual outbreaks of 13 waterborne disease in 22 European and North
36 American countries between 2011 and 2015. The data from 10 socio-economic
37 and environmental indicators between 1998 and 2010 is used to estimate the
38 coefficients of the model (train the model). Results show that with relatively
39 small number of indicators and training data, the proposed model has the

40 ability to forecast the worst cases of outbreaks for less common waterborne
 41 diseases. For more common waterborne disease like Diarrhoea, Pertussis and
 42 Malaria, however, more indicators are needed.

43 The remainder of the paper is organised as follows. The proposed forecast-
 44 ing method is presented in Section 2. Section 3 gives a complete description
 45 of the waterborne disease dataset and indicators used to forecast the disease
 46 outbreaks. The results from the forecasting exercise for waterborne disease
 47 outbreaks are presented in Section 3. Finally, Section 4 concludes the paper.

48 2. Methodology

49 2.1. Value at Risk and Population Loss Value at Risk

50 The Value at Risk (VaR) (Leavens , 1945) is one of the common risk
 51 measures in financial risk analysis. The VaR measure shows the minimum
 52 value of an asset (or its return) with $1 - \alpha$ confidence level, i.e. the probability
 53 that the value of an asset goes under the VaR is α . In other words, the VaR
 54 shows the scenario which with confidence level $1 - \alpha$ worst that that won't
 55 happen (the risk that cases worst than VaR happens in reality is α). Since in
 56 investment problems, the worst cases are always the lower values (e.g. lower
 57 returns, price, or income) the VaR in risk level α (confidence level $1 - \alpha$) is
 58 defined as follows:

$$VaR_\alpha(Y) = \inf\{y \in \mathbb{R} : F_Y(y) = \alpha\}$$

59 where Y is the value (return, price, ...) of the financial asset. The VaR_α is
 60 the α th quantile of the value distribution ($F_Y(y)$), It shows the value of an
 61 asset in risk situations which means with $1 - \alpha$ confidence the VaR_α is the
 62 worst case scenario (for more details on VaR see McNeil et al., 2005).

63 Adopting the VaR concept from finance, we define the Population Loss
 64 Value at Risk (PLVaR), as the worst case scenario in disease outbreak with
 65 risk level α :

$$PLVaR_\alpha(Y) = \inf\{y \in \mathbb{R} : F_Y(y) = 1 - \alpha\}, \quad (1)$$

66 where Y is the number (or ratio) of losses in disease outbreak. Unlike
 67 $VaR_\alpha(Y)$, the $PLVaR_\alpha(Y)$ is the $(1 - \alpha)$ th quantile of the Y , since the
 68 worst case in disease outbreak is the case with largest number (ratio) of

69 losses. In this manner, the $PLVaR_\alpha$ shows the worst case scenario in disease
70 outbreak, with $1 - \alpha$ confidence level.

71 The $PLVaR$ can be used as a risk measure in disease control and out-
72 break prevention planes. The $PLVaR$ has the ability to forecast the disease
73 outbreaks along with the size of the break out. Non-zero Values of $PLVaR$
74 show the outbreak situations, while the larger values show the estimate the
75 larger outbreaks. For instance, the $PLVaR_{0.01} = 0$ means in 0.99 confidence
76 level, there is not a disease outbreak (in other words, it means the chance of
77 disease outbreak is under 1%). Using $PLVaR$ as a risk measure, one may
78 forecast the future values of $PLVaR_\alpha$ in order to forecast the size of the
79 future outbreaks.

80 2.2. Multivariate Singular Spectrum Analysis

81 The Horizontal MSSA Recurrent (HMSSA-R) forecasting algorithm uses
82 following steps to forecast multivariate time series. Those interested in an
83 in-depth explanation of the theory underlying MSSA are directed to Sanei
84 and Hassani (2015). In presenting this algorithm we mainly follow and rely
85 on the notations in Sanei and Hassani (2015).

