# Early warning signals anticipate critical transitions in empirically-based simulations of measles dynamics in Niger

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### **Abstract**

Forecasting the trajectory of infectious disease outbreaks over time is a fundamental challenge facing society.

### Introduction

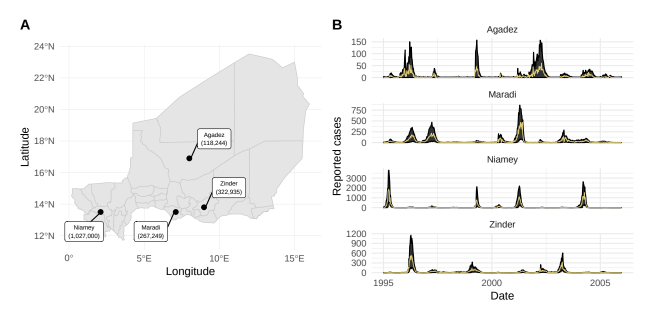
- Theory shows that epidemic transitions can be anticipated by trends in the statistical properties of disease
- 3 time series (AERO papers). The existence of statistical trends in the data that precede critical transitions,
- 4 so-called 'early warning signals' (EWS), imply that we may be able to anticipate disease emergence and
- outbreaks. The end goal is a model-independent detection system, where statistical properties of disease
- 6 surveilence data can trigger warnings of impending outbreaks without the need to fit mechanistic models of
- disease transmission (Han and Drake 2016).
- However, there is currently a gap between the theoretical work, which has relied on knowing the underlying
- 9 disease dynamics, and the eventual goal of applying EWS in real-world situations where the underlying
- disease dynamics may be unknown. Theoretical development of EWS has focused on anticipating when the
- population becomes supercritical, when  $R_0 > 1$ , after which an outbreak is inevitable, perhaps with some
- bifurcation delay (Dibble et al. 2016). Knowing the value of  $R_0$  through time makes it possible to test the
- accuracy of EWS that are estimated from state variables alone. Empirical application of EWS does not
- require knowing the value of  $R_0$  through time, meaning that "tests" require making assumptions about when
- 15 critical transitions occur. Whether EWS track and/or anticipate underlying dynamics of real disease time
- series remains unknown, and is a critical knowledge gap that must be filled before EWS can confidently be
- 17 deployed.
- 8 To fill this gap we will fit a mechanistic model to incidence data of measles in Niger to estimate the temporal
- 19 epidemiology of the disease, yielding the very same parameters that are known in data-free modeling studies.
- In particular, we are interested in the correlation between EWS and the time-varying repreoductive ratio,
- known as the effective reprodutive ratio  $(R_E)$ . If EWS and  $R_E$  are significantly and positively correlated, then
- we have empirical evidence that EWS are applicable in real-world settings. If EWS and  $R_E$  are negatively
- correlated or not significantly posivitely correlated, then we have evidence that EWS may not be applicable
- 24 in certain settings.

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## 25 Materials and Methods

#### 26 Data

- <sup>27</sup> We used weekly measles case report data from four Nigerien cities, collected over an 11 year period
- 28 (1995-2005) (Figure 1). These data are ideal for stress testing EWS because each city has different
- 29 population sizes, has different dynamics in terms of size of outbreaks and length of inter-epidemic periods,
- and each time series has different amounts of demographic stochasticity due to differences in population size.
- The data come from [somewhere/someone], and used here with permission from [somewhere/someone].



**Figure 1:** (A) Locations and population sizes (in parantheses) of our four focal cities in Niger. (B) Time series of reported cases (yellow solid lines) and the 95% prediction intervals for one-step-ahead forecasts from our fitted SEIR models for each city.

