Anticipating the emergence of infectious diseases

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In this paper, we present candidate approaches for anticipating disease emergence prior to large-scale outbreaks. Through use of ideas from the theories of dynamical systems and stochastic processes, we develop approaches which are not specific to a particular disease system or model but instead have general applicability. The indicators of disease emergence detailed in this paper can be classified into two parallel approaches: as early-warning signals based around the theory of critical slowing down and a likelihood-based approach. To test the reliability of these two approaches, we contrast theoretical predictions with simulated data. We find good support for our methods across a range of different model structures and parameter values.

1. Introduction

Tipping points—where small changes in circumstances precipitate dramatic shifts in state—are a feature of many natural systems. These sudden transitions can have devastating consequences, for example, irreversible climate change [1] or ecological collapse [2]. Forewarning, with sufficient time to act, is of clear importance.

In principle, careful scientific research and detailed mechanistic understanding could lead to the formulation of predictive models capable of anticipating such transitions. However, the combination of nonlinearity, non-stationarity, noise and data availability is a barrier to mechanistic modelling for many natural systems. There has been a trend towards looking for alternative methods of anticipating transitions, which do not rely on an empirically validated model [3]. One promising avenue is the development of early-warning signals (EWS) [4], which are summary statistics derived from dynamical systems theory and are calculable directly from observed data.

Dynamical systems theory states that as a stable system approaches a tipping point, the time taken to recover after a perturbation increases; ultimately diverging at the transition [4]. This phenomenon, known as critical slowing down, is expected to be observable in systems which are continually subject to shocks—whose origin can be either extrinsic (for instance, environmental fluctuations) or intrinsic (such as demographic noise). The effects of critical slowing down are manifest in the time series, leading to systematic changes in the summary statistics. For example, due to the persistence of perturbations the time series becomes increasingly correlated, which can be measured by the autocorrelation and correlation time. Because EWS are informed by generic properties derived from dynamical systems theory, they have a key advantage over model fitting approaches: detailed calibration is not required. EWS have been applied to anticipate transitions in a range of complex systems, including ecosystem collapse in lakes and microbial communities [5,6], desertification in arid ecosystems [7] and changes in climate [8].

Identifying reliable warning signals is especially timely for emerging infectious diseases. Examples of pathogens with pandemic potential include influenza virus [9] and SARS-like coronaviruses [10]. Similarly, pathogens which have been successfully controlled can re-emerge, for example, pertussis [11].
Constructing validated models for emerging diseases is complicated in part by complexity, and also by virtue of the pathogen’s novelty in a new host. Key epidemiological determinants, such as the infectious period, the mode of transmission, infectiousness and rate of spillover, are typically unknown or poorly quantified. This makes them a prime candidate for model-free approaches, such as EWS. To date, work on anticipating disease emergence has predominantly focused on specific systems and models, for instance, in studies on the effects of climate change on disease emergence [12], with a previous attempt at applying EWS to simulated data finding the approach to emergence difficult to detect [13].

The scenario considered in this paper is based on the slow emergence of a pathogen. This can occur for a variety of reasons: as a result of changing immunological landscape (increasing the susceptible pool—cf. monkeypox [14] or measles [15]), pathogen adaptation to new host [16], immune evasion [17,18] or long-term transient dynamics (the ‘honeymoon effect’ [19]). These mechanisms all lead to increasing inter-individual transmission of the pathogen—characterized by an increase in $R_0$, a threshold quantity. The primary concern for disease containment is averting the tipping point which occurs when $R_0 = 1$.

Formally, $R_0$ is defined as the average number of secondary infections caused by an infectious individual [20]. From theory we know that if $R_0 < 1$ transmission is subcritical and the pathogen is incapable of long-term persistence.

Instead, continued circulation can only be sustained through demographic. Additionally, we comment on how the transmission dynamics can be well approximated as a supercritical. It is this transition, from subcritical to supercritical, that we seek to anticipate.

The birth–death–immigration (BDI) process can be viewed as the simplest possible model of emerging diseases with direct transmission, modelling spillover of a pathogen from an external reservoir. In this regime, the transmission dynamics are characterized by what are known as ‘stuttering chains’ [21]. On the other hand, if $R_0 > 1$ the disease will typically invade successfully and grow in a fully susceptible population, leading to a large-scale outbreak and possible endemicity [22]. Such dynamics are referred to as supercritical.

As they are derived from a very generic model of disease emergence, the approaches are expected to have applicability across directly transmitted emerging disease systems. We test the robustness of measures to changes in model structure, simulating disease emergence using the SIS and SIRS compartmental models subject to varying immunology and demographics. Additionally, we comment on how the speed at which the transition is crossed affects the reliability of the two approaches, something which has been considered poorly understood in previous work on EWS [24].

2. Early-warning signals using the birth–death–immigration process

Previous work exploring EWS for infectious disease dynamics has shown that performance depends on the direction the critical transition is traversed, with the two contrasting cases shown in figure 1 [13]. In studies employing a combination of analytical results and simulated data, EWS have successfully been identified for diseases approaching elimination—for instance, due to increasing vaccine uptake—where $R_0$ approaches the critical transition from above [13,27]. The case of disease emergence has thus far proven more challenging [13].

