



Article

Estimation of Daily Reproduction Numbers during the COVID-19 Outbreak

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Abstract: (1) Background: The estimation of daily reproduction numbers throughout the contagiousness period is rarely considered, and only their sum R_0 is calculated to quantify the contagiousness level of an infectious disease. (2) Methods: We provide the equation of the discrete dynamics of the epidemic's growth and obtain an estimation of the daily reproduction numbers by using a deconvolution technique on a series of new COVID-19 cases. (3) Results: We provide both simulation results and estimations for several countries and waves of the COVID-19 outbreak. (4) Discussion: We discuss the role of noise on the stability of the epidemic's dynamics. (5) Conclusions: We consider the possibility of improving the estimation of the distribution of daily reproduction numbers during the contagiousness period by taking into account the heterogeneity due to several host age classes.

Keywords: daily reproduction number; COVID-19 outbreak; discrete epidemic growth equation; discrete deconvolution; COVID-19 in several countries



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1. Introduction

1.1. Overview and Literature Review

Following the severe acute respiratory syndrome outbreak caused by coronavirus SARS CoV-1 in 2002 [1] and the Middle East Respiratory Syndrome outbreak caused by coronavirus MERS-CoV in 2012 [2], the COVID-19 disease caused by coronavirus SARS CoV-2 is the third coronavirus outbreak to occur in the past two decades. Human coronaviruses, including 229E, OC43, NL63 and HKU1, are a group of viruses that cause a significant percentage of all common colds in humans [3]. SARS CoV-2 can be transmitted from person to person by respiratory droplets and through contact and fomites. Therefore, the severity of disease symptoms, such as cough and sputum, and their viral load, are often the most important factors in the virus's ability to spread, and these factors can change rapidly within only a few days during an individual's period of contagiousness. This ability to spread is quantified by the basic reproduction number R₀ (also called the average reproductive rate), a classical epidemiologic parameter that describes the transmissibility of an infectious disease and is equal to the number of susceptible individuals that an infectious individual can transmit the disease to during his contagiousness period. For contagious diseases, the transmissibility is not a biological constant: it is affected by numerous factors, including endogenous factors, such as the concentration of the virus in aerosols emitted by the patient (variable during his contagiousness period), and exogenous factors, such as geo-climatic, demographic, socio-behavioral and economic factors governing pathogen transmission (variable during the outbreak's history) [4–8].

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Due to these exogenous factors, R_0 might change seasonally, but these factor variations are not significant if a very short period of time is considered. R_0 depends also on endogenous factors such as the viral load of the infectious individuals during their contagiousness period, and the variations in this viral load [9–15] must be considered in both theoretical and applied studies on the COVID-19 outbreak, in which the authors estimate a unique reproduction number R_0 linked to the Malthusian growth parameter of the exponential phase of the epidemic, during which R_0 is greater than 1 (Figure 1). The corresponding model has been examined in depth, because it is useful and important for various applications, but the distribution of the daily reproduction number R_j at day j of an individual's contagiousness period is rarely considered within a stochastic framework [16–20].

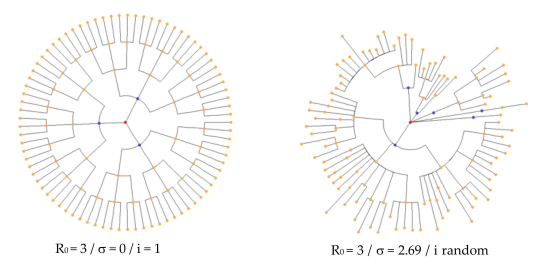


Figure 1. Spread of an epidemic disease from the first infectious "patient zero" (in red), located at the center of its influence sphere comprising the successive generations of infected individuals, for the same value of the reproduction number $R_0 = 3$, with a deterministic dynamic (**left**) and a stochastic one (**right**), with standard deviation σ of the uniform distribution on an interval centered on R_0 and with a random variable time interval i between infectious generations (after [16]).

We therefore defined a partial reproduction number for each day of an individual's contagiousness period, and, assuming initially that this number was the same for all individuals, we obtained the evolution equation for the number of new daily cases in a population. Assuming that the distribution of partial reproduction numbers (referred to as daily for simplicity) was subject to fluctuations, we calculated the consequences for their estimation, and we estimated them for a large number of countries, taking a duration of contagiousness of 3 followed by 7 days.

When this distribution is considered, it is possible to calculate its entropy as a parameter quantifying its uniformity and to simulate the dynamics of the infectious disease either using a Markovian model such as that defined in Delbrück's approach [17] or a classical discrete or ODE SIR deterministic model. In the Markovian case, R_0 can be calculated from the evolutionary entropy defined by L. Demetrius as the Kolmogorov–Sinaï entropy of the corresponding random process [18], which measures the stability of the invariant measure, dividing the population into the subpopulations S (individuals susceptible to but not yet infected with the disease), I (infectious individuals) and R (individuals who have recovered from the disease and now have immunity to it). In the deterministic case, R_0 corresponds to the Malthusian parameter quantifying its exponential growth, and the stability of the asymptotic steady state depends on the subdominant eigenvalue [19,20].

1.2. Calculation of R_0

In epidemiology, there are essentially two broad ways to calculate R_0 , which correspond to the individual-level modeling and to the population-level modeling. At the individual level, if we suppose the susceptible population size constant (hypothesis valid

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during the exponential phase of an epidemic), the daily reproduction rates of an individual are typically non-constant over his contagiousness period, and the calculations we present in the following define a new method for estimating R_0 , as the sum of the daily reproduction rates. This new approach allows us to have a clearer view on the respective influence on the transmission rate by endogenous factors (depending on the level of immunologic defenses of an individual) and exogenous factors (depending on environmental conditions).

2. Materials and Methods

The methodology chosen starts from an attempt to reconstruct an epidemic dynamic from the knowledge of the number R_{ikj} of people infected at day j by a given infectious individual i during the kth day of his period of contagiousness of length r. By summing up the number of new infectious individuals X_{j-k} present on day j -k where started their contagiousness, we find that the number of new infected people on day j is equal to:

$$X_{i} = \sum_{k=1,r} \sum_{i=1} X_{i-k} R_{ikj}$$
 (1)

We will assume in the following that R_{ikj} is the same, equal to R_k , for all individuals I and day j, then depends only on day k. Then, we have:

$$X_{j} = \Sigma_{k=1,r} R_{k} X_{j-k}$$
 (2)

The convolution Equation (2) is the basis of our modelling of the epidemic dynamics.

2.1. The Contagion Mechanism from a First Infectious Case Zero

Let us suppose that the secondary infected individuals are recruited from the centre of the sphere of influence of an infectious case zero and that the next infected individuals remain on a sphere centred on case 0, by just widening its radius on day 2. Therefore, the susceptible individuals C(j), which each infectious on day j-1 can recruit, are on a part of the sphere of influence of case 0 reached at day j (rectangles on Figure 2).

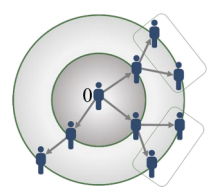


Figure 2. Spread of an epidemic disease from a first infectious case 0 (located at its influence sphere centre) progressively infecting its neighbours in some regions (rectangles) on successive spheres.

2.2. The Biphasic Pattern of the Virulence Curve of Coronaviruses

Mostly, the clinical course of patients with seasonal influenza shows a biphasic occurrence of symptoms with two distinct peaks. Patients have a classic influenza disease followed by an improvement period and a recurrence of the symptoms [11]. The influenza RNA virus shedding (the time during which a person might be contagious to another person) increases sharply one half to one day after infection, peaks on day 2 and persists for an average total duration of 4.5 days, between 3 and 6 days, which explains why we will choose in the following contagiousness duration these extreme values, i.e., either 3 or 6 days, depending on the positivity of the estimated daily reproduction numbers. It is common to consider this biphasic evolution of influenza clinically: after incubation of one day, there is a high fever (39–40 $^{\circ}$ C), then a drop in temperature before rising, hence the

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term "V" fever. The other symptoms, such as coughing, often also have this improvement on the second day of the flu attack: after a first feverish rise (39–39.5 $^{\circ}$ C), the temperature drops to 38 $^{\circ}$ C on the second day, then rises before disappearing on the 5th day, the fever being accompanied by respiratory signs (coughing, sneezing, clear rhinorrhea, etc.). By looking at the shape of virulence curves observed in coronavirus patients [12–16], we often see this biphasic pattern.

2.3. Relationships between Markovian and ODE SIR Approaches

In the following, we suppose that the susceptible population size remains constant, which constitutes a hypothesis valid during the exponential phase of epidemic waves. The Markovian stochastic and ODE deterministic approaches are linked by a common background consisting of the birth and death process approach used in the kinetics of molecular reactions by Delbrück [17], then in dynamical systems theory by numerous authors [18–23], namely in modelling of the epidemic spread in exponential growth. In the ODE approach, the Malthusian parameter is the dominant eigenvalue, and the equivalent in the Markovian approach is the Kolmogorov–Sinai entropy (called evolutionary entropy in [24–26]).

2.3.1. First Method for Obtaining the SIR Equation from a Deterministic Discrete Mechanism

Let us suppose the model is deterministic and denote by X_j the number of new infected cases at day j ($j \ge 1$), and R_k ($k = 1, \ldots, r$) the daily reproduction number at day k of the contagiousness period of length r for all infectious individuals. Then, we have obtained Equation (2) by supposing that the contagiousness behaviour is the same for all the infectious individuals:

$$X_j = \sum_{k=1,r} R_k X_{j-k},$$

which says that the X_{j-k} new infected at day j-k give R_k X_{j-k} new infected on day j, throughout a period of contagiousness of r days, the R_k 's being possibly different or zero. For example, if r=3, for the number X_5 of new cases at day 5, equation $X_5=R_1X_4+R_2X_3+R_3X_2$ means that new cases at day 4 have contributed to new cases at day 5 with the term R_1X_4 , R_1 being the reproduction number at first day of contagiousness of new infected individuals at day 4.

In matrix form, we obtain:

$$X = MR, (3)$$

where $X = (X_j, ..., X_{j-r-1})$ and $R = (R_1, ..., R_r)$ are r-dimensional vectors and M is the following r-r matrix:

$$\mathbf{M} = \begin{bmatrix} X_{j-1}, & X_{j-2}, \dots, & X_{j-r} \\ X_{j-k-1}, & X_{j-k-2}, \dots, & X_{j-k-r} \\ X_{j-r} & X_{j-r-1}, \dots, & X_{j-2r+1} \end{bmatrix}$$
(4)

It is easy to show that, if $X_0 = 1$ and r = 5 (estimated length of the contagiousness period for COVID-19 [12–21]), we obtain:

$$X_5 = R_1^5 + 4R_1^3R_2 + 3R_1^2R_3 + 3R_1R_2^2 + 2R_2R_3 + 2R_1R_4 + R_5$$
 (5)

The length r of the contagiousness period can be estimated from the ARIMA series of the stationary random variables Y_j 's, equal to the X_j 's without their trend, by considering the length of the interval on which the auto-correlation function remains more than a certain threshold, e.g., 0.1 [4]. For example, by assuming r = 3, if $R_1 = a$, $R_2 = b$ and $R_3 = c$, we obtain:

$$X_{0} = 1, X_{1} = a, X_{2} = a^{2} + b + c, X_{3} = a^{3} + 2ab, X_{4} = a^{4} + 3a^{2}b + b^{2} + 2ac,$$

$$X_{5} = a^{5} + 4a^{3}b + 3ab^{2} + 3a^{2}c + 2bc, X_{6} = a^{6} + 5a^{4}b + 4a^{3}c + 6a^{2}b^{2} + 6abc + b^{3} + c^{2},$$

$$X_{7} = a^{7} + 6a^{5}b + 5a^{4}c + 10a^{3}b^{2} + 12a^{2}bc + 4ab^{3} + 3b^{2}c + 3ac^{2}$$
(6)

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If R_1 and R_2 are equal, respectively, to a and b, and if a = b = R/2, c = 0, then, X_5 behaves like:

$$X_5 = R^5/32 + R^4/4 + 3R^3/8 \tag{7}$$

If R = 2, $\{X_j\}_{i=1,\infty}$ is the Fibonacci sequence, and more generally, for R > 0, the generalized Fibonacci sequence. Let us suppose now that b = c = 0 and a depends on the day j: $a_j = > C(j)$, where C(j) represents the number of susceptible individuals, which can be met by one contagious individual at day j. If infected individuals (supposed to all be contagious) at day j are denoted by I_j , we have:

$$X_{j} = \Delta I_{j} / \Delta j = (I_{j+1} - I_{j}) / (j+1-j) = \nu C(j) I_{j}$$
 (8)