#### 32 Stochastic SEIR model

- 33 The model is a discrete-time approximation of a continuous-time SEIR model with limited demography,
- specified as a set of difference equations,

$$S_{t+dt} = n_{S,t} - n_{E,t} \tag{1}$$

$$E_{t+dt} = n_{E,t} - n_{I,t} \tag{2}$$

$$I_{t+dt} = n_{I,t} + n_{O,t} - n_{R,t}, (3)$$

where  $\mathbf{n}_t$  are random variables representing the number of individuals transitioning into or out of each class

- at each timestep  $t \to t + dt$ .  $n_S$  is the number of births,  $n_E$  is the number of newly infected individuals that
- have the disease but are not infectious,  $n_I$  is the number of newly infectious individuals,  $n_O$  is the number of
- $n_R$  is the number of newly recovered individuals who are no longer infectious and
- have life-long immunity. The stochastic random variables are specified as follows:

$$n_{S,t} \sim \text{Poisson}(\mu_t N_t \times dt)$$
 (4)

$$n_{E,t} \sim \text{Binomial}(\lambda_{E,t}, S_t)$$
 (5)

$$n_{Lt} \sim \text{Binomial}(\lambda_{Lt}, E_t)$$
 (6)

$$n_{O,t} \sim \text{Poisson}(\psi \times dt)$$
 (7)

$$n_{R,t} \sim \text{Binomial}(\lambda_{R,t}, I_t),$$
 (8)

- where  $\mu_t$  is the birth rate at time t,  $\psi$  is the rate of imported infections, and  $\lambda_E$ ,  $\lambda_I$ , and  $\lambda_R$  are the probabilities
- of exposure, becoming infectious, and recovery, respectively. These probabilities reflect the processes of
- tranmission, transition from the latent period to the infectious period, and recovery, which we model as:

$$\lambda_{E,t} = 1 - e^{-\frac{\beta_t I_t dt}{N_t}} \tag{9}$$

$$\lambda_{I,t} = 1 - e^{-\eta E_t dt} \tag{10}$$

$$\lambda_{R,t} = 1 - e^{-\gamma I_t dt},\tag{11}$$

where  $\beta_t$  is time-varying rate of transmission,  $\eta$  is time-invariant rate from the exposed class to the infectious class, and  $\gamma$  is time-invariant recovery rate. We model rate of transmission as:

$$\beta_t = \beta \left( 1 + \sum_{i=1}^6 q_i \xi_{i_t} \right) \Gamma_t. \tag{12}$$

 $\beta$  is the mean transmission rate,  $\psi$  accounts for measles infections from external sources that are not part of the local dynamics, and the term  $\sum_{i=1}^{6} q_i \xi_{i_t}$  is a B-spline to model seasonality in transmission. The B-spline bases  $(\xi_{i_t})$  are periodic with a 1 year period. The transmission rate  $(\beta_t)$  is also subject to stochastic process noise at each time step,  $\Gamma_t$ , which we model as a gamma-distributed white (temporally uncorrelated) noise with mean 1 and variance  $\sigma^2$  (Bretó and Ionides 2011).

We do not include a death process in the model because the rate of infection is much faster than the rate of death. Excluding deaths means we can avoid making further assumptions about demographic rates – we are already making assumptions about birth rates (e.g., the rate is the same across cities, but with city-specific population size). We model demographic stochasticity in births and imported infections by drawing time-specific values from Poisson distributions. Transitions in the model are shown in Table 1. In this model, the effective reproductive ratio at time t is:  $R_E(t) = \frac{\beta_t}{\gamma} \frac{S_t}{N_t}$ .

We assume observed case reports (y) are drawn from a Negative Binomial distribution subject to a constant reporting fraction  $(\rho)$  and dispersion parameter  $\tau$ ,

$$y_t \sim \text{Negative Binomial}(\rho I_t, \tau)$$
. (13)

Table 1: Transitions in the SEIR model. We show the determinstic transmission rate for clarity, but our model uses the stochastic tranmission rate.

Transition	$(\Delta S, \Delta E, \Delta I)$	Propensity
birth	(1,0,0)	$N_t \mu_t$
transmission	(-1, 1, 0)	$SI\beta_t/N_t$
(deterministic)		
transmission (stochastic)	(-k, k, 0)	$rac{S}{k}\sum_{j=0}^k inom{k}{j} (-1)^{k-j+1} au_{\mathrm{f}}^{-1} \mathrm{ln}(1+(eta_t I/N_t)) au_{\mathrm{f}}(S-j)$
symptomatic	(0, -1, 1)	$E\eta$
(infectious)	, , , ,	·
imported infections	(0,0,1)	$\psi_t$
recovery	(0,0,-1)	Ιγ