This challenge can be understood from a theoretical perspective. The conventional theory of EWS, as detailed, for example, by Scheffer et al. [4] and used by O’Regan & Drake [13], relies on the presence of critical slowing down, presupposing that the system is at a stable fixed point and subject to exponentially decaying perturbations. When the number of infectious individuals is sufficiently large the transmission dynamics can be well approximated as a linearized system subject to Gaussian noise [28,29].

For endemic diseases prior to elimination the number of infected individuals is large enough for this approximation to be valid, and EWS perform well [13]. Conversely, for subcritical emerging diseases there are typically very few infected individuals present in the population. This means that the effects of demographic stochasticity feature strongly in the dynamics [29]. The stochastic dynamics are highly non-Gaussian and the conventional theory of critical slowing down is invalid. In the light of these findings, we instead develop EWS starting from an alternative model, the BDI process, which explicitly incorporates transmission between individuals. By starting from a model which captures the conditions preceding disease
emergence, we show how EWS can be expected to behave for this class of transition, and assess their prospects as generic indicators.

The BDI process is one of the simplest models of a subcritical disease [23]. It neglects the effects of susceptible depletion, assuming that only a small fraction of the population are infected at any one time, and that the pool of susceptibles is replenished with sufficient speed. The BDI process focuses on three simple transitions: (i) an infected individual is introduced into the population, (ii) the infection can spread to other individuals with a rate proportional to the number of infected individuals present, and (iii) infected individuals may recover from the disease.

We assume that at a constant rate $\zeta$ individuals import the infection due to contact with external sources, for example a zoonotic reservoir. We focus on diseases which are emerging due to changes in the transmission rate of the disease, $\beta(t)$. If there are $n$ infected individuals present in the population, then the total force of infection is $\lambda(t) = \beta(t) n + \zeta$, and the rate of recovery is $\gamma n$. Throughout the paper we assume that changes in $\beta(t)$ occur on a much slower time scale than the typical duration of an infection, $1/\gamma$.

The transition rate $T_{n,m}$ is defined as the probability per unit time of transitioning from a state with $n$ infected individuals to a state with $m$ infected individuals. The BDI process is a one-step stochastic process which consists of two possible transitions; for infectious diseases these are infection and recovery. Respectively, the transition rates for infection and recovery are

\[ T_{n+1,m} = (\beta n + \zeta) \]
\[ T_{n-1,m} = \gamma n. \]

(2.1)

In the BDI process, the chain of transmission stemming from a particular introduced index case is given by a branching process [30]. A particular outbreak can be considered as a superposition of the separate chains of transmission caused by each introduced case during the outbreak [31]. The basic reproductive number is defined as the average number of secondary cases, $R_0 = \beta/\gamma$, found using the offspring distribution of the associated branching process [30,32].

The probability of $n$ individuals being infected at time $t$ is $P_n(t)$. The change in probability with time is found by solving the master equation, a set of coupled linear differential equations built from the transition rates. For the BDI process,

\[
\frac{dP_n(t)}{dt} = T_{n-1,n}P_{n-1}(t) + T_{n+1,n}P_{n+1}(t) - [T_{n+1,n} + T_{n-1,n}]P_n(t) \quad \forall n > 0,
\]

(2.2)

\[
\frac{dP_0(t)}{dt} = T_{0,1}P_1(t) - T_{1,0}P_0(t).
\]

The advantage in considering such a simple model is that the master equation, which determines how the probability distribution of the number of infectious individuals changes in time, can be solved exactly without the need for any approximations [23]. This can be achieved through use of the moment generating function, $Z(\psi, t) = \mathbb{E}[e^{\psi n}^t]$. The variable $\psi$ is used to find the moments and correlation functions of the stochastic process via differentiation, for example, the $i$th moment, $\mu_i = \mathbb{E}[n^i] = \partial^i Z/\partial \psi^i \big|_{\psi=0}$. We find from the master equation that $Z(\psi, t)$ satisfies the partial differential equation

\[
\frac{\partial Z}{\partial t} = (e^\psi - 1) \left( \beta \frac{\partial}{\partial \psi} + \zeta \right) Z + (e^{-\psi} - 1) \gamma \left( \frac{\partial Z}{\partial \psi} \right)^2 \tag{2.3}
\]

Further details of the master equation calculation are presented in appendix A.

Using that $R_0 = \beta/\gamma$, the mean-field behaviour of the system solves $d\mu_i/dt = (R_0 - 1)\gamma \mu_i + \zeta$. If $R_0 > 1$ then $\mu_1$ grows exponentially, whereas if $R_0 < 1$ the disease persists at a low level, $\mu_1 = (\zeta/\gamma)/(1 - R_0)$, sustained by immigration.

Table 1 shows the list of candidate EWS calculated from the moment generating function of the stationary BDI process which will be considered in this paper. The mean, variance, index of dispersion (also known as the variance to mean ratio), and correlation time all diverge as $R_0 \rightarrow 1$. The autocorrelation approaches one, whereas the coefficient of variation remains constant below the transition.

The correlation time gives a measure of the persistence of correlations. Its divergence as $R_0 \rightarrow 1$ implies that correlations persist for extremely long times, and is the mathematical definition of critical slowing down. Intuitively, critical slowing down can be understood by considering the lengths of transmission chains. Although on average each imported case will infect fewer than one individual (as $R_0 < 1$), there is a possibility that a large chain of secondary cases is sparked, which then takes a long time to die out. As $R_0$ increases, the probability of larger chains of transmission increases, in turn leading to an increasingly autocorrelated dynamics. The increased probability of larger chains of transmission also causes an increase in the mean and variance of the number of infected.