86 2.2.1. HMSSA-R Optimal Forecasting Algorithm

- 87 1. Consider M time series with identical series lengths of N_i , such that
88 $Y_{N_i}^{(i)} = (y_1^{(i)}, \dots, y_{N_i}^{(i)})$ ($i = 1, \dots, M$).
- 89 2. For forecasting exercises we would split each time series into three parts
90 leaving $\frac{2}{3}^{rd}$ for model training and testing, and $\frac{1}{3}^{rd}$ for validation.
- 91 3. Beginning with a fixed value of $L = 2$ ($2 \leq L \leq \frac{N}{2}$) and in the pro-
92 cess, evaluating all possible values of L for Y_{N_i} , using the training data
93 construct the trajectory matrix $\mathbf{X}^{(i)} = [X_1^{(i)}, \dots, X_K^{(i)}] = (x_{mn})_{m,n=1}^{L,K_i}$ for
94 each single series $Y_{N_i}^{(i)}$ ($i = 1, \dots, M$) separately.
- 95 4. Then, construct the block trajectory matrix \mathbf{X}_H as follows:

$$\mathbf{X}_H = \left[\mathbf{X}^{(1)} : \mathbf{X}^{(2)} : \dots : \mathbf{X}^{(M)} \right].$$

- 96 5. Let vector $U_{H_j} = (u_{1j}, \dots, u_{Lj})^T$, with length L , be the j^{th} eigenvector
97 of $\mathbf{X}_H \mathbf{X}_H^T$ which represents the SVD.

- 98 6. Evaluate all possible combinations of r ($1 \leq r \leq L - 1$) step by step
 99 for the selected L and construct $\widehat{\mathbf{X}}_H = \sum_{i=1}^r U_{H_i} U_{H_i}^T \mathbf{X}_H$ as the recon-
 100 structed matrix obtained using r eigentriples:

$$\mathbf{X}_H = \left[\widehat{\mathbf{X}}^{(1)} : \widehat{\mathbf{X}}^{(2)} : \dots : \widehat{\mathbf{X}}^{(M)} \right].$$

- 101 7. Consider matrix $\widetilde{\mathbf{X}}^{(i)} = \mathcal{H}\widehat{\mathbf{X}}^{(i)}$ ($i = 1, \dots, M$) as the result of the
 102 Hankelization procedure of the matrix $\widehat{\mathbf{X}}^{(i)}$ obtained from the previous
 103 step for each possible combination of SSA choices.
 104 8. Let $U_{H_j}^\nabla$ denote the vector of the first $L - 1$ coordinates of the eigenvec-
 105 tors U_{H_j} , and π_{H_j} indicate the last coordinate of the eigenvectors U_{H_j}
 106 ($j = 1, \dots, r$).
 107 9. Define $v^2 = \sum_{j=1}^r \pi_{H_j}^2$.
 108 10. Denote the linear coefficients vector \mathcal{R} as follows:

$$\mathcal{R} = \frac{1}{1 - v^2} \sum_{j=1}^r \pi_{H_j} U_{H_j}^\nabla. \quad (2)$$

- 109 11. If $v^2 < 1$, then the h -step ahead HMSSA forecasts exist and is calcu-
 110 lated by the following formula:

$$\left[\hat{y}_{j_1}^{(1)}, \dots, \hat{y}_{j_M}^{(M)} \right]^T = \begin{cases} \left[\tilde{y}_{j_1}^{(1)}, \dots, \tilde{y}_{j_M}^{(M)} \right], & j_i = 1, \dots, N_i, \\ \mathcal{R}^T \mathbf{Z}_h, & j_i = N_i + 1, \dots, N_i + h, \end{cases} \quad (3)$$

111 where, $\mathbf{Z}_h = \left[Z_h^{(1)}, \dots, Z_h^{(M)} \right]^T$ and $Z_h^{(i)} = \left[\hat{y}_{N_i-L+h+1}^{(i)}, \dots, \hat{y}_{N_i+h-1}^{(i)} \right]$
 112 ($i = 1, \dots, M$).