## Model fitting and inference

We fit the SEIR model to time series of case reports from each of our focal cities using Maximization by Iterated particle Filtering (MIF). We estimated 14 parameters for each city: six seasonal transmission 60 parameters  $(q_i)$ , mean transmission rate  $(\beta)$ , three initial conditions  $(S_{(t=0)}, E_{(t=0)}, I_{(t=0)})$ , the number of 61 imported infections  $(\psi)$ , reporting fraction  $(\rho)$ , one parameter accounting for process noise  $(\sigma)$ , and one 62 parameter accounting for measurement noise  $(\tau)$ . To ensure identifiability, and to make the model easier to fit, we assumed the infectious period was fixed at  $1/\eta = 8$  days and the recovery period was fixed at  $1/\gamma = 5$ days. The birth rate  $(\mu_t)$  was multiplied by 0.3 to account for the reported 70% vaccination coverage (Ferrari et al. 2008).

MIF relies on particle filtering, which estimates the likelihood of fixed parameters by integrating state variables of a stochastic system. To narrow in on the maximum likelihood estimates, MIF lets parameters take a random walk during the filtering process and selectively propagates forward parameter sets (i.e., particles) with the highest likelihood. The variance of the random walk decreases at each iteration of MIF, where a MIF iteration means one filtering pass through the time series. This procedure converges toward the 71 maximimum likelihood estimates (MLEs), in theory.

We used the IF2 algorithm (Ionides et al. 2015) implemented in the R (R Core Team 2017) package pomp (King et al. 2016, 2018) to conduct the MIF procedure. To initialize MIF, we generated 5000 parameter sets 74 using Latin Hypercube Sampling over large ranges of the parameter values. We then performed two rounds 75 of MIF, each for 100 iterations, with 10000 particles, and geometric cooling. For the first round of MIF we 76 set cooling.facter = 1. For the second round, which was initialized using the collection of parameter sets from the end of the first round, we set cooling.factor = 0.9. We computed the log likelihood of 5000 final MIF parameter sets (i.e., parameter sets collected after 200 MIF iterations) as the log of the mean likelihoods of 50 replicate particle filters with 10000 particles each. At this stage, we assume the parameter set with highest log likelihood is the MLE. 81

We used profiling to estimate 95% confidence intervals for four parameters of interest: mean transmission rate  $(\beta)$ , the initial fraction of susceptible individuals  $(S_{(t=0)})$ , the number of imported infections  $(\psi)$ , and reporting fraction ( $\rho$ ). For each parameter, we profiled over a range of 100 values using the same MIF and likelihood estimate procedure as described above for finding the MLEs. We replicated the profiling procedure 10 times for each unique value of the focal parameter and then took the log of the mean of those

- 10 profiled likelihoods. These values were used to estimate 95% confidence intervals using the Monte
- 88 Carlo-adjusted profile algorithm described by Ionides et al. (2017).

#### 89 Model simulations

## 90 Early warning signals

- We considered ten candidate early warning signals (EWS; Table 3). EWS were calculated using the function
- spaero::get\_stats() (O'Dea 2018) in R (R Core Team 2017).

**Table 2:** List of candidate early warning signals and their estimating equations. Note that b denotes the bandwidth. See Brett et al. (2018) for details.

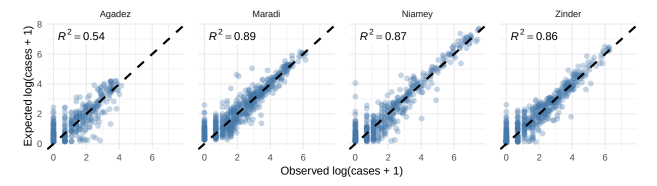
		Theoretical Correlation with		
EWS	Estimator	$R_E(t)$		
Mean	$\mu_t = \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{X_s}{2b-1}$	Positive		
Variance	$\sigma_t^2 = \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} rac{(X_s - \mu_s)^2}{2b-1}$	Positive		
Coefficient of variation	$CV_t = rac{\sigma_t}{\mu_t}$ $ID_t = rac{\sigma_t^2}{\mu_t}$	Null		
Index of dispersion	$ID_t = \frac{\sigma_t^2}{\mu_t}$	Positive		
Skewness	$S_t = \frac{1}{\sigma_t^3} \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{(X_s - \mu_s)^3}{2b-1}$	Positive		
Kurtosis	$K_t = \frac{1}{\sigma_t^4} \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{(X_s - \mu_s)^4}{2b-1}$	Positive		
Autocovariance	$ACov_{t} = \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{(X_{s}-\mu_{s})(X_{s-\delta}-\mu_{s-\delta})}{2b-1}$	Positive		
Autocorrelation	$AC_t = \frac{ACov_t}{\sigma_t \sigma_{t-\delta}}$	Positive		
Decay time	$\overline{ au}_t = -\delta/\ln[\mathrm{AC}_t(\delta)]$	Positive		
First differenced	$\Delta\sigma_t^2 = \sigma_t^2 - \sigma_{t-\delta}^2$	Positive		
variance				