Let us suppose, as in Section 2.1, that the first infectious individual 0 recruits from the centre of its sphere of influence secondary infected individuals remaining in this sphere, and that the susceptible individuals recruited by the I_j infectious individuals present at day j are located on a part of the sphere of centered on the first infectious 0 obtained by widening its radius (Figure 2). Then, we can consider that the function C(j) increases, then saturates due to the fact that an infectious individual can meet only a limited number of susceptible individuals as the sphere grows. We can propose for C(j) the functional form C(j) = S(j)/(c + S(j)), where S(j) is the number of susceptible individuals at day j. Then, we can write the following equation, taking into account the mortality rate μ :

$$X_{j} = \Delta I_{j} / \Delta j = \nu C(j) I_{j} - \mu I_{j} = \nu I_{j} S(j) / (c + S(j)) - \mu I_{j}$$
(9)

This discrete version of epidemic modeling is used much less than the classic continuous version, corresponding to the ODE SIR model, with which we will show a natural link. Indeed, the discrete Equation (9) is close to SIR Equation (10), if the value of c is greater than that of S:

$$dI/dt = \nu IS/(c+S) - \mu I \tag{10}$$

2.3.2. Second Method for Obtaining the SIR Equation from a Stochastic Discrete Mechanism

Another way to derive the SIR equation is the probabilistic approach, which comes from the microscopic equation of molecular shocks by Delbrück [17] and corresponds to a classical birth-and-death process: if at least one event (with rates of contact ν , birth f, death μ or recovering ρ) occurs in the interval (t, t + dt), and by supposing that births compensate deaths, leaving constant the total size N of the population, we have:

$$\begin{split} \text{Probability } (\{S(t+dt)=k, I(t+dt)=N-k\}) &= P(S(t)=k, I(t)=N-k) \left[1-[\mu k + \nu k(N-k) - fk - \rho(N-k)] dt\right] \\ &\quad + P(S(t)=k-1, I(t)=N-k+1) \left[f(k-1) + \rho(N-k+1)\right] dt \\ &\quad - P(S(t)=k+1, I(t)=N-k-1) \left[\mu(k+1) + \nu(k+1) \left(N-k-1\right)\right] dt \end{split} \tag{11}$$

Hence, we have, if $P_k(t)$ denotes Probability($\{S(t) = k, I(t) = N - k\}$):

$$\begin{split} dP_k(t)/d &= [P(S(t+dt)=k, I(t+dt)=N-k) - P(S(t)=k, I(t)=N-k)]/dt \\ &= -P(S(t)=k, I(t)=N-k) \left[\mu k + \nu k \left(N-k\right) - f k - \rho (N-k)\right] \\ &+ P(S(t)=k-1, I(t)=N-k+1) \left[f(k-1) + \rho (N-k+1)\right] \\ &- P(S(t)=k+1, I(t)=N-k-1) \left[\mu (k+1) + \nu (k+1) (N-k-1)\right], \end{split}$$

and we obtain:

$$dP_k(t)/dt = -[\mu k + \nu k(N-k) - fk - \rho(N-k)]P_k(t) + [f(k-1) + \rho(N-k+1)]P_{k-1}(t) - [\mu(k+1) + \nu(k+1)(N-k1)]P_{k+1}(t) + [\mu(k+1) + \nu(k+1)(N-k)]P_k(t) + [$$

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Then, by multiplying by s^k and summing over k, we obtain the characteristic function of the random variable S. If births do not compensate deaths, we have:

Probability (
$$\{S(t+dt) = k, I(t+dt) = j\}$$
) = $P(S(t) = k, I(t) = j)$ (1 - $[\mu k + \nu kj - fk - \rho j]dt$)
+ $P(S(t) = k - 1, I(t) = j + 1)$ [$f(k - 1) + \rho(j + 1)$] dt (12)
- $P(S(t) = k + 1, I(t) = j - 1)$ [$\mu(k + 1) + \nu(k + 1)(j - 1)$] dt

If S and I are supposed to be independent and if the coefficients ν , f, μ and ρ are sufficiently small, S and I are Poisson random variables [27], whose expectations E(S) and E(I) verify:

$$dE(S)/dt = fE(S) - \nu E(SI) - \mu E(S) + \rho E(I)$$
or, if $f = \mu$, $dE(S)/dt \approx E(I) [-\nu E(S) + \rho]$, (13)

leading to the SIR equation for the variables S, I and R considered as deterministic:

$$dS/dt = -\nu SI + \rho R, dI/dt = \nu SI - kI - \mu I, dR/dt = kI - \rho R$$
(14)

3. Results

3.1. Distribution of the Daily Reproduction Numbers R_j 's along the Contagiousness Period of an Individual. A Theoretical Example Where They Are Supposed to Be Constant during the Epidemics

If R_0 denotes the basic reproduction number (or average transmission rate) in a given-population, we can estimate the distribution V (whose coefficients are denoted $V_j = R_j/R_o$) of the daily reproduction numbers R_j along the contagious period of an individual, by remarking that the number X_j of new infectious cases at day j is equal to $X_j = I_j - I_{j-1}$, where I_j is the cumulated number of infectious at day j, and verifies the convolution equation (equivalent to Equation (2)):

$$X_{j} = \sum_{k=1,r} R_{k} X_{j-k}, \text{ giving in continuous time}: \ X(t) = \int_{1}^{r} R(s) X(t-s) ds, \qquad (15)$$

where r is the duration of the contagion period, estimated by $1/(\rho + \mu)$, ρ being the recovering rate and μ the death rate in SIR Equation (14). r and S can be considered as constant during the exponential phases of the pandemic, and we can assume that the distribution V is also constant; then, V can be estimated by solving the linear system (equivalent to Equation (3)):

$$R = M^{-1}X \tag{16}$$

where M is given by Equation (4). Equation (16) can be solved numerically, if the pandemic is observed during a time greater than $1/(\rho + \mu)$. We will first demonstrate an example of how the matrix M can be repeatedly calculated for consecutive periods of length equal to that of the contagiousness period (supposed to be constant during the outbreak), giving matrix series M_1, M_2, \ldots Following Equation (4), we put the values of X_i 's in the two matrices below, with r=3 for two periods, the first from day 1 to day 3 and the second from day 4 to day 6.

$$M_1 = \begin{bmatrix} X_4 & X_3 & X_2 \\ X_3 & X_2 & X_1 \\ X_2 & X_1 & X_0 \end{bmatrix}, \ M_2 = \begin{bmatrix} X_6 & X_5 & X_4 \\ X_5 & X_4 & X_3 \\ X_4 & X_3 & X_2 \end{bmatrix}, \dots,$$

where, after Equation (6), M_1 and M_2 can be calculated from the R_i 's as:

$$M_1 = \left[\begin{array}{ccc} R_1^4 + 3R_1^2R_2 + 2R_1R_3 + R_2^2 & R_1^3 + 2R_1R_2 + R_3 & R_1^2 + R_2 \\ R_1^3 + 2R_1R_2 + R_3 & R_1^2 + R_2 & R_1 \\ R_1^2 + R_2 & R_1 & 1 \end{array} \right] \text{,}$$

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and M₂ is given by:

$$\left[\begin{array}{cccc} R_1^6 + 5R_1^4R_2 + 4R_1^3R_3 + 6R_1R_2R_3 + 6R_1^2R_2^2 + R_2^3 + R_3^2 & R_1^5 + 4R_1^3R_2 + 3R_1^2R_2 + 2R_2R_3 + 3R_3R_1^2 & R_1^4 + 3R_1^2R_2 + 2R_1R_3 + R_2^2 \\ R_1^5 + 4R_1^3R_2 + 3R_1^2R_2 + 2R_2R_3 + 3R_3R_1^2 & R_1^4 + 3R_1^2R_2 + 2R_1R_3 + R_2^2 & R_1^3 + 2R_1R_2 + R_3 \\ R_1^4 + 3R_1^2R_2 + 2R_1R_3 + R_2^2 & R_1^3 + 2R_1R_2 + R_3 \end{array}\right]$$

Additionally, from Equation (2), if, for instance, j = 8 and r = 3, then we have the expression below, which means that the new cases on the 8th day depend on the new cases detected on the previous days 7, 6 and 5, supposed to be in a period of contagiousness of 3 days:

$$X_8 = \sum_{k=1,3} R_k X_{8-k} = R_1 X_7 + R_2 X_6 + R_3 X_5$$
 (17)

Let us suppose now that the initial R_i 's on a contagiousness period of 3 days, are equal

$$\begin{bmatrix} R_1 \\ R_2 \\ R_3 \end{bmatrix} = \begin{bmatrix} 2 \\ 1 \\ 2 \end{bmatrix}, \text{ then matrix M defined by } M_{ij} = X_{7-(i+j)} \text{ gives the } R_j\text{'s from Equation (16),}$$
hence allows the calculation of $X_i = \sum_{i=1,2}^{n} R_i X_{i-1}$.

hence allows the calculation of $X_j = \sum_{k=1,3} R_k X_{j-k}$.

The inverse of M is denoted by M^{-1} and verifies: $R = M^{-1}X$, where $X = (X_6, X_5, X_4)$, with $X_1 = 1$, $X_2 = 2$, $X_3 = 5$, $X_4 = 14$, $X_5 = 37$, $X_6 = 98$ and we obtain:

$$\mathbf{M}_{1}^{-1} = \begin{bmatrix} 37 & 14 & 5 \\ 14 & 5 & 2 \\ 5 & 2 & 1 \end{bmatrix}^{-1} = \begin{bmatrix} -1/4 & 1 & -3/4 \\ 1 & -3 & 1 \\ -3/4 & 1 & 11/4 \end{bmatrix},$$

and a deconvolution gives the resulting R_i's:

$$\begin{bmatrix} -1/4 & 1 & -3/4 \\ 1 & -3 & 1 \\ -3/4 & 1 & 11/4 \end{bmatrix} \begin{bmatrix} 98 \\ 37 \\ 14 \end{bmatrix} = \begin{bmatrix} 2 \\ 1 \\ 2 \end{bmatrix} = \begin{bmatrix} R_1 \\ R_2 \\ R_3 \end{bmatrix}, \text{ thanks to the following calculation:}$$

$$R_1 = -49/2 + 37 - 21/2 = 2$$

$$R_2 = 98 - 111 + 14 = 1$$

$$R_3 = -147/2 + 37 + 77 = 2$$

We obtain for the resulting distribution of daily reproduction numbers the exact replica of the initial distribution. We obtain the same result by replacing M_1 by the matrix M_2 .

3.2. Distribution of the Daily Reproduction Numbers R_i 's When They Are Supposed to Be Random

Let us consider a stochastic version of the deterministic toy model corresponding to Equation (17), by introducing an increasing noise on the R_i's, e.g., by randomly choosing their values following a uniform distribution on the three intervals: [2 - a, 2 + a], [1 - a/2, 1 + a/2] and [2 - a, 2 + a] (for having a U-shape behavior), with increasing values of a, from 0.1 to 1, in order to see when the deconvolution would give negative resulting R_i 's, with conservation of the average of their sum R_0 , if the random choice of the values of the R_i's at each generation is repeated, following the stochastic version of Equation (2): $X_i = \Sigma_{k=1,r} (R_k + \varepsilon_k) X_{i-k}$, where r is the contagiousness period duration and ε_k is a noise perturbing R_k , whose distribution is chosen uniform on the interval [0, 2a] for k = 1,3, and [0, a] for k = 2. This choice is arbitrary, and the main reason of the randomization is to show that the deconvolution can give negative results for R_k 's, as those observed for increasing values of a, from 0.1 to 1, with explicit calculations for three consecutive periods, from day 1 to day 3, from day 4 to day 6, and from day 7 to day 9.

For each random choice of the values of the daily reproduction numbers R_i 's, we can calculate a matrix M_1 corresponding to Equation (3). Its inversion into the matrix M_1^{-1} makes it possible to solve the problem of deconvolution of Equation (2)—that is to say, to Computation 2021, 9, 109 8 of 31

obtain new R_j 's as a function of the observed X_k 's. We can then calculate a new matrix M_2 from these new R_j 's and thus continue during an epidemic the estimation of the daily reproduction numbers R_j 's from the successive matrices M_1, M_2, \ldots , and observed X_k 's.

1. For a = 0.1, let us randomly and uniformly choose the initial distribution of the daily reproduction numbers R_1 in the interval [1.9, 2.1], R_2 in [0.95, 1.05] and R_3 in [1.9, 2.1] as R_1 = 2.1, R_2 = 0.95, R_3 = 2.1. Then, the transition matrix M_1 is equal to:

$$M_1 = \left[\begin{array}{ccc} 41.7391 & 15.351 & 5.36 \\ 15.351 & 5.36 & 2.1 \\ 5.36 & 2.1 & 1 \end{array} \right]$$
 and we have:

$$\mathbf{M}_{1}^{-1} = \begin{bmatrix} -0.2154195 & 0.92857143 & -0.7953515 \\ 0.92857143 & -2.95 & 1.2178571 \\ -0.7953515 & 1.2178571 & 2.705584 \end{bmatrix}$$

From X_6 = 113.491, X_5 = 41.7391, X_4 = 15.351, resulting R_j 's are: R_1 = 2.1, R_2 = 0.95, R_3 = 2.1.