- 113 12. Seek the combination of L and r which minimises a loss function, \mathcal{L}
 114 and thus represents the optimal HMSSA-R choices for decomposing
 115 and reconstructing in a multivariate framework.
 116 13. Finally use the selected optimal L to decompose the series comprising
 117 of the validation set, and then select r singular values for reconstructing
 118 the less noisy time series, and use this newly reconstructed series for
 119 forecasting the remaining $\frac{1}{3}^{rd}$ observations (or the test set as relevant
 120 to this study).

121 *2.3. Quantile Regression*

122 The Quantile Regression (QR) models the τ th quantile of the response
123 variable using a regression line:

$$Q_\tau = \beta_{0,\tau} + \sum_{i=1}^p \beta_{i,\tau} x_i + \varepsilon_\tau,$$

124 where x_1, \dots, x_p are independent variables and Q_τ is the τ th quantile of re-
125 sponse variable y with cumulative distribution function $F_Y(\cdot)$:

$$Q_\tau = \inf\{y \in \mathbb{R} : F_Y(y) = \tau\}, \quad 0 < \tau < 1.$$

126 The coefficients of the model can be estimated by minimizing the loss function
127 $L_\tau(e) = (\tau - I_{(e < 0)}) e$ where $I_{(e < 0)}$ is the Indicator function (for more details
128 on QR see Davino et al., 2014):

$$I_{(e < 0)} = \begin{cases} 1 & \text{if } e < 0 \\ 0 & \text{otherwise} \end{cases}$$

129 The QR model is a simple tool for risk analysis. For instance, one may
130 use the QR model to estimate the VaR (or PLVaR) for response variable
131 y based on given situation (indicators) x_1, \dots, x_p . On the other hand, one
132 may use the QR model to control the worst case scenario using the control
133 variables x_1, \dots, x_p .

134 *2.4. MSSA-QR model for PLVaR forecasting*

135 In order to forecast the PLVaR, we propose a two stage model. At the
136 first stage, we use MSSA to forecast the indicators in the model. The second
137 stage, uses forecasted values of indicators, to estimate the outbreak risk. It
138 should be noted that in first stage, not all the variables need to be forecasted
139 using MSSA. The future values of some indicators are already forecasted
140 (for instance the population structure and population growth rates for dif-
141 ferent countries are forecasted using Birth/Death models and are available
142 from <http://www.un.org/en/development/desa/population/>). Further-
143 more, some of the indicators are related to governments policies and can be
144 forecasted based on governments announced policies. The MSSA-QR model
145 for PLVaR h step ahead forecasting follows these steps:

146 **First Stage:** Forecasting the indicators

- 147 1. Use data available from the past ($t = 1, \dots, N$) for M countries/regions
 148 and the birth/death models to calculate h step ahead forecast for pop-
 149 ulation indicators (e.g. population structure, growth etc.).
 150 2. Assess the government's announced policies and use data available from
 151 the past ($t = 1, \dots, N$) to forecast the indicators related to govern-
 152 ment's policies (like infrastructural developments) for the desired time
 153 horizon.
 154 3. Use the HMSSA-R algorithm and calculate the h step ahead forecasts
 155 for the rest of the indicators, based on historical data (each indicator is
 156 a M -variate time series where M is the number of countries/regions).

157 **Second Stage:** Forecasting the PLVaR for a given risk level α

- 158 1. Use the data available in time period $t = 1, \dots, N$ and countries/regions
 159 $i = 1, \dots, M$ to fit the QR model as:

$$PLVaR_{\alpha}(Y_{t,i}) = Q_{1-\alpha} = \beta_{0,1-\alpha} + \sum_{j=1}^p \beta_{j,1-\alpha} x_{j,t,i} + \varepsilon_{1-\alpha,t,i},$$

160 where $Y_{t,i}$ is the number (or ratio) of deaths caused by disease outbreak
 161 at time t and country/region i . The $x_{j,t,i}$ is the j th indicator observed
 162 value at time t and country/region i . The $\varepsilon_{\alpha,t,i}$ is the innovation term
 163 with mean zero and constant variance σ_{α}^2 .