In addition to the EWS derived from the BDI process presented in table 1 we also study two signals from information theory: the Kolmogorov complexity, and the Shannon entropy [33]. The Kolmogorov complexity is a measure whose origin lies in the length of the minimal computer program needed to reproduce the sequence of data, with longer computer programs implying more complex data [33–35]. The increased correlation near the transition may therefore result in a decrease in complexity, which can serve as an EWS. The Shannon entropy is a measure of the information content of the time series. As $R_0$ approaches one, the probability of rare large outbreaks increases, which suggests the entropy will also increase near the transition. For details of how these two measures are computed, see appendix B.

<table>
<thead>
<tr>
<th>early-warning signal</th>
<th>formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>mean</td>
<td>$\mu_1 = \frac{\zeta}{\gamma} + \frac{\beta \psi}{\gamma}$</td>
</tr>
<tr>
<td>variance</td>
<td>$\sigma^2 = \frac{1}{\gamma}$</td>
</tr>
<tr>
<td>coefficient of variation</td>
<td>$\sigma/\mu_1 = \frac{1}{1 - R_0}$</td>
</tr>
<tr>
<td>index of dispersion</td>
<td>$\frac{\sigma^2}{\mu_1} = \frac{1}{1 - R_0}$</td>
</tr>
<tr>
<td>correlation time</td>
<td>$\tau = \frac{1}{1 - R_0}$</td>
</tr>
<tr>
<td>autocorrelation</td>
<td>$AC(\tau) = \exp(-\tau/\tau)$</td>
</tr>
</tbody>
</table>

Table 1. Measurable quantities calculated from the BDI process. All expressions are valid for $0 \leq R_0 < 1$. See equations (A 3), (A 4) and (A 7) for details.
Two simplifying assumptions are implicit in the model. Firstly, the BDI process neglects susceptible depletion. Secondly, the speed with which $b$ changes is ignored, assuming that it varies infinitely slowly. The effects of relaxing the first of these assumptions will be investigated in the next section. Disease emergence over faster time scales will be investigated in §5.

3. Comparison with models featuring susceptible depletion

For EWS to be useful they must be robust to changes in model structure, and not just specific to the BDI process. Using results from simulations of common disease models, in this section we explore the sensitivity of the EWS to a range of demographic and immunological properties.

The SIS model is a model of infectious disease spread in which individuals move from the susceptible class to the infected class upon infection, and then return to the susceptible class when they cease being infectious. The total population size is assumed constant. The SIRS model is similar to the SIS model, but in addition the pathogen confers temporary immunity to reinfection, which wanes at rate $r$ [20]. The dynamics of both models can be represented using a master equation, as with the BDI process. Exact numerical realizations of these models can be obtained using the next reaction method (NRM) algorithm [25, 26].

The candidate EWS for the SIS and SIRS models are shown in figure 2. The results for the BDI process listed in table 1 are also plotted.

![Figure 2. EWS for the stationary SIS and SIRS models. Red lines correspond to the theoretical results found from the BDI model (table 1). Symbols depict the results of simulations with the duration of immunity as indicated in the figure. Independent simulations are performed for each value of $R_0$. EWS are calculated by time averaging over $4 \times 10^3$ weeks of data (see appendix D). The force of infection in the SIS and SIRS models is $\lambda = (\beta n + \zeta)/N$, where $n$ is the number of infected and $N$ the total population size. The remaining model parameters are $\gamma = 1$ and $\zeta = 1$ week$^{-1}$. The autocorrelation is calculated at lag one week, $AC(1)$. (Online version in colour.)](#)
is to decrease the responsiveness of a signal to increases in $R_0$. The mean and variance always increase prior to the transition; however, the magnitude of the increase is reduced. The value of $R_0$ for which the variance is maximized is seen to increase. A similar effect is also observed for the index of dispersion, autocorrelation and correlation time.

Note that there is a large distinction in behaviour between models above the transition; for instance the coefficient of variation strictly decreases for the SIS model, whereas for the SIRS model it may decrease, increase or remain approximately flat depending on the model parameters. The generality of the EWS is strictly for disease emergence.

4. Cox’s $\delta$: a likelihood-ratio test

An alternative model-specific approach to detecting critical transitions has been proposed by Boettiger and Hastings, focusing on anticipating the loss of stability of fixed points [24]. They propose likelihood-ratio testing, using a statistic they refer to as Cox’s $\delta$. The test determines whether the dynamics are approaching a transition or are instead stationary. Each hypothesis is represented by a model, and a maximum-likelihood estimate (MLE) for the data is found. The value of Cox’s $\delta$ provides a measure of how much disease emergence is favoured. The procedure by which Cox’s $\delta$ is calculated also allows for an estimation of the time remaining until the transition is reached.

Boettiger and Hastings assume that the Ornstein–Uhlenbeck (OU) process is an appropriate model for the dynamics. The OU process is a continuous-time stochastic process in which there is deterministic reversion to the mean, with Gaussian white noise perturbing the system [28]. It has a known mathematical expression for the likelihood of a time series, which Boettiger and Hastings use to efficiently calculate Cox’s $\delta$ and thereby determine whether the system is approaching a transition [24].