The next initial R_i 's are chosen as: $R_1 = 2$, $R_2 = 0.95$, $R_3 = 1.9$ and we have:

$$X_7 = 2X_6 + 0.95X_5 + 1.9X_4 = 226.982 + 39.652 + 29.17 = 295.8$$

$$X_8 = 2X_7 + 0.95X_6 + 1.9X_5 = 591.6 + 107.816 + 79.304 = 778.72$$

Then, we obtain the matrices M_2 and M_2^{-1} :

$$\mathbf{M}_2 = \left[\begin{array}{cccc} 295.8 & 113.491 & 41.7391 \\ 113.491 & 41.7391 & 15.351 \\ 41.7391 & 15.351 & 5.36 \end{array} \right]$$

$$\mathbf{M}_{2}^{-1} = \begin{bmatrix} -0.07779371 & 0.20964295 & 0.00524305 \\ 0.20964295 & -1.0123552 & 1.26721348 \\ 0.00524305 & 1.26721348 & -3.48354228 \end{bmatrix}$$

Then, the resulting R_j 's equal: $R_1 = 2.0279$, $R_2 = 7.6158$, $R_3 = -16.426$. The next initial R_j 's are: $R_1 = 2$, $R_2 = 1.05$, $R_3 = 1.9$ and we have:

$$X_9 = 2X_8 + 1.05X_7 + 1.9X_6 = 1557.44 + 310.59 + 215.63 = 2083.66$$

$$X_{10} = 2X_9 + 1.05X_8 + 1.9X_7 = 4167.32 + 817.656 + 562.02 = 5546.996$$

From these values of X_9 and X_{10} , we obtain the matrices M_3 and M_3^{-1} :

$$M_3 = \begin{bmatrix} 2083.66 & 778.72 & 295.8 \\ 778.72 & 295.8 & 113.491 \\ 295.8 & 113.491 & 41.7391 \end{bmatrix}$$

$$\mathbf{M}_3^{-1} = \begin{bmatrix} 0.02596375 & -0.05192766 & -0.04280771 \\ -0.05192766 & 0.0256605 & 0.29823273 \\ -0.04280771 & 0.29823273 & -0.48358035 \end{bmatrix}$$

Then, the resulting R_i 's equal: $R_1 = 2.486$, $R_2 = -2.33$, $R_3 = 7.38769$.

2. For a = 1, let us choose the initial R_1 in [1, 3], R_2 in [0.5, 1.5] and R_3 in [1, 3], e.g., R_1 = 1, R_2 = 1.355 and R_3 = 1.1. Then, the transition matrix M_1 is equal to:

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$$M_1 = \left[\begin{array}{ccc} 9.101 & 4.81 & 2.355 \\ 4.81 & 2.355 & 1 \\ 2.355 & 1 & 1 \end{array} \right] \text{ and its inverse is given by:}$$

$$\mathbf{M}_{1}^{-1} = \begin{bmatrix} -1.11983471 & 2.02892562 & 0.60828512 \\ 2.02892562 & -2.93801653 & -1.84010331 \\ 0.60828512 & -1.84010331 & 1.40759184 \end{bmatrix}$$

New cases are: $X_6 = 18.209$, $X_5 = 9.101$, $X_4 = 4.81$, $X_3 = 2.355$, $X_2 = 1$, $X_1 = 1$, and by deconvoluting, we obtain the resulting R_j 's equal to: $R_1 = 1$, $R_2 = 1.355$, $R_3 = 1.1$, i.e., the exact initial distribution.

Let us now consider new initial R_j 's: $R_1 = 1$, $R_2 = 1$, $R_3 = 1$. That gives a new matrix M_2 , with new X_7 and X_8 calculated from the new initial R_j 's, by using the former values of X_6, \ldots, X_2 :

$$X_7 = X_6 + X_5 + X_4 = 18.209 + 9.101 + 4.81 = 32.12$$

 $X_8 = X_7 + X_6 + X_5 = 32.12 + 18.209 + 9.101 = 59.43$

Hence, we obtain:

$$\begin{split} M_2 = \begin{bmatrix} & 32.12 & 18.209 & 9.101 \\ & 18.209 & 9.101 & 4.81 \\ & 9.101 & 4.81 & 2.36 \end{bmatrix} \text{ and } \\ M_2^{-1} = \begin{bmatrix} & -0.35061537 & 0.1839519 & 0.97925345 \\ & 0.1839519 & -1.47916605 & 2.31025157 \\ & 0.97925345 & 2.31025157 & -8.0783421 \end{bmatrix} \end{split}$$

and the resulting R_j 's equal: $R_1 = 2.90$, $R_2 = 5.4888$, $R_3 = -14.696$. We calculate X_9 and X_{10} using new initial R_j 's: $R_1 = 3.0$, $R_2 = 0.5$, $R_3 = 2.9$:

$$X_9 = 3X_8 + 0.5X_7 + 2.9X_6 = 178.29 + 16.06 + 52.81 = 247.16$$

$$X_{10} = 3X_9 + 0.5X_8 + 2.9X_7 = 741.48 + 29.715 + 93.148 = 864.343$$

Hence, we obtain:

$$\begin{split} M_3 = \begin{bmatrix} 247.16 & 59.43 & 32.12 \\ 59.43 & 32.12 & 18.209 \\ 32.12 & 18.209 & 9.101 \end{bmatrix} \text{ and } \\ M_3^{-1} = \begin{bmatrix} 0.00718287 & -0.00805357 & -0.00923703 \\ -0.00805357 & -0.22288084 & 0.47435642 \\ -0.00923703 & 0.47435642 & -0.80659958 \end{bmatrix} \end{split}$$

and the resulting R_i 's equal: $R_1 = 3.66898$, $R_2 = -33.857$, $R_3 = 61.32$.

More precise simulation results are given in Table 1, which summarizes computations made for random choices of R_j 's distributions, for a=0.1 and a=1 and until time 20. These simulations show a great sensitivity to noise, but a qualitative conservation of their U-shaped distribution along the contagiousness period of individuals. More precisely, because of the presence of noise on the R_j 's, we cannot always obtain positive values from the data for the R_j 's by applying the deconvolution, which explains the presence of negative values in empirical examples, as in the theoretical noised examples. A way to solve this problem could be to suppose that noise is stationary during all of the growth period of a wave, then calculate the R_j 's for all running time windows of length equal to the contagiousness duration and then obtain the mean of the R_j 's corresponding to these windows. As this stationary hypothesis is not widely accepted, we prefer to keep negative values and focus on the shape of the distribution of the R_j 's.

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Table 1. Simulation results obtained for extre	me noises $a = 0.1$ and $a = 1$, showing great variations of deconvoluted distribution of	
daily reproduction numbers X_i 's and a qualitative	tive conservation of their U-shaped distribution along contagiousness period.	

a	Initial R _j 's	t	X _t	X _{t+1}	X _{t+2}	Resulting R' _j s	R ₀	Distribution Shape, Sign R ₀
0.1	2.1; 0.95; 2.1	4	15.35	31.74	113.5	2.1; 0.95; 2.1	5.15	U-shape, positive
	2; 0.95; 1.9	6	113.5	295.8	778.7	2.03; 7.6; -16.4	-6.77	Inverted U-shape, negative
	2; 1.06; 1.9	8	778.7	2083.7	5547	2.49; -2.33; 7.39	7.55	U-shape, positive
	1.9; 1.05; 1.9	10	5547	14,207	36,776	2.69; -16.7; 43.8	29.8	U-shape, positive
	1.9; 0.95; 1.9	12	36,776	93,910	240,359	2.92; 1.68; -6.7	-2.1	Decreased shape, negative
	1.9; 1; 1.9	14	240,359	622,149	1,605,227	2.3; -4.83; 14.3	11.8	U-shape, positive
	2; 1.05; 1.9	16	1,605,227	4,331,630	11,561,153	2.76; 27; -70	-40.2	Inverted U-shape, negative
	1.9; 1; 1.95	18	11,561,153	29,558,395	76,502,587	2.5; -6.48; 17.9	13.9	U-shape, positive
	2; 1; 2.1	20	76,502,587	2,076,519	556,226,772	2.67; -7.6; 19.7	14.8	U-shape, positive
1	1; 1.355; 1.1	4	4.81	9.1	18.21	1; 1.355; 1.1	3.455	Inverted U-shape, positive
	1; 1; 1	6	18.21	32.12	59.43	2.9; 5.49; -14.7	-6.31	Inverted U-shape, negative
	3; 0.5; 2.9	8	59.43	247.16	864.34	3.7; -33.9; 61.3	31.1	U-shape, positive
	2.6; 0.7; 2.6	10	864.34	2574.82	7942	3; -1.79; 7.14	8.35	U-shape, positive
	2.5; 0.75; 1.5	12	7942.2	23,083.1	67,526.6	3.35; 2.54; -11.6	-5.71	Decreased shape, negative
	2.4; 0.8; 2.4	14	67,526.6	199,590	588,437	2.58; -0.5; 4.8	6.88	U-shape, positive
	2; 1; 2	16	588,437	1,511,517	4,010,652	2.72; -1.08; 3.19	4.83	U-shape, positive
	2.3; 1.15; 2.3	18	4,010,652	12,316,150	36,415,885	2.88; -7.9; 21.7	16.7	U-shape, positive
	2.8; 0.6; 2	20	36,415,885	117,375,471	375,133,150	3.7; 4.1; -17	-9.2	Inverted U-shape, negative

3.3. Distribution of the Daily Reproduction Numbers R_i's. The Real Example of France

Figure 3 gives the effective transmission rates R_e calculated between 20–25 October 2020 just before the second lockdown in France [28,29]. As the second wave of the epidemic is still in its exponential phase, it is more convenient (i) to consider the distribution of the marginal daily reproduction numbers and (ii) to calculate its entropy and simulate the epidemic dynamics using a Markovian model [4]. By using the daily new infected cases given in [30], we can calculate, as in Section 3.1, the inverse matrix M^{-1} for the period from 20 to 25 October 2020 (exponential phase of the second wave), by choosing 3 days for the duration of contagiousness period and the following raw data for new infected cases: 20,468 for 20 October, then 26,676, 41,622, 42,032, 45,422 and 52,010 for 25 October. Then, for France between 15 February and 27 October 2020, we obtain the daily reproduction numbers given in Figure 3 with a U-shape as observed for influenza viruses.

We have:

$$\mathbf{M}^{-1} = \begin{bmatrix} 45,422 & 42,032 & 41,622 \\ 42,032 & 41,622 & 26,676 \\ 41,622 & 26,676 & 20,468 \end{bmatrix}^{-1} = \begin{bmatrix} -0.0000163989812 & -0.0000292188776 & 0.00007142863 \\ -0.0000292188776 & 0.0000938161392 & -0.0000628537817 \\ 0.00007142863 & -0.0000628537817 & -0.00001447698 \end{bmatrix}$$

Hence, we can deduce the daily R_i 's, i.e., the vector (R_1, R_2, R_3) :

$$\begin{bmatrix} -0.0000163989812 & -0.0000292188776 & 0.00007142863 \\ -0.0000292188776 & 0.0000938161392 & -0.0000628537817 \\ 0.00007142863 & -0.0000628537817 & -0.00001447698 \end{bmatrix} \begin{bmatrix} 52,010 \\ 45,422 \\ 42,032 \end{bmatrix} = \\ \begin{bmatrix} -0.852911911949567 & -1.32717986039119 & 3.00228812555347 \\ -1.51967382631645 & 4.26131667592337 & -2.64187015405365 \\ 3.71500298367996 & -2.85494447414886 & -0.60849658654673 \end{bmatrix} = \begin{bmatrix} 0.82219725466 \\ 0.0997726955533 \\ 0.2515619229844 \end{bmatrix} = \begin{bmatrix} R_1 \\ R_2 \\ R_3 \end{bmatrix}$$

The effective reproduction number is equal to $R_0 \approx 1.174$, a value close to that calculated directly (Figure 3), giving V = (0.7, 0.085, 0.215), with a maximal daily reproduction number the first day of the contagiousness period. The entropy H of V is equal to:

$$H = -\Sigma_{k=1,r} V_k Log(V_k) = 0.25 + 0.21 + 0.33 = 0.79.$$

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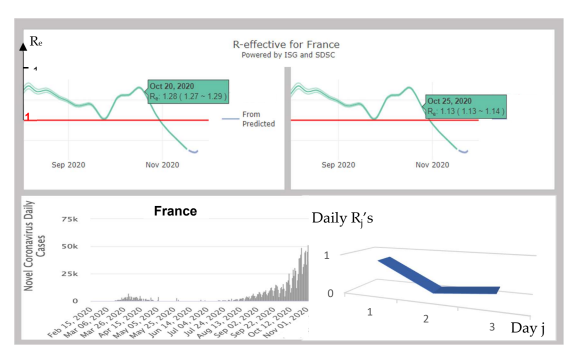


Figure 3. Top: estimation of the effective reproduction number R_e 's for 20 October and the 25 October 2020 (in green, with their 95% confidence interval) [28,29]. **Bottom left**: daily new cases in France between 15 February and 27 October [30]. **Bottom right**: U-shape of the evolution of the daily R_i 's along the 3-day contagiousness period of an individual.