- 164 2. Use the fitted QR model and forecasted values of indicators (from the
 165 First Stage) to forecast future PLVaRs:

$$\widehat{PLVaR}_{\alpha}(Y_{t+k,i}) = \widehat{\beta}_{0,1-\alpha} + \sum_{j=1}^p \widehat{\beta}_{j,1-\alpha} \widehat{x}_{j,t+k,i}, \quad k = 1, \dots, h$$

166 *2.5. Model accuracy measures*

167 **Root mean squared error:** The common accuracy measure in time
 168 series forecasting models, is the Root Mean Square Error (RMSE). For M -
 169 variate time series the RMSE is formulated as follows:

$$RMSE = \sqrt{\sum_{i=1}^M \sum_{t=1}^N (y_{t,i} - \widehat{y}_{t,i})^2},$$

170 where $\widehat{y}_{t,i}$ is the forecasted value of time series.

171 **Exceedance rate:** Suppose \widehat{Q}_τ is the estimated value of τ th quantile
 172 based on observations y_1, \dots, y_N . The exceedance rate of \widehat{Q}_τ is the relative
 173 frequency of the observations greater than $\widehat{Q}_\tau(Y)$. If the estimation of τ
 174 quantile is accurate, the exceedance rate should be close to $1 - \tau$. In risk
 175 assessment applications, the exceedance rate is used to evaluate the accuracy
 176 of estimated VaR. If the exceedance rate is less than $1 - \tau$ the estimated VaR
 177 will present the worst case scenario accurately.

178 In this research, the exceedance rate is used to investigate the accuracy
 179 of QR in PLVaR forecasting (with risk level α).

$$ER_\alpha = \frac{1}{N} \sum_{i=1}^M \sum_{t=1}^N I_{(y_{t,i} > PLVaR_\alpha(Y_{t,i}))}$$

180 where $I_{(\cdot)}$ is Indicator function. Exceedance rate lower than α means the risk
 181 of using $PLVaR_\alpha(Y_{t,i})$ as the worst case scenario is less than α .

182 3. Data Description and Results

183 In order to forecast the waterborne and disease outbreak risk, we use the
 184 input dataset, published by World Health Organization (WHO) and used to
 185 calculate the 2000-2016 Disease burden and mortality estimates. The dataset
 186 contains the annual number of deaths cussed by 13 waterborne diseases be-
 187 tween 1998 and 2016, for 22 European and North American countries (WHO,
 188 2018)¹. The annual number of deaths per million, cussed by each disease, is
 189 a measure of disease outbreak for that disease.

190 Table 1 shows the list of waterborne disease considered in this study whilst
 191 Table 2 shows the list of countries involved. The $PLVaR_\alpha$ is considered
 192 as the $(1 - \alpha)$ th quantile of the annual number of deaths per million. The
 193 $PLVaR$ is forecasted using water related environmental and socio-economic
 194 indicators. The description of the indicators are as follows:

- 195 • **FSS:** This indicator is based on an assessment of the percentage of fish
 196 stocks caught within a countrys Exclusive Economic Zone (EEZ) that
 197 are overexploited or collapsed(Wendling et al., 2018; YCELP, 2018).

¹The dataset is available from World Health Organization (http://www.who.int/healthinfo/global_burden_disease/estimates/en/). The original dataset contains 47 countries from Europe and North America. The countries with no records of water- or disease-related environmental indicators, in that period, are dropped from this study.

Table 1: Waterborne diseases in this study.