## 3 Results

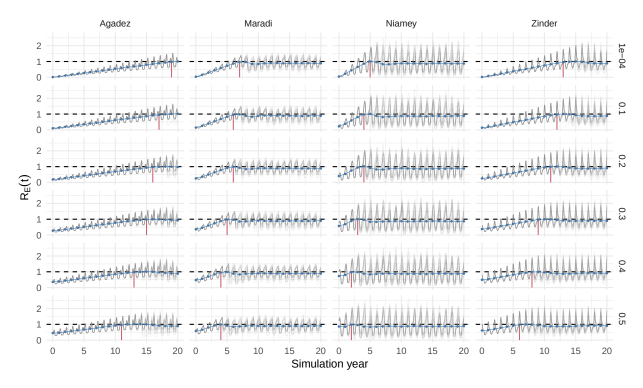
- $^{94}$  The fitted models adequately reproduced observed dynamics (Figure 1), with  $R^2$ s ranging from 0.54 to 0.89
- 95 (Figure 2).
- Stochastic simulations from sub-critical ( $R_E(t) \ll 1$ ) to critical dynamics ( $R_E(t) = 1$ ) differed among the
- 97 four cities (Figure 3).

 Table 3: Maximum likelihood estimates for select epidemiological parameters.

City	Log likelihood (S.E.)	β	σ	Ψ	ρ	$S_{t=0}$
Agadez	-960.65 (0.11)	171.47	0.10	7.80	0.77	0.23
Maradi	-1746.56 (0.16)	483.09	0.06	24.88	0.33	0.10
Niamey	-1454.73 (0.15)	370.63	0.09	23.28	0.26	0.11
Zinder	-1415.52 (0.09)	180.09	0.08	10.47	0.36	0.22



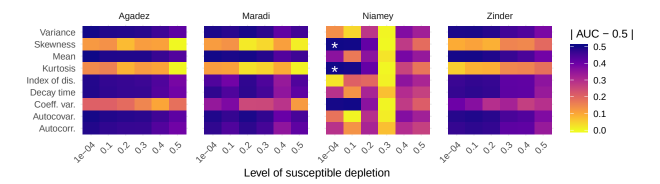
**Figure 2:** Comparison of in-sample model predictions and observations for each city. Expected cases are one-step-ahead predictions from the fitted models. The dashed line shows 1:1.



**Figure 3:** Yearly average  $R_E(t)$  (blue lines) across 500 models simulations at the MLE parameters and 20 representations simulations (grey lines) for each city. The horizontal dashed line shows where  $R_E(t) = 1$ . The vertical solid red lines show the time point at which yearly average  $R_E(t) \ge 1$ . The time periods before the red line for each city were used for testing early warning signals.

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**Figure 4:** Performance of EWS calculated over two windows (far from and near  $R_E(t) = 1$ ) from the time series of 500 simulated dynamics, shown as a heatmap of AUC values minus 0.5. The two windows were defined as equally-sized windows over the course of the time series up to  $R_E(t) = 1$  (red lines in Figure 3). The asterisks (\*) in the heatmap for Niamey note EWS with high AUC, but for the wrong reason: skewness and kurtosis *decrease* as  $R_E(t)$  approaches 1 rather than increase. This occurs because of the negative binomial sampling in the measurement component of our SEIR model.

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