3.4. Calculation of the Rj's for Different Countries

3.4.1. Chile

-0.01613675

0.03539322

By using the daily new infected cases given in [30], we can calculate M^{-1} for the period from 1 to 12 November 2020 (endemic phase), by choosing 6 days for the duration of the contagiousness period and the following 7-day moving average data for the new infected cases (Figure 4): 1400 for 1 November, then 1370, 1382, 1359, 1362, 1405, 1389, 1385, 1384, 1387, 1394 and 1408 for 12 November.

-0.00452917

-0.00686198

We have: 1394 1384 1389 1405 1387 1385 1384 1385 1405 1362 1387 1389 1384 1385 1389 1405 1362 1359 $M^{-1} =$ 1382 1385 1389 1405 1362 1359 1389 1405 1362 1359 1382 1370 1405 1362 1359 1382 1370 1400 -0.057142220.01016059 -0.009016640.01474588 0.00640175 0.03539322 0.01016059 -0.018272910.0106261 -0.007633630.02139586 -0.01613675-0.009016640.0106261 -0.005440510.02150289 -0.01468484-0.002863910.01474588 -0.007633630.02150289-0.01796266-0.00553414-0.005098010.00640175 0.02139586 -0.01468484-0.00553414-0.00305831-0.00452917

-0.00509801

-0.00286391

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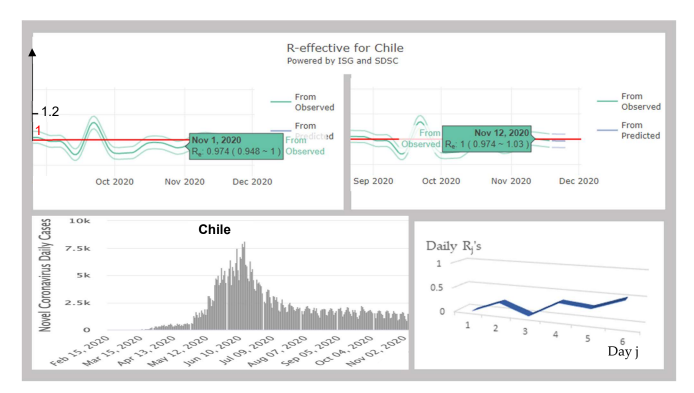


Figure 4. Top: estimation of the effective reproduction number R_e 's for the 1 November and the 12 November 2020 (in green, with their 95% confidence interval) [28,29]. **Bottom left**: Daily new cases in Chile between 1 November and 12 November [30]. **Bottom right**: U-shape of the evolution of the daily R_i 's along the infectious 6-day period of an individual.

Hence, after deconvolution, we obtain:

$$R = \begin{bmatrix} -0.36256122 \\ 0.22645436 \\ 0.01488726 \\ 0.33918287 \\ 0.28557502 \\ 0.50696243 \end{bmatrix}$$

The effective reproduction number is equal to $R_0 \approx 1.011$, a value close to that calculated directly, with a maximal daily reproduction number the last day of the contagiousness period. Due to the negativity of R_1 , we cannot derive the distribution V and therefore calculate its entropy. As entropy is an indicator of non-uniformity, an alternative could be to calculate it by shifting values of R_1 's upwards by the value of their minimum.

The quasi-endemic situation in Chile since the end of August, which corresponds to the increase of temperature and drought at this period of the year [4], gives a cyclicity of the new cases occurrence whose period equals the length of the contagiousness period of about 6 days, analogue to the cyclic phenomenon observed in simulated stochastic data of Section 3.2. with a similar U-shaped distribution of the R_i 's.

3.4.2. Russia

By using the daily new infected cases given in [30], we can calculate M⁻¹ for the period from 30 September to 5 October 2020 (exponential phase of the second wave), by choosing 3 days for the duration of the contagiousness period and the following raw data for new infected cases (Figure 5): 7721 for 30 September, then 8056, 8371, 8704, 9081, 9473 for 5 October.

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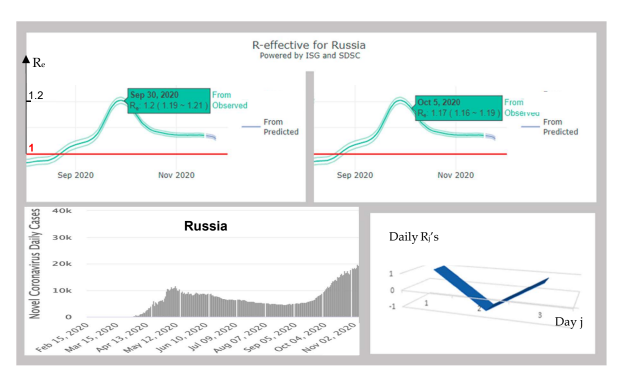


Figure 5. Top: estimation of the effective reproduction number R_e's for 30 September and the 5 October 2020 (in green, with their 95% confidence interval) [28,29]. **Bottom left**: Daily new cases in Russia between 15 February and 21 November [30]. **Bottom right**: U-shape of the evolution of the daily R_i's along the 3-day contagiousness period.

We have:
$$M^{-1} = \begin{bmatrix} 9081 & 8704 & 8371 \\ 8704 & 8371 & 8056 \\ 8371 & 8056 & 7721 \end{bmatrix}^{-1} \text{ and } \\ 8704 & 8371 & 8056 & 7721 \end{bmatrix}$$
 and
$$\begin{bmatrix} 0.031553440566948 & -0.027594779248393 & -0.005417732076268 \\ -0.027594779248393 & -0.00482333528665 & 0.034950483895551 \\ -0.005417732076268 & 0.034950483895551 & -0.030463575061795 \end{bmatrix} \begin{bmatrix} 9473 \\ 9081 \\ 8704 \end{bmatrix} = \begin{bmatrix} R_1 \\ R_2 \\ R_3 \end{bmatrix},$$
 where:
$$R_1 = 298.905742490698404 - 250.588190354656833 - 47.155939991836672 = 1.161612144205$$

$$R_2 = -261.405343820026889 - 43.80070773806865 + 304.209011826875904 = -0.997039731220$$

 $R_3 = -51.322175958486764 + 317.385344255498631 - 265.15495733786368 = 0.90821095914$

The effective reproduction number is equal to $R_0 \approx 1.073$, a value close to that calculated directly, with a maximal daily reproduction number the first day of the contagiousness period. Due to the negativity of R_2 , we cannot derive the distribution V and therefore calculate its entropy. The period studied corresponds to a local slow increase of new infected cases at the start of the second wave in Russia, which looks like a staircase succession of slightly inclined 4-day plateaus followed by a step: at the beginning of October, in Russia, new tightened restrictions (but avoiding lockdown) appeared [31], which could explain the change of the value of the slope observed in the new daily cases [30].

3.4.3. Nigeria

By using the daily new infected cases given in [30], we can calculate M⁻¹ for the period from 5 November to 10 November (endemic phase), by choosing 3 days for the duration of the contagiousness period and the following raw data for the new infected cases (Figure 6): 141 for 5 November, then 149, 133, 161, 164, and 166 for 10 November.

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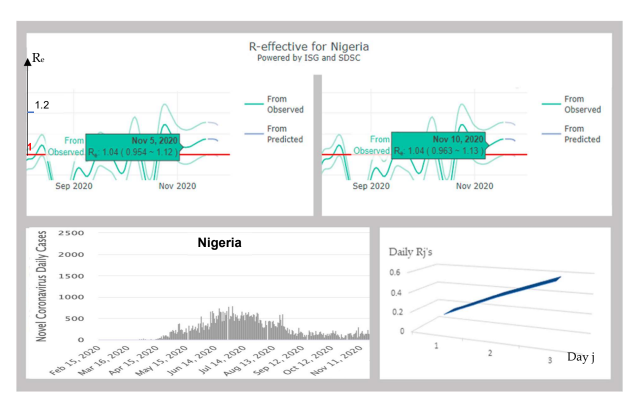


Figure 6. Top: estimation of the effective reproduction number R_e 's for 5 November and 10 November 2020 (in green, with their 95% confidence interval) [28,29]. **Bottom left**: Daily new cases in Nigeria between 15 February and 21 November [30]. **Bottom right**: increasing evolution of the daily R_i 's along the 3-day contagiousness period of an individual.

We have:

$$\mathbf{M}^{-1} = \begin{bmatrix} 164 & 161 & 131 \\ 161 & 131 & 149 \\ 131 & 149 & 141 \end{bmatrix}^{-1} = \begin{bmatrix} 0.01796807 & 0.01502897 & -0.03283028 \\ 0.01502897 & -0.02832263 & 0.01575332 \\ -0.03283028 & 0.01575332 & 0.02141264 \end{bmatrix}$$

After deconvolution, we obtain:

$$R = \left[\begin{array}{c} 0.16177513 \\ 0.38618314 \\ 0.58115333 \end{array} \right]$$

The effective reproduction number is equal to $R_0 \approx 1.129$, value close to that calculated directly, with a maximal daily reproduction number the last day of the contagiousness period. The distribution V equals (0.143, 0.342, 0.515) and its entropy H is equal to:

$$H = -\Sigma_{k=1,r} V_k Log(V_k) = 0.29 + 0.37 + 0.34 = 1.$$

In Appendix C, Table A1 gives the shape of the R_i's distribution for 194 countries.

3.5. Weekly Patterns in Daily Infected Cases

Daily new infected cases are highly affected by weekdays, such that case numbers are lowest at the start of the week and increase afterwards. This pattern is observed at the world level, as well as at the level of almost every single country or USA state. Hence, in order to estimate biologically meaningful reproduction numbers, clean of weekly patterns due to administrative constraints, analyses have to be restricted to specific periods shorter than a week, or at rare occasions when patterns escape the administrative constraints. This weekly phenomenon occurs during exponential increase as well as decrease phases of the pandemic and during endemic periods in numbers of daily cases (Figure 6). In

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addition, the daily new infected case record is discontinuous for many countries/regions, which frequently publish, on Monday or Tuesday, a cumulative count for that day and the weekend days. For example, Sweden typically publishes only four numbers over one week, the one on Tuesday cumulating cases for Saturday, Sunday and the two first weekdays. Discontinuity in records further limits the availability of data enabling detailed analyses of daily reproduction numbers and can be considered as extreme weekday effects on new case records due to various administrative constraints.

We calculated Pearson correlation coefficients r between a running window of daily new case numbers of 20 consecutive days and a running window of identical duration with different intervals between the two running windows. These Pearson correlation coefficients r typically peak with a lag of seven days between the two running windows.

The mean of these correlations are for each day of the week from Tuesday (data making up for the weekend underestimation) to Monday: 0.571, 0.514 (0.081), 0.383 (0.00008), 0.347 (0.00003), 0.381 (0.00006), 0.468 (0.000444) and 0.558 (0.03916), with, in parentheses, the p-value of the one-tailed paired t-test showing that the correlation observed with running windows starting Tuesday are more than the others (see also supplementary material). This could reflect a biological phenomenon of seven infection days. However, examination of the frequency distributions of lags for r maxima reveals, besides the median lag at 7 days, local maxima for multiples of 7 (14, 21, 28, 35, etc.). About 50 percent of all local maxima in r involve lags that are multiples of seven (seven included).

This excludes a biological causation, except if data periodicity comes from an entrainment by the weekly "Zeitgeber" of census, near the duration of the contagiousness interval. We tried to control for weekdays using two methods, and combinations thereof. For the first method, we calculated z-scores for each weekday, considering the mean number of cases for each weekday, and subtracted that mean from the observed number for a day (Figure 7). This delta was then divided by the standard deviation of the number of cases for that weekday. The mean and standard variation are calculated across the whole period of study for each weekday.

The second method implies data smoothing using a running window of 5 consecutive days, where the mean number of new cases calculated across the five days is subtracted from the number of new cases observed for the third day. Hence, data for a given day are compared to a mean including two previous, and two later days (Figure 8).

We constructed two further datasets, where z-scores are applied in the first to data after smoothing from the second method and are applied in the second data after smoothing from the first method (not shown) (Figures 9 and 10).

These four datasets from daily new cases database [30] transformed according to different methods and combinations thereof designed to control for weekday were analysed using the running window method. Despite attempts at controlling for weekday effects, the median lag was always seven days across all four transformed datasets, and local maxima in lag distributions were multiples of seven. After data transformations, about 50 percent of all local maxima were lags that are multiples of seven, seven included.