1	Chlamydia	8	Dengue
2	Diarrhoeal Diseases	9	Japanese Encephalitis
3	Pertussis	10	Trachoma
4	Poliomyelitis	11	Ascariasis
5	Malaria	12	Trichuriasis
6	Schistosomiasis	13	Hookworm Disease
7	Onchocerciasis		

Table 2: List of countries in this study.

1	Canada	9	Guatemala	17	Puerto Rico
2	Croatia	10	Iceland	18	Republic of Moldova
3	Denmark	11	Ireland	19	Sweden
4	Estonia	12	Italy	20	Switzerland
5	Finland	13	Latvia	21	United Kingdom
6	France	14	Netherlands	22	United States of America
7	Germany	15	Panama		
8	Greece	16	Poland		

- 198 • **FPRO:** Fisheries production (Total) (tonnes)²(FAO, 2018)
- 199 • **FWP:** Freshwater KBAs completely covered by protected areas (SDG
200 15.1.2) (Percentage)(BirdLife Internationa, 2018)
- 201 • **POP14:** Child population 0-14 (% of total) (% of population)(UNPD,
202 2018)
- 203 • **POP65:** Elderly population 65 and above (% of total) (% of popula-
204 tion)(UNPD, 2018)
- 205 • **POPG:** Population growth (Percentage)(UNPD, 2018)
- 206 • **IS_R:** Access to improved sanitation: rural (% of rural population)
207 (UNMDG, 2018)
- 208 • **IS_U:** Access to improved sanitation: urban (% of urban population)
209 (UNMDG, 2018)
- 210 • **IWS_R:** Access to improved water sources: rural (% of rural popula-
211 tion) (UNMDG, 2018)
- 212 • **IWS_U:** Access to improved water sources: urban (% of urban popula-
213 tion) (UNMDG, 2018)

214 The FSS, FPRO and FWP indicators, are the environmental indicators
215 related to the freshwater disease risk. For instance, the countries with larger
216 FSS (and relatively lower FPRO) has a higher risk of freshwater disease
217 (Peeler and Feist , 2011). Indicators POP14, POP65 and POPG, indicate
218 the structure of the population. These indicators are included in the study
219 due to the fact that on one hand, child and elderly populations are more
220 vulnerable in disease outbreaks. On the other hand, the larger child popu-
221 lation increase the risk of break out since they usually are cureless while the
222 elderly population are more cautious and usually more experienced. Indica-
223 tors IS_R, IS_U, IWS_R and IWS_U are related to government policies and
224 infrastructural developments related to clean water resources.