A barrier to using Cox’s $\delta$ as a method for anticipating disease emergence is this reliance on the OU process to calculate the likelihood. Dynamically, the OU process does not constrain the number of infected individuals to be a non-negative integer. When there are a small number of infected individuals present in the data, this means biologically impossible paths are included in the likelihood calculation. The problem can be overcome by instead using the BDI process, which presents a more biologically plausible model of disease emergence.

An exact solution to the transition probability exists for the BDI process from which the likelihood of a time series can be constructed (see appendix C). We assume $R_0(t)$ can be decomposed into two parts, a baseline which is set as $R_0(0)$ and a linear trend, $\Delta R_0$, i.e.

$$R_0(t) = R_0(0) + \Delta R_0 t. \quad (4.1)$$

For the test hypothesis (the disease is emerging), the MLE is calculated from a likelihood surface allowing both of these parameters to vary. In calculating the MLE for the null hypothesis (no emergence), we fix $\Delta R_0 = 0$, still allowing $R_0(0)$ to vary. For an observed time series $\{n_t\}_{t=0}^T$ the log-likelihood of the MLE for the test model is

$$L_{\text{Test}} = \max_{R_0(0),\Delta R_0} \sum_{t=0}^{T-\delta} \ln P_{\text{BDI}}(n_{t+\delta} | n_t, R_0(t)) \quad (4.2)$$

and for the null model is

$$L_{\text{Null}} = \max_{R_0(0)} \sum_{t=0}^{T-\delta} \ln P_{\text{BDI}}(n_{t+\delta} | n_t, R_0(0)). \quad (4.3)$$

The expressions for $P_{\text{BDI}}$ are presented in appendix C. Cox’s $\delta$ statistic is defined as twice the difference in the log-likelihoods of the MLEs for the two nested models,

$$\delta = 2(\ln L_{\text{Test}} - \ln L_{\text{Null}}). \quad (4.4)$$

Because the models are nested $\delta \geq 0$. Increasingly positive $\delta$ implies stronger support for the test model over the null model. A $\delta > 2$ is taken as the criterion for concluding that the disease is emerging, on the basis of the AIC score [24].
100 grid of uniformly spaced data points with linearly from 0 to 1.2 over 12 years. Data are output weekly. The likelihood surface is calculated using only data from the first 6 years for parameters on a 100 × 100 grid of uniformly spaced data points with ΔR₀(0) ∈ [0, 0.008] and R₀(0) ∈ [0, 0.8]. The gold star indicates the location of the true parameter set and the red dot indicates the parameters corresponding to the maximum-likelihood estimate (MLE). Simulation performed using the modified next reaction method [26]. The remaining model parameters are fixed to their true values to reduce computation time; 1/μ = 1 week, ζ = 1 week⁻¹ and N = 10⁴ individuals. The MLE is that R₀(0) = 0.00 and ΔR₀ = 1.83 × 10⁻³ week⁻¹. The true values are R₀(0) = 0.00 and ΔR₀ = 1.92 × 10⁻³ week⁻¹. Side panels show the likelihood profiles. (Online version in colour.)

Figure 3. Estimating the rate of emergence using maximum-likelihood methods. Simulation generated using an SIS model with R₀(t) = R₀(0) + ΔR₀t increasing linearly from 0 to 1.2 over 12 years. Data are output weekly. The likelihood surface is calculated using only data from the first 6 years for parameters on a 100 × 100 grid of uniformly spaced data points with ΔR₀(0) ∈ [0, 0.008] and R₀(0) ∈ [0, 0.8]. The gold star indicates the location of the true parameter set and the red dot indicates the parameters corresponding to the maximum-likelihood estimate (MLE). Simulation performed using the modified next reaction method [26]. The remaining model parameters are fixed to their true values to reduce computation time; 1/μ = 1 week, ζ = 1 week⁻¹ and N = 10⁴ individuals. The MLE is that R₀(0) = 0.00 and ΔR₀ = 1.83 × 10⁻³ week⁻¹. The true values are R₀(0) = 0.00 and ΔR₀ = 1.92 × 10⁻³ week⁻¹. Side panels show the likelihood profiles. (Online version in colour.)

Figure 3 shows the log-likelihood surface calculated using the BDI process for a time series generated using the SIS model. The MLE of the test model is R₀(0) = 0.00 and ΔR₀ = 1.83 × 10⁻³ week⁻¹. The critical transition is predicted to occur after 10.52 years, very close to the true value of 10 years. Cox’s δ statistic is 10.78, implying that the hypothesis of disease emergence is statistically supported [24]. One caveat is that this strong result has been achieved under idealized model conditions, assuming knowledge of the rates of immigration and recovery, and calculated using a long sequence of data (312 weeks of data). In the following section, we explore further the reliability of Cox’s δ at anticipating transitions and compare with the EWS.

5. Impact of time scale on reliability of predictors

In §3, we presented results for stationary dynamics, appropriate for a disease emerging over an extremely long time scale. In practice, we are interested in anticipating disease emergence over faster time scales, as shown in figure 3. In this section, we investigate how the EWS and Cox’s δ perform as indicators of disease emergence. Results for the EWS are shown in figures 4 and 5, and results for Cox’s δ are shown in figures 6, 7 and 8.