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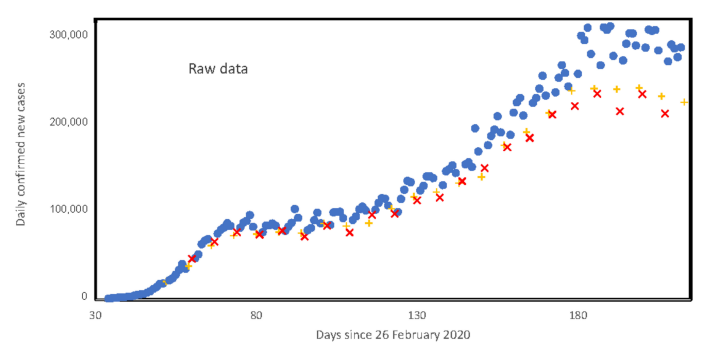


Figure 7. Confirmed world daily new cases (from [30]) as a function of days since 26 February until 23 August 2020 + indicates Sundays, X indicates Mondays.

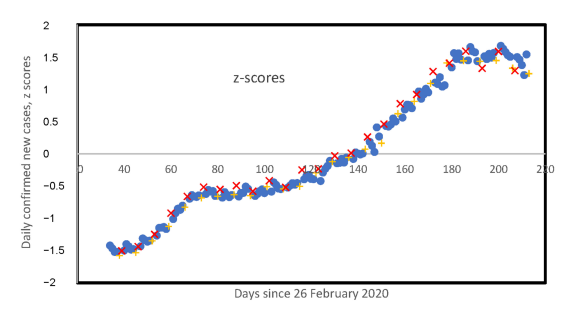


Figure 8. Z-transformed scores of confirmed world daily new cases [30], from Figure 6, as a function of days since 26 February 2020 until 23 August 2020 + indicates Sundays, X indicates Mondays. Z-transformations are specific to each weekday.

Visual inspection of plots of these transformed data versus time for daily new infected cases from the whole world shows systematic local biases in daily new infected cases (after transformation) on Sundays and Mondays, for all four transformed datasets, with Sundays and/or Mondays as local minima and/or local maxima, according to which method or combination thereof was applied to the data. Hence, the methods we used failed to neutralize the weekly patterns in daily new cases due to administrative constraints. This issue highly limits the data available for detailed analyses of daily new cases aimed at estimating biologically relevant estimates of reproduction numbers at the level of short temporal scales.

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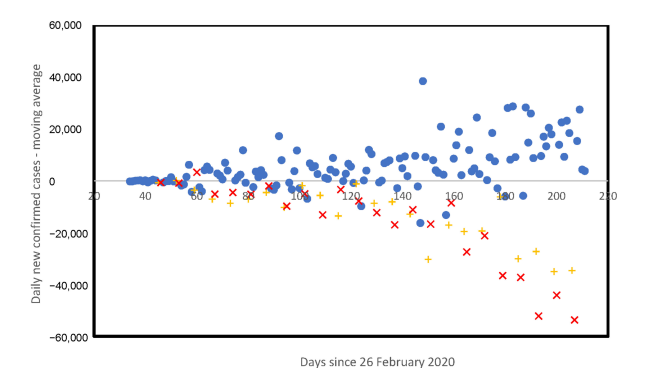


Figure 9. Smoothed confirmed world daily new cases [30], from Figure 7, as a function of days since 26 February 2020 until 23 August 2020 + indicates Sundays, X indicates Mondays. For each specific day j, the mean number of confirmed daily new cases calculated for days j - 1, j - 2, j, j + 1 and j + 2 is subtracted from the number for day j.

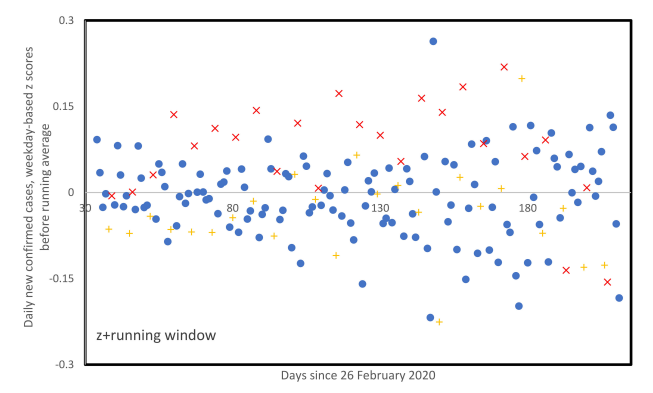
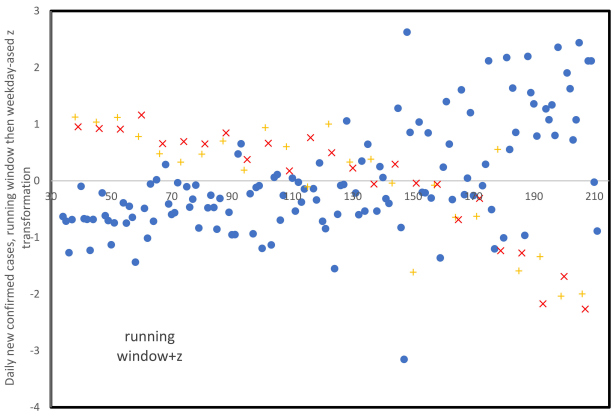


Figure 10. Smoothed confirmed world daily new cases [30] applied to z-scores from Figure 8, as a function of days since 26 February 2020 until 23 August 2020 + indicates Sundays, X indicates Mondays. Z-transformations are specific to each weekday. For specific day j, the mean number of confirmed new cases calculated for days j - 1, j - 2, j, j + 1, j + 2 is subtracted from the number for day j.

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By smoothing on five consecutive days of raw data (confirmed world daily new infected cases [24]) and applying the z-transformation, we obtain a better result in Figure 11 than in Figure 10 in order to neutralize the weekly pattern. The reason is that the smoothing largely eliminates the counting defect during weekends due either to fewer hospital admissions and/or less systematic PCR tests or to a lack of staff at the end of the week to perform the counts.



Days since 26 February 2020

Figure 11. Z-transformed scores of smoothed confirmed world daily new cases [30] smoothed data from Figure 9, as a function of days since 26 February 2020 until 23 August 2020. + indicates Sundays, X indicates Mondays. Z-transformations are specific to each weekday.

4. Discussion

The duration of the contagiousness period, as well as the daily virulence, are not constant over time. Three main factors, which are not constant during a pandemic, can explain this:

- In the virus transmitter, the transition between the mechanisms of innate (the first defense barrier) and adaptive (the second barrier) immunity may explain a transient decrease in the emission of the pathogenic agent during the phase of contagiousness [15],
- In the environmental transmission channel, many geophysical factors that vary over time can influence the transmission of the virus (temperature, humidity, altitude, etc.) [4–8],
- In the recipient of the virus, individual or public policies of prevention, protection, eviction or vaccination, which evolve according to the epidemic severity and the awareness of individuals and socio-political forces, can change the sensitivity of the susceptible individuals [32].

It is therefore very important to seek to estimate the average duration of the period of contagiousness of individuals and the variations, during this phase of contagiousness, of the associated daily reproduction numbers [33–39]. If the duration of the contagiousness

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phase is more than 3–5 days, for example ± 7 days, the periodicity of seven days observed for the new daily cases could result of an entrainment of the dynamics of new cases driven by the social "Zeitgeber" represented by the counting of new cases, less precise during the weekend (probably underestimated in many countries not working at this time). That questions the deconvolution over 3 and 5 days, giving some negative R_j . In a future work, we will compare our results with those obtained by deconvolutions on contagiousness durations between 3 and 12 days in order to obtain possibly more realistic values for the R_j 's, and hence, have perhaps a double explanation for the 7 days periodicity, both sociological and biological. Before this future work, we have extended our study using a duration r=3 of contagiousness to r=7. The results are given in Appendix B: they show the same existence of identical variations of U-shape type but they specify the values of R_j 's, more often positive and of more realistic magnitude, while keeping a sum approximately equal to R_0 .

Rhodes and Demetrius have pointed out the interest of the distribution of the daily reproduction numbers [24] with respect to the classical unique R_0 , even time-dependent [25]. In particular, they found that this distribution was generally not uniform, which we have confirmed here by showing many cases where we observe the biphasic form of the virulence already observed in respiratory viruses, such as influenza. The entropy of the distribution makes it possible to evaluate the intensity of its corresponding U-shape. This entropy is high if the daily reproduction numbers are uniform, and it is low if the contagiousness is concentrated over one or two days. If some R_j are negative, it is still possible to calculate this uniformity index, by shifting their distribution by a translation equal to the inverse of the negative minimum value.

We have neglected in the present study the natural birth and death rates by supposing them identical, but we could have taken into account the mortality due to the COVID-19. The discrete dynamics of new cases can be considered as Leslie dynamics governed by the matrix equation:

$$X_j = L X_{j-1},$$

where X_i is the vector of the new cases living at day j and L is the Leslie matrix given by:

$$L = \begin{bmatrix} R_1 & R_2 & R_3 & \dots & \dots & R_r \\ b_1 & 0 & 0 & \dots & \dots & 0 \\ 0 & b_2 & 0 & \dots & \dots & 0 \\ \vdots & \vdots & \ddots & \dots & \dots & \vdots \\ \vdots & \vdots & \vdots & \ddots & \dots & \vdots \\ 0 & 0 & 0 & \dots & b_{r-1} & 0 \end{bmatrix} \text{ and } X_{j-1} = \begin{bmatrix} X_{j-1} \\ X_{j-2} \\ X_{j-3} \\ \vdots \\ X_{j-r} \end{bmatrix},$$

where $b_j=1-\mu_j\leq 1, \ \forall \ i=1,\ldots,r$, is the recovering probability between days j and j+1. The dynamical stability for L^2 distance to the stationary infection age pyramid $P=\lim_j X_j/\Sigma_{i=j,j-r+1}X_i$ is related to $|\lambda-\lambda'|$, the modulus of the difference between the dominant and sub-dominant eigenvalues of L, namely $\lambda=e^R$ and λ' , where R is the Malthusian growth rate and P is the left eigenvector of L corresponding to λ . The dynamical stability for the distance (or symmetrized divergence) of Kullback–Leibler to P considered as stationary distribution is related to the population entropy P [26–32], which is defined if P is and P is and P is and P is follows:

$$H = -\sum_{i=1,r} p_i \, \text{Log}(p_i) / \sum_{i=1,r} p_i$$
 (18)

The mathematical characterization by the population entropy defined in Equation (16) of the stochastic stability of the dynamics described by Equation (16) has its origin in the theory of large deviations [40–42]. This notion of stability pertains to the rate at which the system returns to its steady state after a random exogenous and/or endogenous

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perturbation and it could be useful to quantify further the variations of the distribution of the daily reproduction numbers observed for many countries [43–53].

In summary, the main limitations of the present study are:

- The hypothesis of spatio-temporal stationarity of the daily reproduction numbers is no longer valid in the case of rapid geo-climatic changes, such as sudden temperature rises, which decrease the virulence of SARS CoV-2 [4], or mutations affecting its transmissibility.
- The still approximate knowledge of the duration r of the period of contagiousness necessitates a more in-depth study at variable durations, by retaining the value of r, which makes all of the daily reproduction numbers positive.
- The choice of uniform random fluctuations of the daily reproduction numbers is based on arguments of simplicity. A more precise study would undoubtedly lead to a unimodal law varying throughout the contagious period, the average of which following a U-shaped curve, of the type observed in the literature on a few real patients [10,54–58].

5. Conclusions and Perspectives

Concerning contagious diseases, public health physicians are constantly faced with four challenges. The first concerns the estimation of the basic reproduction number R₀. The systematic use of R₀ simplifies the decision-making process by policymakers, advised by public health authorities, but it is too much of a caricature to account for the biology behind the viral spread. We have observed in the COVID-19 outbreak that it was non-constant during an epidemic wave due to exogenous and endogenous factors influencing both the duration of the contagiousness period and the daily transmission rate during this phase [54–56]. Then, the first challenge concerns the estimation of the mean duration of the infectious period for infected patients. As for the transmission rate, realistic assumptions made it possible to obtain an upper limit to this duration [45], mainly due to the lack of viral load data in large patient cohorts (see Figure A1 in Appendix A from [57-59]), in order to better guide the individual quarantine measures decided by the authorities in charge of public health. This upper bound also makes it possible to obtain a lower bound for the percentage of unreported infected patients, which gives an idea of the quality of the census of cases of infected patients, which is the second challenge facing specialists of contagious diseases. The third challenge is the estimation of the daily reproduction number over the contagiousness period, which was precisely the topic of the present paper. A fourth interesting challenge for this community is the extension of the methods developed in the present paper to the contagious non-infectious diseases (i.e., without causal infectious agent), such as social contagious diseases [59-61], the best example being that of the pandemic linked to obesity, for which many concepts and modelling methods remain available.