²The rest is downloaded from <http://environmentlive.unep.org/downloader>

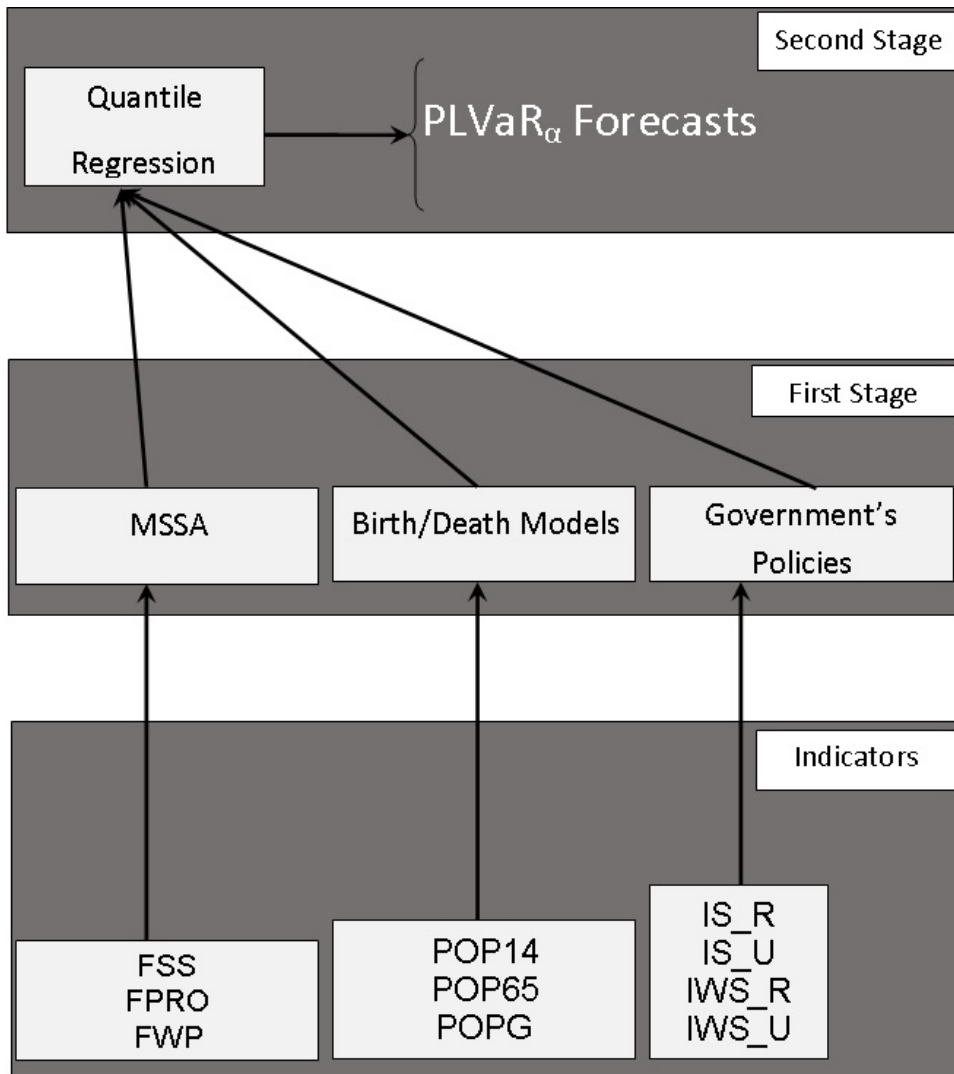


Figure 1: MSSA-QR model for waterborne disease PLVaR forecasting

225 The *PLVaR* is forecasted using the MSSA-QR model for confidence levels
 226 0.9, 0.95 and 0.99 (risk levels $\alpha = 0.1, 0.05, 0.01$). Figure 1 shows the diagram
 227 of the model.

228

229 In the first stage, MSSA is applied to FSS, FPRO and FWP as environ-
 230 mental indicators. The number of components in MSSA is selected based on
 231 minimum in-sample RMSE, using the data available before 2011. Since we do
 232 not have access to government policies on water and sanitation resources (i.e.
 233 IS_R, IS_U, IWS_R and IWS_U) in all of these 22 countries, MSSA is used to

Table 3: Out-of-sample RMSE produced by HMSSA-R, the number of components and window length in MSSA.

Indicator	RMSE					r [†]	L [‡]
	2011	2012	2013	2014	2015		
FSS	11.9396	16.1707	16.3747	16.587	. ^a	2	31
FPRO	1.69E+05	1.78E+05	1.57E+05	1.92E+05	1.58E+05	1	10
FWP	11.1997	13.5262	19.1778	21.2637	23.5379	1	10
IS_R	2.0862	2.5818	3.0744	3.505	3.5473	1	7
IS_U	0.5605	0.5576	0.56	0.5676	0.9434	1	7
IWS_R	2.0018	2.3421	2.6736	2.9148	2.9185	1	11
IWS_U	0.6103	0.6915	0.787	0.8225	0.8248	1	11

.[†] Number of components selected based on minimum in-sample RMSE

.[‡] Window length selected based on minimum in-sample RMSE

.^a The RMSE is not calculated since the 2015 observation is not available for any of the countries.

234 forecast these indicators too. The out-of-sample RMSE is calculated based
 235 on the forecasts for 2011 to 2015. Table 3 shows the out-of-sample RMSE
 236 for each year and indicator. As mentioned before, the POP14, POP65 and
 237 POPG indicator forecasts are available based on Berth/Death models from
 238 <http://www.un.org/en/development/desa/population/>.