Figure 4 shows the EWS calculated for the SIS model, with the stationary BDI process results also shown for comparison. The EWS are calculated from the time-series data; for simplicity, we use unweighted moving averages (see appendix D). The time series were generated by an SIS model with linearly increasing R₀; from 0 to 2 over T = 10, 20, 30 and 40 years. The moving window average is calculated using three window sizes: W = 100, 200 and 300 weeks.

As shown in figure 4, all EWS bar one have a clear response to the approaching transition as long as the time scale is longer than 10 years (T > 10). For instance, the variance increases for all window sizes and speeds prior to R₀ = 1, in line with predictions. The exception is the coefficient of variation, which performs poorly as an indicator of transitions, remaining largely unchanged until R₀ > 1. It is also highly sensitive to the choice of window size, the sharp peak following the transition observed for T = 10 and 20 years increases with window size. The autocorrelation performs particularly well, reaching AC(1) > 0.85 for T = 20, 30 and 40 years. It does not appear particularly sensitive to window size below the transition.

As the speed of emergence is increased, the lag behind the BDI process results increases. Overall, we see that the transition speed has three notable effects which adversely impact all EWS. The first effect is dynamical: the incidence
level is continually responding to the changing equilibrium resulting from increasing $R_0$, leading to a lag between the values of the EWS calculated at equilibrium and those observed in the time series. Combined with demographic stochasticity, this leads to an apparent ‘bifurcation delay’ as the transition is traversed [36,37].

The second effect arises in computing the EWS. The theoretical results for the EWS are calculated by finding the moments and autocorrelation of a stochastic process. Owing to the absence of multiple replications, these are calculated over a moving window [4]. However, when the disease emerges over a finite time scale, the process is non-stationary, and therefore also non-ergodic, due to changing $R_0(t)$. In practice, this means that averages are calculated using data points which were generated at different values of $R_0$; for example, if $T = 10$ years and $W = 100$ weeks, the difference in $R_0$ between the start and end of the window is 0.38. This reduces the responsiveness of the EWS, also contributing to the lag behind the BDI process results. Furthermore, there can be other artifacts arising from non-ergodicity, for instance, increased variance of the data. Increasing both the window size and transition speed exacerbates these effects due to the larger range in values of $R_0$.

Thirdly, there is Monte Carlo error due to the limited number of data points used. This is manifest in the stochastic variation of the observed EWS as $R_0$ changes. This is particularly apparent in the first column of figure 4, where a chance drop in prevalence immediately prior to $R_0 = 1$ leads to a failure of most EWS to behave as theory predicts. Increasing the size of the window has the effect of reducing this error (evidenced by the smoother curves for $W = 300$), however, at the expense of the signal’s responsiveness, as detailed above.

Given the existence of Monte Carlo error, to quantify the performance of the EWS further, we compute 90% CIs for the EWS for disease emergence over 20 years and over 100 years using a window size of 52 weeks (figure 5a). A measure of how effective an EWS is at detecting emergence is the range in $R_0$ over which a specific reading of an EWS lies within the 90% CI. A wide range means that the EWS gives little information about the value of $R_0$. For the variance, a
A reading of \( \sigma^2 = 40 \) falls within the 90% CI for \( 0.8 \leq R_0 \leq 1.1 \), implying a strong chance that the system is near the transition. A reading of \( K = 0.8 \) for the Kolmogorov complexity falls within the 90% CI for all values of \( R_0 \) shown, implying a poor predictive power.

To quantify this further, figure 5b shows an estimate of the probability that \( R_0 > 1 \), given an observed value of an EWS. The noisiness in the estimate for more extreme values of the EWS is due to the infrequency of these events. (Online version in colour.)

![Figure 5](http://rsif.royalsocietypublishing.org/) (a) 95% CIs of EWS for the SIS model with \( N = 1000 \). \( R_0 \) increases linearly from 0 to 2. The total simulation time is \( T = 20 \) years and \( T = 100 \) years. Data are output weekly, EWS are calculated using a window size \( W = 52 \) weeks for both columns. The average of the replicates is shown in black and a sample replicate is shown in dark blue. (b) Numerical estimate of the probability that \( R_0 > 1 \), given an observed value of an EWS. The noisiness in the estimate for more extreme values of the EWS is due to the infrequency of these events. (Online version in colour.)

reading of \( \sigma^2 = 40 \) falls within the 90% CI for \( 0.8 \leq R_0 \leq 1.1 \), implying a strong chance that the system is near the transition. A reading of \( K = 0.8 \) for the Kolmogorov complexity falls within the 90% CI for all values of \( R_0 \) shown, implying a poor predictive power.

To quantify this further, figure 5b shows an estimate of the probability that \( R_0 > 1 \) given a particular reading of an EWS, \( y \) (appendix E provides details on how this estimate is calculated). The sensitivity of \( P(R_0 > 1 \mid y) \) to changes in \( y \) depends heavily on the EWS. Large values for the mean, variance and entropy strongly indicate that transmission of the pathogen is supercritical. On the other hand, for EWS which peak near \( R_0 = 1 \), such as the index of dispersion, autocorrelation and correlation time, there is low indication whether the disease is supercritical or not, unless the observed value is especially large.