Eventually, our approach using marginal daily reproduction numbers involving a certain level of noise in the dynamics of new daily infected cases defines a stochastic framework which describes phenomenologically the exponential phase as our results show for countries such as France, Russia, Sweden, etc. This stochastic modelling allows a better understanding of the role of the contagiousness period length and of the heterogeneity (e.g., the U-shape) of its daily reproduction number distribution in the COVID-19 outbreak dynamics [62–65]. On the medical level, the important message about the U-shape is that COVID-19 is similar to other viral diseases, such as influenza, with two successive reactions from the two immune defense barriers, innate cellular immunity first, which is not sufficient if symptoms persist, then adaptive immunity (cellular and humoral), which results in a transient decrease in contagiousness between the two phases. The medical recommendations are, in this case, never to take a transient improvement for a permanent disappearance of the symptoms. One could indeed, for a public health use, be satisfied after estimating the sum of the R_i's, that is to say, R₀ or the effective R_e. For an individual health use, it is important to know the existence of a minimum of the Ri's, which generally corresponds to a temporary clinical improvement, after the partial success of the innate

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immune defenses. This makes it possible to prevent the patient from continuing to respect absolute isolation and therapeutic measures, even if a transient improvement occurs; otherwise, they risk, as in the flu, a bacterial pulmonary superinfection (a frequent cause of death in the case of COVID-19). On the theoretical level, the interest of the proposed method is its generic character: it can be applied to all contagious diseases, within the very general framework of Equation (1), which makes no assumption about the spatial heterogeneity or the longitudinal constancy of the daily reproduction numbers. The deconvolution of Equation (1) poses a new theoretical problem when it is offered in this context, and our future research will propose new avenues of research in this field.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10 .3390/computation9100109/s1. Table S1. Presentation of the Pearson correlation coefficients between 20 numbers of world daily new cases observed between the days 34 to 53 after the 24 January 2020 (date of the start of the Covid-19 outbreak with confirmed cases in Europe) and series of 20 numbers of world daily new cases observed in running windows of length 20 days until day 213.

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Conflicts of Interest: The authors declare no conflict of interest.

Appendix A

Figure A1 shows a U-shaped evolution for the viral load in real [57] and in simulated [58] COVID-19 patients, and in real influenza-infected animals for the viral load and the body temperature [59].

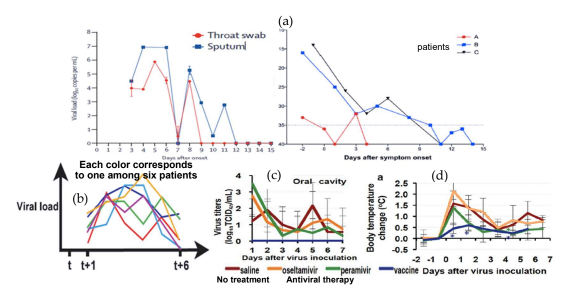


Figure A1. (a) Viral load in real COVID-19 patients [10], (b) in influenza-simulated patients [57] and (c) in real influenza-infected animals (red curve [58]), and (d) body temperature in real influenza-infected animals (red curve [58]).

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Appendix B

1. Beginning of the pandemic in France from 21 February 2020 to 9 March 2020

The numbers of new cases are:

21 February 2, 4, 19, 18, 39, 27, 56, 20, 67, 126, 209, 269, 236, 185 **9 March** Then, the matrix M is defined by:

$$M = \begin{bmatrix} 236 & 269 & 209 & 126 & 67 & 20 & 56 \\ 269 & 209 & 126 & 67 & 20 & 56 & 27 \\ 209 & 126 & 67 & 20 & 56 & 27 & 39 \\ 126 & 67 & 20 & 56 & 27 & 39 & 18 \\ 67 & 20 & 56 & 27 & 39 & 18 & 19 \\ 20 & 56 & 27 & 39 & 18 & 19 & 4 \\ 56 & 27 & 39 & 18 & 19 & 4 & 2 \end{bmatrix}$$

and we have:

Because,
$$X = \begin{bmatrix} 185 \\ 236 \\ 269 \\ 209 \\ 126 \\ 67 \\ 20 \end{bmatrix}$$
 , hence $R = M^{-1} X = \begin{bmatrix} 0.239 \\ 0.052 \\ -0.783 \\ -0.295 \\ 1.189 \\ 3.060 \\ 3.122 \end{bmatrix}$ and we can represent

the evolution of X_i 's on Figure A2.

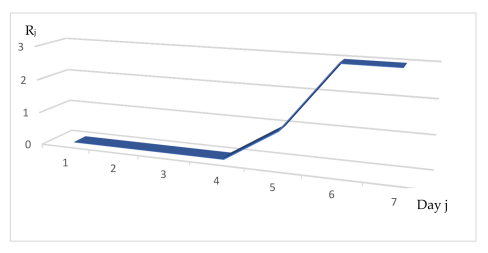


Figure A2. Values of the daily reproduction numbers R_j along the period of contagiousness of length 7 days.

The evolution of the Xj's along the period of contagiousness shows at day 4 a sharp increase and a saturation.

2. Exponential phase in France from 25 October 2020 to 7 November 2020 The numbers of new cases are:

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7 November 83,334, 58,581, 56,292, 39,880, 35,912, 51,104, 45,258, 33,447, 46,185, 44,705, 34,194, 31,360, 25,123, 48,808 25 **October**

Then, the matrix M is defined by:

$$M = \begin{bmatrix} 58,581 & 56,292 & 39,880 & 35,912 & 51,104 & 45,258 & 33,447 \\ 56,292 & 39,880 & 35,912 & 51,104 & 45,258 & 33,447 & 46,185 \\ 39,880 & 35,912 & 51,104 & 45,258 & 33,447 & 46,185 & 44,705 \\ 35,912 & 51,104 & 45,258 & 33,447 & 46,185 & 44,705 & 34,194 \\ 51,104 & 45,258 & 33,447 & 46,185 & 144,705 & 34,194 & 31,360 \\ 45,258 & 33,447 & 46,185 & 44,705 & 34,194 & 31,360 & 25,123 \\ 33,447 & 46,185 & 44,705 & 34,194 & 31,360 & 25,123 & 48,808 \end{bmatrix}$$

and we obtain

$$R = \begin{bmatrix} 2.867 \\ -1.231 \\ 1.351 \\ -2.705 \\ -0.155 \\ 0.223 \\ 0.769 \end{bmatrix}$$

The Figure A3 shows an evolution of the Xj's with a U-shape on the three first days along the period of contagiousness with a sum of R_j 's equal to 1.11, close to the effective reproduction number $R_e = 1.13$ [28].

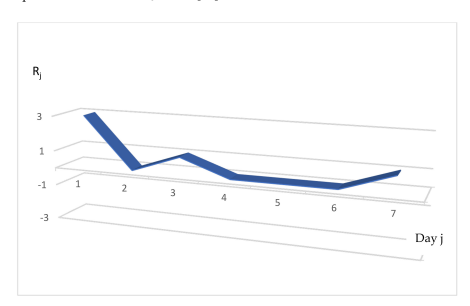


Figure A3. Values of the daily reproduction numbers R_j along the period of contagiousness of length 7 days.

3. Beginning of the pandemic in the USA from 21 February 2020 to 5 March 2020 The number of new cases are: **21 February** 20, 0, 0, 18, 4, 3, 0, 3, 5, 7, 25, 24, 34, 63 **5 March** Then, we have:

$$R = \begin{bmatrix} 0.466 \\ 0.584 \\ 1.547 \\ -1.044 \\ 0.174 \\ 0.297 \\ 0.692 \end{bmatrix}$$

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The evolution of the Xj's shows in Figure A4 a U-shape on day 4 with a sum of R_j 's equal to 2.72, less than the effective reproduction number $R_e = 3.27$ [28].

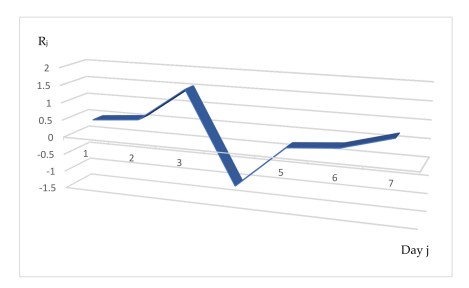


Figure A4. Values of the daily reproduction numbers R_j along the period of contagiousness of length 7 days.

4. USA exponential phase from 1 November 2020 to 4 November 2020

The numbers of new cases are:

N 14 163,961, 183,792, 167,665, 150,535, 159,565, 120,924, 108,248, 135,385, 136,292, 129,663, 113,709, 105,745, 86,030, 75,285 **N 1**

Then, we have:

$$R = \begin{bmatrix} 0.020 \\ -0.439 \\ 0.583 \\ -0.367 \\ 0.497 \\ -0.056 \\ 1.113 \end{bmatrix}$$

The evolution of the Xj's shows in Figure A5 a U-shape on the four last days with a sum of R_j 's equal to 1.35, close to the effective reproduction number R_e = 1.24 [28].

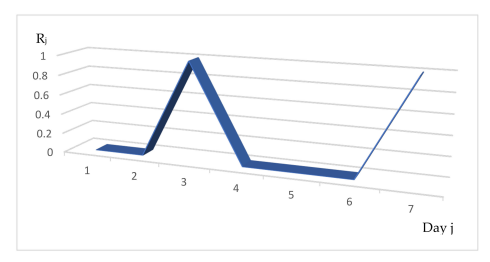


Figure A5. Values of the daily reproduction numbers R_j along the period of contagiousness of length 7 days.

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5. Beginning of the pandemic in the UK from 23 February 2020 to 7 March 2020

The number of new cases are:

23 February 4, 0, 0, 0, 3, 4, 3, 12, 3, 11, 33, 26, 43, 41 **7** March

Then, we have:

$$R = \begin{bmatrix} -0.388 \\ -1.189 \\ 1.334 \\ 1.960 \\ 4.862 \\ -0.170 \\ 3.479 \end{bmatrix}$$

Figure A6 shows an evolution of the Xj's with a U-shape on the three last days along the period of contagiousness with a sum of R_j 's equal to 9.88, higher than the effective reproduction number R_e = 2.95 [28].

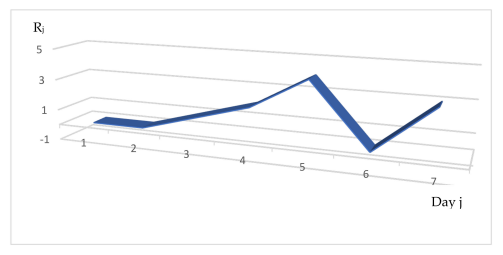


Figure A6. Values of the daily reproduction numbers R_j along the period of contagiousness of length 7 days.

6. UK exponential phase from 17 October 2020 to 30 October 2020

The numbers of new cases are:

30 October 24,350, 23,014, 24,646, 22,833, 20,843, 19,746, 22,961, 20,484, 21,195, 26,624, 21,282, 18,761, 16,943, 16,133 **17 October**

Then, we have:

$$R = \begin{bmatrix} 0.020 \\ 0.334 \\ 0.462 \\ -0.098 \\ -0.134 \\ -0.043 \\ 0.526 \end{bmatrix}$$

Figure A7 shows an evolution of the Xj's with a U-shape on the five last days along the period of contagiousness with a sum of R_{j} 's equal to 1.07, close to the effective reproduction number R_{e} = 1.06 [28].

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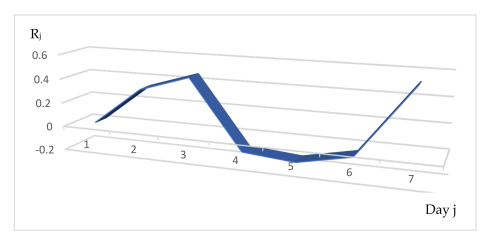


Figure A7. Values of the daily reproduction numbers R_j along the period of contagiousness of length 7 days.

Appendix C

Table A1 is built from new COVID-19 cases at the start of the first and second waves for 194 countries; it shows 42 among these 194 countries having a U-shape evolution of their daily R_j 's twice, for 12.12 ± 6 expected with 0.95 confidence ($p < 10^{-12}$), and 189 times, a U-shape evolution for all countries and waves (397), for 99.3 \pm 9 expected with 0.95 confidence ($p < 10^{-24}$). Hence, the U-shape is the most frequent evolution of daily R_j 's, which confirms the comparison with the behavior of seasonal influenza (see Section 2.2).

Table A1. Calculation of the daily R_i's and shape of their distribution for 194 countries and for the two first waves.