239 In the second stage, the data from 1998 to 2010 are used to estimate the
 240 QR model coefficients in each confidence level. Table 4 shows the exceedance
 241 rate (ER_α) in each disease and confidence level for the estimated PLVaR. The
 242 out-of-sample ER_α for forecasted PLVaR (from 2011 to 2015) are given in
 243 Tables 5 and 6.

244 According to the Table 4, the in-sample ER_α is less than the risk level
 245 for most diseases. In more common diseases, (i.e. Diarrhoea, Pertussis and
 246 Malaria), however, the ER_α is slightly larger than the risk level. We record
 247 similar results during the out-of-sample forecasting exercise. Tables 5 and 6
 248 show that in all time horizons (from 2011 to 2015), for less common diseases,
 249 the ER_α does not exceed the risk level.

Table 4: In-sample Exceedance rate (ER_α) for estimated PLVaR based on 1998-2010 data.

Disease	Confidence Level [†]			Disease	Confidence Level [†]		
	0.9	0.95	0.99		0.9	0.95	0.99
Chlamydia	0.0185	0.0185	0.0074	Dengue	0.0296	0.0185	0.0000
Diarrhoeal Diseases	0.1148	0.0704	0.0074	Japanese Encephalitis	0.0185	0.0185	0.0037
Pertussis	0.1000	0.0556	0.0333	Trachoma	0.0185	0.0185	0.0037
Poliomyelitis	0.0741	0.0667	0.0000	Ascariasis	0.0333	0.0222	0.0148
Malaria	0.0807	0.0526	0.0246	Trichuriasis	0.0037	0.0037	0.0037
Schistosomiasis	0.0741	0.0519	0.0185	Hookworm	0.0222	0.0148	0.0000
Onchocerciasis	0.0037	0.0037	0.0037				

.[†] Confidence Level is $1 - \alpha$ where α is risk level.

250 Overall, according to these results, it is evident that the MSSA-QR model
 251 and the forecasted PLVaR values can be used as useful measures for fore-
 252 casting the worst case scenario in waterborne disease control and prevention.
 253 The model is not without its weaknesses, as we notice that it struggles at
 254 forecasting the more common disease like Diarrhoea, Pertussis and Malaria.
 255 However, we believe the performance for these diseases could be improved
 256 using more indicators. This is because the more common diseases are usually
 257 affected by more socioeconomic and environmental variables. For instance,
 258 the climatological and economic-development variables could affect the risk
 259 of a Malaria outbreak.

260 4. Conclusion

261 In this paper, a new model for forecasting the disease outbreak risk is
 262 proposed. In order to quantify the risk, we adopt a risk measure from finan-
 263 cial risk analysis and develop the Population Loss Value at Risk (PLVaR)
 264 as a measure of disease outbreak risk. The larger values of PLVaR show
 265 the bigger risk of disease outbreak. The PLVaR is forecasted using a two
 266 stage model based on Multivariate Singular Spectrum Analysis and Quantile
 267 Regression (MSSA-QR model). The proposed risk measure (PLVaR) and

Table 5: Out-of-sample Exceedance rate (ER_α) for estimated PLVaR.