Because of their narrow confidence intervals, the variance and entropy are exquisitely sensitive, with \( P(R_0 > 1 \mid y) \) increasing from near 0 to 1 over a small range of \( y \). Others, such as the index of dispersion and autocorrelation, have an intermediary range of values of \( y \) where \( P(R_0 > 1 \mid y) \approx 0.5 \).

We see a trade-off between how reliable an EWS is at classifying whether \( R_0 > 1 \) and the range of values of \( R_0 \) over which \( P(R_0 > 1 \mid y) \) notably increases. A practical conclusion from this is that sequential readings of an EWS are needed to ascertain whether a disease is emerging.

Using a moving window, in figure 6 we show Cox’s \( \delta \) calculated for the same set of time series studied in figure 4. As the transition is approached Cox’s \( \delta \) typically increases in significance, with large responses for window sizes \( W = 200 \) and 300 weeks. There are spurious increases in Cox’s \( \delta \) for the stationary time series, due to the intrinsic variability of the dynamics, but these are dampened by using a larger window.

To test its reliability as an indicator of disease emergence, figure 7 shows the probability that \( \delta > 2 \) as the transition is approached.
Large window sizes lead to an improved performance, as does a faster speed of emergence. For faster speeds of emergence ($T < 20$ years) and larger windows ($W > 200$ weeks), Cox’s $\delta$ performs well, with $P(\delta > 2) > 0.75$ prior to the transition. From figure 7, the recommendation is to use the largest window size feasible given the data.

The effect of the speed of emergence and the window size on the performance of Cox’s $\delta$ is more subtle than for the EWS shown in figure 4. For EWS, the performance improves when the disease emerges at a slower rate due to the smaller change in $R_0$ between the start and end of the window. The opposite is the case for Cox’s $\delta$. Instead, slower emergence reduces the difference in likelihood between the emerging and non-emerging models, reducing the power of Cox’s $\delta$.

From the estimates of the baseline $R_0$ and $\Delta R_0$ the time to emergence can be estimated (figure 8). The mean estimated time to emergence generally does agree well with the true value, with typically marginally better agreement for smaller window sizes. However, the 95% CIs are large, in line with the variation in the MLE of $\Delta R_0$ seen in figure 6. Increasing the window size reduces the confidence intervals, with a more pronounced reduction in the probability of underestimating the time to emergence, cf. the difference between $W = 100$ and 300 weeks for $T = 20$ years. With sufficiently large window size given the speed of emergence, the estimated time to emergence can be taken as an overestimate.

### 6. Discussion/conclusion

This work is intended to provide a theoretical base for the development of methods to anticipate disease emergence. We have presented two parallel approaches by which this may be accomplished. The first approach is to extend the burgeoning literature on EWS, reformulating them in a manner appropriate for emerging diseases. The second approach makes use of Cox’s $\delta$, a likelihood-ratio test for emergence.

For both approaches, the application to emerging diseases is achieved through use of the BDI process, a well-understood model from the theory of stochastic processes.

Both the EWS and Cox’s $\delta$ perform best when calculated using a large number of data points as this reduces statistical uncertainty. The time scales imposed by real-world disease emergence likely mean that available data are more limited. We therefore apply both approaches to weekly time series data of diseases emerging over a range of different intervals.
We find that—apart from the coefficient of variation—all EWS undergo similar behaviour regardless of the time scale of emergence. More advanced time averaging methods may be needed for very fast emergence to detect a strong signal. Although larger window sizes do reduce uncertainty they are typically not necessary, with $W = 100$ weeks being sufficient. Cox’s $\delta$ performs best with the largest window size possible so that there is the largest possible difference in $R_0$ between the start and end of the window.

Overall we conclude that the two methods are complementary, performing best under differing conditions. Cox’s $\delta$ reliably detects emergence over fast time scales, where the null hypothesis is strongly disfavoured. EWS perform best at detecting long term trends in the time series, where there is a more gradual approach to the epidemic transition. Additionally, our results favour the use of some EWS over others. The behaviour of the coefficient of variation and Kolmogorov complexity prior to the transition means they both perform poorly as indicators of emergence. The remaining EWS (the mean, variance, index of dispersion, autocorrelation, correlation time and entropy) have a behaviour which is resilient to parameter changes and reliable in the face of stochasticity, making them strong candidate EWS.

By grounding our results in theory, rather than a detailed model for a particular disease we expect our results will have applicability to a wide range of emerging infectious diseases.

Figure 7. Probability that Cox’s $\delta > 2$, calculated from 200 replicates. Cox’s $\delta$ computed using the likelihood-ratio test and BDI process solution, see §4. Fixed model parameters are $N = 1000$, $\gamma = 1$ and $\zeta = 1$. (Online version in colour.)

Figure 8. Mean and 95% CIs of the remaining time to emergence. Time to emergence calculated using the MLE, $T_e = W + (1 - R_0(0))/\Delta R_0$, where $W$ is the window size. True value of the time to emergence shown in black. Calculated using the same data as in figure 7. (Online version in colour.)
This generality is of particular importance for emerging infectious diseases, where there are likely many unknowns regarding transmission of the pathogen.