	All Countries		First Wa	ve		Second Way	⁄e	
No	Country Name	$\mathbf{R_0}$	$\mathbf{R}_{\mathbf{j'}}\mathbf{s}$	U-Shape	$\mathbf{R_0}$	R_j 's	U-Shape	
1	AFGHANISTAN	0.65	0.17; 0.09; 0.39	YES	0.04	-1.38; -0.36; 1.78	INCR	
2	ALGERIA	1.25	3.93; -6.21; 3.53	YES	0.91	1.28; -1.06; 0.69	YES	
3	ARUBA	5.46	10.31; -39.32; 34.47	YES	1.10	1.54; -1.60; 1.16	YES	
4	ANDORRA	1.36	1.00; 0.79; -0.43	DECR	0.12	4.34; -1.63; -2.59	DECR	
5	ANGOLA	0.63	0.33; 1.42; -1.12	INV	1.70	9.22; -1.58; -5.94	DECR	
6	ANTIGUA	1.92	0.00; 1.25; 0.67	INV	2.13	-0.40; 1.33; 1.20	INV	
7	ALBANIA	0.96	0.48; 0.50; -0.02	INV	0.66	1.98; -0.56; -0.76	DECR	
8	ARGENTINA	0.73	0.57; -1.28; 1.44	YES	0.36	1.27; 0.75; -1.66	DECR	
9	ARMENIA	4.43	17.99; -36.99; 23.43	YES	0.86	1.41; -0.97; 0.42	YES	
10	AUSTRALIA	2.79	-1.02; 3.47; 0.34	YES	1.50	-0.88; 0.68; 1.70	INCR	
11	AUSTRIA	1.17	-1.78; -0.05; 3.00	INCR	2.08	0.62; -3.55; 5.01	YES	
12	AZERBAIJAN	1.16	1.23; -1.32; 1.25	YES	0.37	10.36; -6.45; -3.54	YES	U-shaped
13	BAHAMAS	0.57	-0.13; -0.98; 1.68	YES	1.22	0.22; -0.86; 1.86	YES	5
14	BAHRAIN	1.10	-0.74; 0.28; 1.56	INCR	1.14	1.98; -2.69; 1.85	YES	a constant
15	BANGLADESH	1.04	2.37; -2.97; 1.64	YES	0.99	0.86; -0.69; 0.82	YES	YES
16	BARBADOS	1.86	0.86; -0.64; 1.64	YES	1.14	0.22; -0.81; 1.73	YES	Decreasing
17	BELARUS	1.57	-2.37; -4.58; 8.52	YES	1.07	-0.33; 0.24; 1.16	INCR	
18	BELGIUM	0.43	11.66; -15.63; 4.41	YES	2.23	1.17; -2.39; 3.45	YES	5
19	BELIZE	0.99	0.80; 0.42; -0.23	DECR	0.51	1.77; -0.21; -1.05	DECR	DECR
20	BENIN	0.85	0.81; 0.47; -0.43	DECR	0.85	1.17; 0.22; -0.54	DECR	Onej
21	BHUTAN	15.00	14.00; 15.00; -14.00	INV	1.08	0.80; 0.57; -0.29	DECR	Inverted
22	BOLIVIA	2.17	8.47; -1.17; -5.13	DECR	1.61	0.96; -0.30; 0.95	YES	Biphasic
23	BOSNIA	0.09	-1.06; -1.05 ; 2.20	INCR	1.56	-0.57; -0.51; 2.64	INCR	AT .
24	BOTSWANA	28.47	0.22; 0.00; 28.25	YES	28.43	0.22; -0.05; 28.26	YES	INV
25	BRAZIL	0.77	0.31; 1.08; -0.62	INV	0.46	1.21; 0.16; -0.91	DECR	Increasing
26	BRUNEI	1.08	0.10; -0.15; 1.13	YES	1.00	1.00; -1.00; 1.00	YES	Hicreasing
27	BULGARIA	5.06	14.73; -66.02; 56.35	YES	0.75	1.34; -0.98; 0.39	YES	
28	BURKINA FASO	1.08	0.72; -0.34; 0.70	YES	0.94	0.31; 0.24; 0.39	YES	INCR
29	BURUNDI	1.33	1.33; -0.67; 0.67	YES	2.18	0.53; 1.80; -0.15	INV	n vek
30	CABO VERDE	0.82	-0.08; -0.26; 1.16	YES	0.19	0.56; 1.37; -1.74	INV	
31	CAMBODIA	0.34	0.08; 0.25; 0.01	INV	0.27	0.06; 0.15; 0.06	INV	
32	CAMEROON	2.17	2.36; 1.25; -1.44	DECR	2.48	0.50; -0.25; 2.23	YES	
33	CANADA	1.10	-0.55; -0.72; 2.37	YES	0.44	2.36; -0.44; -1.48	DECR	
34	CAR	1.66	-0.07; 0.64; 1.09	INCR	0.33	0.44; -0.22; 0.11	YES	
35	CHAD	1.19	0.77; -1.15; 1.57	YES	0.77	1.19; 0.25; -0.67	DECR	
36	CHILE	1.00	0.72; 0.17; 0.11	DECR	1.64	0.37; -4.45; 5.72	YES	
37	CHINA	1.10	0.90; -0.49; 0.69	YES	0.87	1.16; 0.60; -0.89	DECR	

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 Table A1. Cont.

	All Countries	All Countries First Wave Second Wave					
No	Country Name	R_0	R _{j′} s	U-Shape	R_0	R _j 's	U-Shape
38	COLUMBIA	1.00	1.75; -0.86; 0.11	YES	1.47	-1.14; 3.08; -0.47	INV
39	COMOROS	3.75	0.00; -2.75; 6.5	YES	1.65	-0.58; 1.24; 0.99	INV
40	CONGO DEM	0.03	-0.37; -0.39; 0.79	YES	0.88	0.66; 0.74; -0.52	INV
41	CONGO REP	0.92	0.92; 0.92; -0.92	DECR	0.39	-0.12; 0.19; 0.32	INCR
42	COSTA RICA	0.50	-2.79; -3.84; 7.13	YES	1.26	1.21; -0.85; 0.90	YES
43	COTE D'VOIRE	1.18 0.75	-0.49; -0.63; 2.30	YES	2.09 0.57	4.32; -7.09; 4.86	YES YES
44 45	CROTIA CUBA		0.53; 0.79; -0.57	INV	0.57	0.68; -0.64; 0.53	YES
46	CURACAO	0.48 0.50	-37.25; 16.17; 21.56 3.00; -1.00 ; -1.50	INCR DECR	4.19	0.34; -0.73; 1.17 1.93; -4.01; 6.27	YES
47	CYPRUS	0.69	0.27; 2.49; -2.07	INV	0.45	-0.42; 1.76; -0.89	INV
48	CZECH	0.16	-0.16; 3.88; -3.56	INV	0.88	1.88; -1.41; 0.41	YES
49	DENMARK	0.80	-0.11; 0.41; 0.50	INCR	0.64	-0.03; 4.65; -3.98	INV
50	DJIBOUTI	0.17	1.23; 0.24; -1.30	DECR	0.36	0.64; 0.41; -0.69	DECR
51	DOMINICAN	1.02	1.05; -0.31; 0.28	YES	1.57	0.32; -0.06; 1.31	YES
52	DOMINICA	7.75	2.00; -4.00; 9.75	YES	0.67	-0.36; 0.72; 0.31	INV
53	ECUADOR	1.46	-0.47; 1.06; 0.87	INV	1.14	0.73; -0.14; 0.55	YES
54	EGYPT	0.84	0.30; 0.37; 0.17	INV	0.51	11.99; -3.76; -7.72	DECR
55	EL SALVADOR	1.70	-0.20; 0.59; 1.31	INCR	0.66	-0.76; -14.49; 15.91	YES
56	EQUITORIAL G.	0.38	0.85; -0.20; -0.27	DECR	1.48	0.81; -0.66; 1.33	YES
57	ERITREA	1.18	1.44; -0.05; -0.21	DECR	0.80	1.02; 0.20; -0.42	DECR
58	ESTONIA	0.87	1.96; 0.82; -1.91	DECR	3.04	-0.70; -1.80; 5.54	YES
59	ESWATINI	0.94	1.41; -1.42; 0.95	YES	0.71	-0.02; 1.52; -0.79	INV
60	ETHIOPIA	0.80	-0.56; -1.45 ; 2.81	YES	1.24	0.34; 0.13; 0.77	YES
61	FIJI	2.00	0.00; 1.00; 1.00	INCR	0.50	0.75; -0.50; 0.25	YES
62 63	FINLAND FRANCE	1.14 1.17	0.91; -0.42; 0.65	YES YES	2.41 2.17	0.56; -2.38; 4.23	YES YES
63 64	GABON	0.97	0.82; 0.10; 0.25 0.20; 0.47; 0.30	INV	0.19	0.88; -0.86; 2.15 -0.51; 0.00; 0.70	INCR
65	GAMBIA	0.83	-0.25; 0.43; 0.65	INCR	0.19	-0.31; 0.00; 0.70 -0.38; 0.00; 0.75	INCR
66	GEORGIA	1.23	0.16; 0.43; 0.64	INCR	0.79	1.52; -0.49; -0.24	YES
67	GERMANY	0.73	0.15; -1.04 ; 1.62	YES	0.79	1.15; -0.56; 0.20	YES
68	GHANA	1.48	0.55; 0.70; 0.23	INV	0.62	0.13; -0.81; 1.30	YES
69	GREECE	0.71	0.33; -0.27; 0.65	YES	0.71	0.95; 0.28; -0.52	DECR
70	GRENADA	14.00	-5.00; 3.00; 16.00	INCR	0.10	-0.15; 0.00; 0.25	INCR
71	GUADELOUPE	1.35	0.00; 0.76; 0.59	INV	1.35	0.00; 0.76; 0.59	YES
72	GUATEMALA	0.25	2.01; -0.70; -1.06	YES	0.27	1.19; -0.11; -0.81	DECR
73	GUIANA FRENCH	0.88	1.30; -0.38; -0.04	YES	0.43	0.99; 0.27; -0.83	DECR
74	GUINEA	0.46	0.65; -0.56; 0.37	YES	1.68	0.21; 0.68; 0.79	INCR
75	GUINEA BISSAU	1.14	0.06; 1.59; -0.51	INV	4.20	-0.11; 0.04; 4.27	INCR
76	GUYANA	2.38	-3.45; -0.20; 6.03	INCR	4.23	-0.53; 0.58; 4.18	INCR
77	HAITI	0.60	0.30; -0.13; 0.43	YES	0.61	0.32; 0.42; -0.13	INV
78 70	HONDURAS	0.57	-2.94; 3.12; 0.39	INV	1.64	0.13; 0.54; 0.97	INCR
79	HONGKONG	0.04	0.95; -0.69; -0.22	YES	0.24	2.50; -8.79; 6.53	YES
80 81	HUNGARY ICELAND	0.90 2.28	0.66; -0.12; 0.36	YES INV	1.93 0.66	1.91; -2.72; 2.74	YES NO
82	INDIA	0.98	-0.85; 3.93; -0.80 1.82; 0.53; -1.37	DECR	0.66	0.84; 0.22; -0.40 1.08; -0.57; 0.45	YES
83	INDONESIA	0.98	0.67; 0.88; -0.60	INV	0.96	1.06; -0.03 ; $0.451.06$; -0.03 ; -0.03	YES
84	IRAN	1.04	1.73; -0.67; -0.02	YES	0.99	6.62; -6.62; 0.90	YES
85	IRAQ	0.77	0.15; -0.35; 0.96	YES	0.96	0.77; -0.40; 0.59	YES
86	IRELAND	2.16	-2.83; -5.64; 10.63	YES	1.12	1.12; -0.39; 0.39	YES
87	ISRAEL	0.21	-1.39; 1.08; 0.52	INV	1.16	-0.16; 0.44; 0.88	INCR
88	ITALY	1.04	2.24; -1.85; 0.65	YES	3.69	1.65; -7.89; 9.93	YES
89	JAMAICA	0.43	0.13; 0.06; 0.24	YES	2.47	-0.34; 2.06; 0.75	INV
90	JAPAN	1.02	0.69; 0.88; -0.55	INV	1.16	0.61; 0.42; 0.13	DECR
91	JORDAN	2.53	10.82; -18.20; 9.91	YES	0.93	1.28; 0.57; -0.92	DECR
92	KAZAKHSTAN	0.60	0.53; -5.45; 5.52	YES	2.06	-0.05; 2.37; -1.26	INV
93	KENYA	1.14	0.05; 0.65; 0.44	INV	1.18	0.47; 1.34; -0.63	INV
94	KOREA REP.	1.00	0.12; 0.87; 0.01	INV	1.04	0.60; -0.03; 0.47	YES
95	KOSOVO	1.02	1.00; 1.02; -1.00	INV	0.99	1.31; -0.29; -0.03	YES
96	KUWAIT	0.88	0.5; -0.34; 0.67	YES	1.10	0.58; -0.84; 1.36	YES
97	KYRGYZSTAN	0.17	-0.73; 0.26; 1.64	INCR	1.05	0.28; -0.32; 1.09	YES
98	LAO PDR	0.50	0.50; 0.50; -0.50	DECR	0.15	0.33; 0.74; -0.92	INV
99 100	LATVIA	0.74	1.97; -0.76; -0.47	YES	0.50	0.40; -0.22; 0.32	YES
100	LEBANON	1.03	0.57; 0.12; 0.34	YES	0.90	0.23; 0.06; 0.61	YES
101 102	LESOTHO	7.08	-2.86; 7.22; 2.72	INV VES	1.42	0.37; 1.51; -0.46	INV INV
102	LIBERIA LIBYA	0.31 0.96	0.18; -0.04; 0.17 0.19; -0.71; 1.48	YES YES	4.56 0.79	0.14; 4.61; -0.19 -0.42; 0.56; 0.65	INV INCR
103	LITHUANIA	0.96	0.56; 0.11; 0.16	YES	2.49	-0.42; 0.56; 0.65 -0.90; -0.52; 3.91	INCR
104	LUXEMBOURG	0.33	-8.55; -3.75; 12.54	INCR	1.48	1.16; -0.91; 1.23	YES
	LCALINDOUNG	U.4T	0.00, 0.70, 12.04	11 (CI)	1.10	1.10, 0.71, 1.40	110

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 Table A1. Cont.