Disease	Confidance Level [†]	ER_α				
		2011	2012	2013	2014	2015
Chlamydia	0.9	0.0000	0.0000	0.0000	0.0000	0.0000
	0.95	0.0000	0.0000	0.0000	0.0000	0.0000
	0.99	0.0000	0.0000	0.0000	0.0000	0.0000
Diarrhoeal Diseases	0.9	0.4091	0.3636	0.3636	0.2857	0.3684
	0.95	0.3182	0.3182	0.2727	0.2857	0.2632
	0.99	0.2727	0.2273	0.2273	0.1905	0.2105
Pertussis	0.9	0.1364	0.1818	0.2727	0.2857	0.2105
	0.95	0.1364	0.1364	0.1818	0.2381	0.1053
	0.99	0.0909	0.1364	0.1364	0.1905	0.1053
Poliomyelitis	0.9	0.0455	0.0909	0.0909	0.0476	0.1053
	0.95	0.0000	0.0455	0.0455	0.0000	0.1053
	0.99	0.0000	0.0455	0.0455	0.0000	0.0000
Malaria	0.9	0.0455	0.1364	0.0455	0.0476	0.1053
	0.95	0.1364	0.1364	0.1818	0.1905	0.2632
	0.99	0.0909	0.0909	0.0455	0.0000	0.1053
Schistosomiasis	0.9	0.0000	0.1364	0.0000	0.0476	0.0000
	0.95	0.0000	0.1364	0.0000	0.0476	0.0000
	0.99	0.0000	0.0000	0.0000	0.0000	0.0000
Onchocerciasis	0.9	0.0000	0.0455	0.0455	0.0000	0.0000
	0.95	0.0000	0.0455	0.0455	0.0000	0.0000
	0.99	0.0000	0.0455	0.0455	0.0000	0.0000

[†] Confidence Level is $1 - \alpha$ where α is risk level.

Table 6: Out-of-sample Exceedance rate (ER_α) for estimated PLVaR.

Disease	Confidance	ER_α				
	Level [†]	2011	2012	2013	2014	2015
Dengue	0.9	0.0455	0.0455	0.1364	0.1429	0.1053
	0.95	0.0455	0.0455	0.0909	0.0952	0.0526
	0.99	0.0000	0.0000	0.0000	0.0000	0.0000
Japanese Encephalitis	0.9	0.0000	0.0455	0.0000	0.0476	0.0000
	0.95	0.0000	0.0455	0.0000	0.0476	0.0000
	0.99	0.0000	0.0455	0.0000	0.0000	0.0000
Trachoma	0.9	0.0000	0.0000	0.0000	0.0000	0.0526
	0.95	0.0000	0.0000	0.0000	0.0000	0.0526
	0.99	0.0000	0.0000	0.0000	0.0000	0.0526
Ascariasis	0.9	0.0000	0.0909	0.0000	0.0000	0.0000
	0.95	0.0000	0.0909	0.0000	0.0000	0.0000
	0.99	0.0000	0.0000	0.0000	0.0000	0.0000
Trichuriasis	0.9	0.0455	0.0000	0.0000	0.0000	0.0000
	0.95	0.0455	0.0000	0.0000	0.0000	0.0000
	0.99	0.0455	0.0000	0.0000	0.0000	0.0000
Hookworm	0.9	0.0455	0.0000	0.0000	0.0476	0.0526
	0.95	0.0455	0.0000	0.0000	0.0476	0.0526
	0.99	0.0000	0.0000	0.0000	0.0000	0.0526

.[†] Confidence Level is $1 - \alpha$ where α is risk level.

268 forecasting model (MASS-QR) is used to forecast the worst cases of water-
269 borne disease outbreaks in 22 European and North American countries based
270 on socio-economic and environmental indicators. The results show that the
271 proposed method perfectly forecasts the worst case scenario for less com-
272 mon waterborne diseases. According to our findings, the forecasting of more
273 common diseases needs more socio-economic and environmental indicators.

274 We evidence that the proposed method has the ability to forecast the
275 worst case scenarios in disease outbreak and provides a practical tool for
276 policy makers and health institutions to control and prevent the outbreaks.
277 Furthermore, introducing a PLVaR as a risk measure adopted from finan-
278 cial risk analysis opens a new door to epidemiological and environmental
279 risk analysis using other risk analysis tools in finance. For instance, us-
280 ing PLVaR, one may adopt the copula method to investigate the relations
281 between different outbreaks. Moreover, more research is required into devel-
282 oping and evaluating the accuracy of the proposed PLVar, MSSA-QR model
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