The key simplifying assumption of the BDI process which makes it mathematically solvable is that it neglects the impact of disease transmission on the availability of susceptible individuals. In reality, infection diminishes the size of the susceptible pool and can also confer immunity to reinfection, delaying the individual’s return to susceptibility. Our results show how these demographic and immunological considerations impact on the performance of EWS. Although the signals’ strengths are reduced, for all parameters considered the performance is still informative for a population of 10^6 individuals.

Our work is not the first to investigate the signature of criticality prior to the epidemic transition; in particular, we note the work of Jansen and collaborators [15]. These previous works seek to identify the approach to the critical transition through changes in the tail of the outbreak size distribution [15,38]. In contrast, both approaches in this paper are based around summary statistics. This has advantages: the sizes of individual distinct outbreaks can be difficult to identify, a large number of outbreaks are required to accurately reconstruct the tail of the outbreak size distribution, and summary statistics calculated using moving windows clearly display temporal variation in the dynamics.

The SIS and SIRS models considered in this paper have only one susceptible and one infected class, assuming all individuals are equally infectious and interact homogeneously. Risk structure has been shown to significantly change the necessary conditions for epidemics, with R_0 = 1 being an underestimate for the epidemic threshold [39]. For diseases emerging in populations structured as many loosely connected smaller community units, such as Ebola in sub-Saharan Africa, this may significantly impact on predictability. R_0 is a measure based on the average context; however heterogeneities in the contact structure of individuals can lead to instances of superspreading [22], particularly relevant for sexually transmitted diseases and something not captured in the BDI process. The large variations in transmission rates reduce the worth of R_0 as a measure, and a universal increase in R_0 is unlikely to be the driver of emergence. Superspreading events have been identified for SARS and more recently for MERS [21,40]. In addition, there is the challenge of incomplete and unreliable data, modelled through explicit inclusion of a reporting process [41]. Further work is ongoing studying EWS and Cox’s δ for these dynamics.

This work lays theoretical foundations, but more research is needed to develop actionable technologies applicable to actual disease data. For instance, improved reporting would lead to an increase in the mean number of case reports, but not an increased autocorrelation. A simultaneous increase in both presents stronger evidence of disease emergence. The development of ways in which readings of multiple EWS and also Cox’s δ can be combined and leveraged to improve the quality of predictions is therefore desirable. One motivation in studying EWS is to ultimately develop comprehensive EWS software packages. These packages would be plug and play ready; public health practitioners could use them to give quantifiable information about the risk of a critical transition and disease emergence in a particular context.

As a final note, while this paper has focused on anticipating infectious disease emergence this does not limit the applicability of our methods. Emerging diseases are an example of an invasion process, where a new class of individual attempts to establish itself in a population. Other examples are found across the biological sciences. The BDI process is an appropriate model for a large range of invasion processes. We propose that the methods in this paper will have a similarly general status for anticipating successful invasion. Our work, therefore, opens up a new class of transitions which it may be possible to anticipate.

Appendix A. Critical slowing down for the stationary birth–death–immigration process

The master equation for the BDI process, equation (2.3), can be solved both when R_0, γ and ζ are constant or when they are functions of time [23]. For mathematical simplicity, we will focus on the case in which the disease is emerging over a long time scale relative to the infectious period. Under these conditions, we can find exact theoretical values for the EWS. If there are $y$ infectious individuals initially present then $Z(\psi, 0) = e^{\psi y}$ and equation (2.3) has the solution [23]

$$Z_\gamma(\psi, t) = \frac{(1 - R_0)^{\psi + 1/\gamma} (A(\psi) + B(\psi, t))^{\psi}}{(A(\psi) + R_0 B(\psi, t))^{1 + 1/\gamma}}, \quad \text{(A1)}$$

with $A(\psi) = (1 - R_0 e^{\psi})$ and $B(\psi, t) = e^{(R_0 - 1)e^{\psi} - 1}$.

We are interested in dynamics prior to the emergence of disease so restrict ourselves to considering $R_0 < 1$. As $\tau \to \infty$, we see that $B(\psi, t) \to 0$. The generating function converges onto the stationary solution

$$Z^* = \left(\frac{1 - R_0}{1 - R_0 e^{\psi}}\right)^{\psi/\gamma}, \quad \text{(A2)}$$

for any number of initial infected. This is the moment generating function of a negative binomial distribution with mean and variance

$$\mu_1 = \frac{\zeta / \gamma}{1 - R_0} \quad \text{(A3)}$$

and

$$\sigma^2 = \frac{(\zeta / \gamma)}{(1 - R_0)^2}. \quad \text{(A4)}$$

Both equation (A3) and equation (A 4) diverge as $R_0 \to 1$. Using equations (A 1) and (A 2), we can calculate the correlation function in the stationary state. The conditional mean, $E[P(\psi) | n(0)]$, is

$$\frac{\partial Z_{\psi}(0)}{\partial \psi} \bigg|_{\psi = 0} = \mu_1 (1 - e^{(R_0 - 1)\psi})$$

$$+ \eta(0) e^{(R_0 - 1)\psi}, \quad \text{(A5)}$$
and therefore the correlation function \( C(\tau) = \mathbb{E}[n(t)n(0)] \) is
\[
C(\tau) = \mathbb{E}[|n(t)| | n(0)|] = \sigma^2 e^{(\mu-1)\gamma t} + \mu^2.
\]
(A 6)
The autocorrelation, defined for a stationary process as
\( AC(\tau) = (C(\tau) - \mu^2)/\sigma^2\), is
\[
AC(\tau) = e^{-(1-R_0)\gamma t}.
\]
(A 7)
As the critical transition is approached, the correlation time \( \tau_c \) diverges. This is the hallmark of critical slowing down.