No	All Countries Country Name	\mathbf{R}_{0}	First Wa R _{j′} s	ve U-Shape	R_0	Second Wav R _j 's	re U-Shape
106	MACAO	0.29	1.14; 2.29; -3.14	INV	-	-	-
107	MADAGASCAR	0.94	0.61; -0.16; 0.49	YES	0.75	0.38; -1.54; 1.91	YES
108	MALAWI	1.12	-0.23; 0.53; 0.82	INCR	6.46	-0.41; 0.99; 5.88	INCR
109	MALAYSIA	1.25	0.38; 2.79; -1.92	INV	1.30	-0.57; 1.82; 0.05	INV
110	MALDIVES	0.83	0.60; -0.53; 0.76	YES	1.05	-0.27; 0.70; 0.62	INV
111	MALI	0.64	0.59; 0.42; -0.37	DECR	7.78	-2.64; -4.96; 15.38	YES
112	MALTA	1.06	1.15; 0.24; -0.33	DECR	0.99	-0.73; 1.81; -0.09	INV
113	MAURITANIA	1.76	-0.94; 0.29; 2.41	INCR	1.14	0.73; -0.41; 0.82	YES
114	MAURITIUS	4.49	-4.05; 0.36; 8.18	INCR	0.35	1.41; 0.53; -1.59	DECR
115	MAYOTTE	5.46	-9.46; -2.50; 17.42	INCR	1.05	0.72; -0.17; 0.50	YES
116	MEXICO	0.86	-1.39; 3.07; -0.82	INV	2.53	-0.55; 0.10; 2.98	INCR
117	MOLDOVA	1.03	2.73; -0.67; -1.03	DECR	0.36	1.27; 0.66; -1.57	DECR
118	MONACO	3.15	0.52; -1.93; 4.56	YES	0.54	1.02; -0.12; -0.36	DECR
119	MONGOLIA	10.25	1.25; 19.25; -10.25	INV	0.68	0.91; 0.25; -0.48	DECR
120	MONTENEGRO	1.37	2.94; -3.90; 2.33	YES	0.66	2.36; 0.26; -1.96	DECR
121	MOROCCO	0.90	0.36; 1.41; -0.87	INV	0.95	0.95; -0.15; 0.15	YES
122	MOZAMBIQUE	0.72	0.92; 0.001; -0.20	DECR	0.70	2.46; -2.45; 0.69	YES
123	MYANMAR	1.12	-0.75; 1.07; 0.80	INV	1.15	-1.36; -2.17; 4.68	YES
124	NAMIBIA	0.68	1.37; -1.82; 1.13	YES	1.22	-0.26; 0.95; 0.53	INV
125	NEPAL NETHERLAND	0.74	0.35; 0.76; -0.37	INV YES	0.78 1.04	0.11; 0.58; 0.09	INV YES
126 127	NEW CALEDONIA	1.19 5.00	0.11; 0.11; 0.97 -2.00; 2.00; 5.00		1.04	1.05; -0.99; 0.98 1.00; -1.00; 1.00	YES
127	NEW CALEDONIA NEW ZEALAND	0.74	-2.00; 2.00; 5.00 2.30; -3.40; 1.84	YES YES	0.72	-0.52; 0.43; 0.81	INCR
129	NICARAGUA	0.74	-0.03; 0.97; 0.03	INV	1.02	0.86; 0.14; 0.02	DECR
130	NIGER	0.63	-0.03; 0.97 ; $0.030.28$; -0.12 ; 0.47	YES	2.21	-0.14; 0.39; 1.96	INCR
131	NIGER	1.13	0.16; 0.39; 0.58	INCR	1.02	1.38; -0.65; 0.29	YES
132	MACEDONIA	0.74	1.83; -1.16; 0.07	YES	0.74	1.26; -0.10; -0.42	DECR
133	NORWAY	0.77	-0.19; -0.61; 1.57	YES	2.13	6.02; -10.80; 6.91	YES
134	OMAN	3.70	0.39; 0.12; 3.19	YES	9.80	-16.87; 39.41; -12.74	INV
135	PAKISTAN	1.22	-0.61; 1.07; 0.76	INV	1.19	0.55; -0.11; 0.75	YES
136	PALESTINE	0.96	-0.18; -0.23 ; 1.37	YES	1.06	-0.21; 0.18; 1.09	INCR
137	PANAMA	0.96	0.16; 0.56; 0.24	INV	0.79	1.22; -0.16; -0.27	DECR
138	PAPAU NEW G.	0.49	0.35; -1.96; 2.10	YES	0.88	-0.39; 0.04; 1.23	INCR
139	PARAGUAY	0.59	-1.52; 1.90; 0.21	INV	1.20	-3.20;3.06; 1.34	INV
140	PERU	0.89	8.30; -2.47; -4.94	DECR	0.53	3.98; -4.72; 1.27	YES
141	PHILLIPPINES	1.15	0.89; -0.08; 0.34	YES	1.54	0.07; 2.84; -1.37	INV
142	POLAND	0.92	2.32; -1.89; 0.49	YES	1.31	1.71; -1.63; 1.23	YES
143	POLYNESIA	0.66	0.22; 0.20; 0.24	YES	0.21	-1.05; 1.09; 0.17	INV
144	PORTUGAL	1.56	-1.34; -8.29; 11.19	YES	3.89	1.13; -4.00; 6.76	YES
145	QATAR	0.80	-0.84; -1.99; 3.63	YES	1.03	0.62; 0.61; -0.20	INV
146	ROMANIA	0.88	0.90; 0.06; -0.08	DECR	0.95	1.23; -0.48; 0.20	YES
147	RUSSIA	1.07	1.16; -1.00; 0.91	YES	0.87	0.83; -5.77; 5.81	YES
148	RWANDA	1.80	3.20; 2.20; -3.60	DECR	0.14	3.93; -2.75; -1.04	YES
149	SAO TOME	1.44	0.44; 0.64; 0.36	INV	2.67	2.25; -3.45; 3.87	YES
150	SAN MARINO	5.10	0.28; 1.14;3.68	INCR	0.26	-0.05; 2.32; -2.01	INV
151	SAUDI ARABIA	0.90	-1.70; 2.94; -0.34	INV	0.98	-1.05; 0.54; 1.49	INCR
152	SENEGAL	0.72	-0.19; 1.48; -0.57	INV	1.59	0.73; 0.23; 0.63	YES
153	SERBIA	1.62	-0.40; 0.47; 1.55	INCR	0.82	2.02; -0.94; -0.26	YES
154	SEYCHELLES	0.48	0.30; 0.51; -0.33	INV	0.54	0.38; -0.19; 0.35	YES
155 156	SIERRA LEONE	2.23	-2.93; -0.80; 5.96	INCR	1.37 2.83	0.95; -1.25; 1.67 1.61; -2.44; 3.66	YES YES
156 157	SINGAPORE	1.33 0.99	1.15; 0.51; -0.33 -2.67; 1.90; 1.76	DECR INV	2.83 0.74	1.61; -2.44; 3.66 0.97; -0.73; 0.50	YES
157	SLOVAK SLOVENIA	0.99	-2.67; 1.90; 1.76 1.56; -0.71; -0.10	DECR	0.74 0.64	0.97; -0.73 ; $0.501.47$; -0.47 ; -0.36	YES
158	SOMALIA	1.18	-0.16; 1.51; -0.17	INV	0.64	0.86; 0.57; -1.14	DECR
160	SOUTH AFRICA	0.87	-0.16; 1.51 ; $-0.170.22$; 0.73 ; -0.08	INV	1.49	0.86; 0.57; -1.14 0.20; -0.04; 1.33	YES
161	SOUTH AFRICA SOUTH SUDAN	0.58	0.22; 0.73; -0.08	INV	1.49	0.20; -0.04; 1.33	YES
162	SPAIN	0.38	-0.18; 0.27; 0.29	INCR	0.51	1.21; -0.86; 0.16	YES
163	SRI LANKA	2.13	2.73; -0.75; 0.15	YES	0.51	0.42; 1.00; -0.63	INV
164	ST KITTS NEVIS	2.13	0.00; 1.00; 1.00	INCR	1.07	0.25; 0.18; 0.64	YES
165	ST LUCIA	1.13	-0.53; -0.04 ; 1.70	INCR	1.00	1.00; -1.00; 1.00	YES
166	ST VINCENT	0.04	-0.29; 0.24; 0.10	INV	0.69	-0.24; 0.35; 0.58	INCR
167	SUDAN	0.36	-1.46; 2.34; -0.52	INV	2.00	0.00; 2.00; 0.00	INV
168	SURINAME	10.34	2.70; 18.77; -11.13	INV	1.63	2.95; -1.25; -0.07	YES
169	SWEDEN	0.56	0.58; -1.20; 1.18	YES	1.21	0.67; -0.91; 1.45	YES
170	SWITZERLAND	1.21	1.25; 0.13; -0.17	DECR	0.28	0.89; 1.18; -1.79	INV
171	SYRIA	1.43	1.39; 4.13; -4.09	INV	0.18	0.31; -0.68; 0.55	YES
172	TAIWAN	1.88	-0.13; 1.38; 0.63	INV	0.66	-5.21; 13.83; -7.96	INV
173	TAJIKISTAN	1.02	0.71; -0.60; 0.91	YES	1.49	1.83; -0.17; -0.17	YES
174	TANZANIA	0.91	-1.50; 0.18; 2.23	INCR	1.89	3.42; 14.26; -15.79	INV

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All Countries			First Wa	ve		Second Wave			
No	Country Name	R_0	$\mathbf{R}_{\mathbf{j'}}\mathbf{s}$	U-Shape	R_0	R_j 's	U-Shape		
175	THAILAND	0.69	0.42; 0.07; 0.20	YES	2.71	-1.77; -0.75; 5.23	INCR		
176	TIMOR LESTE	5.00	1.00; 0.00; 4.00	YES	1.33	0.00; 1.00; 0.33	INV		
177	TOGO	0.08	6.05; -6.18; 0.21	YES	1.14	0.18; 0.09; 0.87	YES		
178	TRINIDAD	0.32	-0.26; 1.46; -0.88	INV	0.55	0.26; 0.03; 0.26	YES		
179	TUNISIA	1.53	0.77; -0.04; 0.80	YES	2.77	-3.21; -2.41; 8.39	INCR		
180	TURKEY	1.15	-1.50; -1.13; 3.78	INCR	2.21	19.82; -47.90; 30.29	YES		
181	UAE	0.97	2.07; -1.11; 0.01	YES	1.15	1.25; -0.64; 0.54	YES		
182	UGANDA	0.95	0.87; -0.37; 0.45	YES	0.64	0.44; -0.06; 0.26	YES		
183	UKRAINE	0.96	1.35; -1.04; 0.65	YES	0.30	3.10; 1.07; -1.73	DECR		
184	UK	0.76	-0.02; -0.76; 1.54	YES	1.03	0.43; 0.82; -0.22	INV		
185	USA	8.42	31.42; -99.18; 76.18	YES	0.49	3.32; -0.38; -2.45	DECR		
186	URUGUAY	0.63	0.71; 0.31; -0.39	DECR	1.03	-0.23; 0.35; 0.91	INCR		
187	UZBEKISTAN	0.95	0.04; 0.10; 0.81	INCR	0.90	-0.03; -0.39; 1.32	YES		
188	VENEZUELA	1.54	1.65; 2.95; -3.06	INV	0.82	1.09; -2.53; 2.26	YES		
189	VIETNAM	3.29	-0.84; -0.39; 4.52	YES	1.43	0.76; -0.11; 0.78	YES		
190	VIRGIN ISLANDS	0.51	0.01; -0.06; 0.56	YES	0.33	0.44; -0.22; 0.11	YES		
191	WEST GAZA	1.00	-1.00; -2.00; 4.00	YES	0.98	0.59; -0.11; 0.50	YES		
192	YEMEN	0.70	-0.34; 0.17; 0.86	INCR	1.50	1.00; 0.00; 0.50	YES		
193	ZAMBIA	0.75	0.25; -0.13; 0.63	YES	1.12	1.11; -0.44; 0.45	YES		
194	ZIMBABWE	1.44	0.24; 0.60; 0.60	INCR	1.62	1.08; -1.12; 1.66	YES		

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