Appendix B. Shannon entropy and Kolmogorov complexity

The Shannon entropy is a measure of the information content of a random variable, and is defined as
\( H(t) = -\sum_i p_i(t) \log p_i(t) \) [33]. For time series data, an approximation to the probability distribution over \( n \) is calculated numerically via \( p_i(t) \approx \sum_{n=1}^W n_i(t)/W \), where \( n_i(t) \) is the number of infected individuals present at time \( s \) and \( n_i(t) \) is the indicator function, equal to one if \( n = n_i(t) \) and zero otherwise. As mentioned above, when the process is non-stationary this method is inexact.

Another information theoretic measure is the Kolmogorov complexity [33]. The Kolmogorov complexity of a sequence of data is the shortest computer algorithm required to reconstruct it. A quantification of the Kolmogorov complexity was proposed by Kaspar & Schuster [34]. Before applying their algorithm the data are converted to sequence of bits, 0 or 1. Their algorithm proceeds by considering two operations: ‘copy’ and ‘insert’. Each operation adds one to the complexity measure. The Kolmogorov complexity is the minimum number of operations required to reconstruct the data sequence. For further details, see [34]. The specific algorithm used to calculate the Kolmogorov complexity can be found in [35]. As with the other EWSs, we calculate the Kolmogorov complexity over a moving window. To convert the time series to a sequence of bits, we perform a linear detrending over the data within the window. The residual time series is discretized into 1 whenever the residual is larger than zero, and 0 otherwise.

Appendix C. Transition probability distribution and likelihood

The generating function given in equation (A 1) can also be used to find the probability of there being \( n_{t+\delta} \) infected at time \( t+\delta \) given \( n_t \) infected at time \( t \) [28]. Assuming that \( R_0 \) does not vary significantly between time \( t \) and \( t+\delta \), the transition probability for the BDI process is
\[
P_{BDI}(n_{t+\delta} | n_t, R_0) = \sum_{m=0}^{n_t+n_{\delta}} P_{BDI}(n_{t+\delta} - m | n_t, R_0)
\times P_{BDI}(m | 0, R_0),
\]
(C 1)
where
\[
P_{BDI}(m | n, R_0) = \sum_{k=0}^{\min(n, m)} \binom{n}{k} \binom{m-k}{n-k} R_0^k (1-R_0)^{m-k}
\]
and
\[
P_{BDI}(m | 0, R_0) = \left(\frac{m + \zeta + \gamma}{m}\right) R_0^m (1-R_0)^m
\]
with \( \rho = [1 \quad AC(\delta)]/[1 \quad R_0AC(\delta)] \) and \( \zeta = \mu_1/\mu_1 + \zeta/\gamma = 1/(2 \quad R_0) \). Given a time series \( \{n_t\}_{t=0}^T \), the likelihood of the data for a given rate of emergence, \( R_0(t) = R_0(0) + \Delta R_0 t \), is
\[
L(R_0(0), \Delta R_0) = \prod_{t=0}^{T-\delta} P_{BDI}(n_{t+\delta} | n_t, R_0(t)).
\]
(C 4)
The MLE for \( (R_0(0), \Delta R_0) \) is given by \( \arg \max_{R_0(0), \Delta R_0} L(R_0(0), \Delta R_0) \).

Appendix D. Calculating early-warning signals from single time-series data

If the BDI process is stationary then the process is ergodic, allowing expectation values to be calculated from a time series by time averaging. For example, the correlation function is
\[
C(\tau) = \mathbb{E}[n_{t+\delta} n_t]
\approx \frac{1}{T-\delta} \sum_{s=t}^{t+\delta} \frac{n_{s+\delta} n_s}{W}.
\]
(D 1)
The sum is performed over integer multiples of the timestep \( \delta \), with the approximation becoming exact as \( W \to \infty \). Although not formally equivalent, and therefore potentially introducing errors (see the discussion in §5), we will also use this approach to calculate expectation values for non-stationary processes, in line with the early-warnings literature.

Appendix E. Estimating the probability of supercriticality

Suppose we observe that EWS \( \Theta \) has a value \( y \). We would like to know the probability that \( R_0 > 1 \) given this observation, \( P(R_0 > 1 | \Theta = y) \).

Monte Carlo sampling allows for the estimation of \( P(\Theta = y | R_0 = r) \). Through use of Bayes’ theorem,
\[
P(R_0 > 1 | \Theta = y) = \int_0^1 dr P(\Theta = y | R_0 = r) P(R_0 = r)
\]
(E 1)
Assuming a prior \( P(R_0 = r) = 1/\Lambda \) if \( r \in [0, \Lambda] \) and 0 otherwise,
\[
P(R_0 > 1 | \Theta = y) = \int_0^1 dr P(\Theta = y | R_0 = r) P(R_0 = r).
\]
(E 2)
Finally, \( P(R_0 > 1 | \Theta = y) \) can be found numerically by binning recorded values of \( \Theta \) and approximating the integrals as summations over all bins.