Critical slowing down anticipates re-emergence and elimination of measles in Niger

Abstract

Introduction

- 2 Forecasts of the emergence and re-emergence of infectious diseases have the potential to save lives, money,
- and human productivity by allowing for proactive, rather than reactive, preparedness measures (1). Similarly,
- 4 indicators of the elimination of infectious diseases can signal the effectiveness of "end game" strategies
- 5 aimed at disease eradication (2). Predicting (re)emergence and elimination is possible with complex
- 6 mathematical models of disease transmission, but their success relies on detailed understanding of the
- underlying transmission and pathogen dynamics. We often do not have enough information to parameterize
- 8 such models. An alternative approach is to use model-independent statistical signals that portend infectious
- 9 disease (re)emergence and elimination by detecting critical slowing down as the system approaches a critical transition (3).
- Emergence and elimination of an infectious disease both involve a critical transition. The transition typically
- occurs at the critical point where the basic reproduction number (R_0 , the number of secondary cases that arise
- from a single infected case in a fully susceptible population) is equal to one (4). Thus, subcritical ($R_0 < 1$)
- and supercritical $(R_0 \ge 1)$ systems represent alternative modes of fluctuation (5, 6).
- 15 Critical transitions in stochastic systems, such as disease transmission systems, are often associated with
- critical slowing down, a reduction in the resilience of a system to perturbations (7). Critical slowing down
- 17 (CSD), in turn, is associated with changes in the dynamical features of the system, so-called early warning
- signals (EWS) such as an increase in the variance and autocorrelation (5, 8). Recent theoretical work
- suggests that CSD occurs as disease dynamics approach $R_0 = 1$ from below (emergence) (3, 9) and from
- above (elimination) (2, 3, 10), and that several EWS anticipate the critical transition (11-13). Empirical tests
- 21 of EWS and associated CSD are, however, scarce. Documenting CSD in real disease dynamics is an essential
- 22 first step toward the development of model-independent outbreak detection systems (1).
- 23 Here, we use empirically-based model simulations of measles dynamics to test whether CSD anticipates
- 24 critical transitions in real disease dynamics. We focus on two scenarios: (1) the re-emergence of measles
- 25 following a large outbreak, a situation typical of measles dynamics in sub-Saharan Africa, and (2) the
- elimination of measles by a vaccination campaign. We seek to answer two related questions. First, can CSD
- 27 distinguish between time series of disease incidence when the underlying dynamics are far from and near to a
- ²⁸ critical transition? If so, then CSD can anticipate disease re-emergence and elimination. Second, how does
- 29 the distance to and the rate of approaching the threshold impact the anticipatory skill of CSD?
- To answer these questions, we fit mechanistic models of disease transmission to time series of measles
- incidence in four Nigerien cities. We then use the fitted models to perform model experiments designed to

test the performance of several EWS, which quantify CSD, at anticipating re-emergence and elimination. Our results confirm theoretical expectations about several EWS and associated CSD. In particular, we show that CSD before a critical transition is detectable by several EWS in realistic scenarios, and they do so using much shorter time series than used in theoretical studies. However, our study highlights the limitations of EWS in situations where disease re-emergence and elimination occurs rapidly. Moreoever, and contrary to theoretical expectations (*3*), we find that EWS perform better at detecting CSD before re-emergence than before elimination.

39 Materials and Methods

40 Data

- We used weekly measles case report data from four Nigerien cities: Agadez, Maradi, Niamey, and Zinder.
- The data were collected over an 11 year period from 1995-2005 (Figure 1). These data are ideal for testing
- theory on CSD in disease dynamics because each city has different 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. Such differences provide a natural
- gradient of "noise" that may obscure CSD (citation?). The data come from [somewhere/someone], and used
- here with permission from [somewhere/someone].

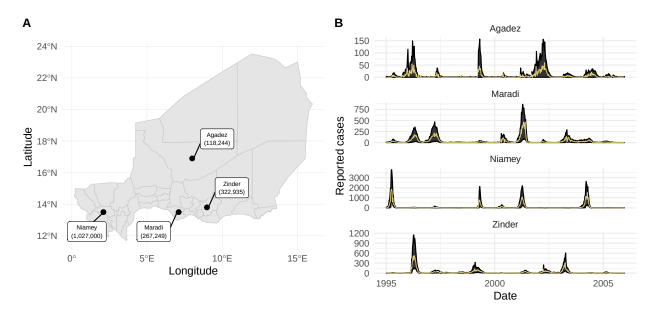


Fig. 1. Locations of data sources and observed and predicted measles dynamics. (**A**) Locations and population sizes (in parantheses) of our four focal cities in Niger. (**B**) Time series of weekly reported cases (yellow solid lines) and the 95% prediction intervals (black ribbons) for one-step-ahead predictions from our fitted SEIR models for each city.

48 Stochastic SEIR model

- 49 The model is a discrete-time approximation of a continuous-time SEIR model with limited demography,
- 50 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 imported infections, and 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)

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

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

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

where μ_t is the birth rate at time t, ψ is the rate of imported infections, and λ_E , λ_I , and λ_R are the probabilities of exposure, becoming infectious, and recovery, respectively. These probabilities reflect the processes of transmission, 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 β_t is time-varying rate of transmission, η is time-invariant rate from the exposed class to the infectious class, and γ is time-invariant recovery rate. We model rate of transmission as:

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

 β is the mean transmission rate, ψ 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 (ξ_{i_t}) are periodic with a 1 year period. The transmission rate (β_t) is also subject to stochastic process noise at each time step, Γ_t , which we model as gamma-distributed white (temporally uncorrelated) noise with mean 1 and variance σ^2 (14).

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: $R_E(t) = \frac{\beta_t}{\gamma} \frac{S_t}{N}$.
- We assume observed case reports (\mathbf{y}) are drawn from a Negative Binomial distribution subject to a constant reporting fraction (ρ) and dispersion parameter τ ,

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

Table 1. Transitions in the SEIR model. We show the determinstic transmission rate for clarity, but our model uses the stochastic transmission 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)	$\frac{S}{k} \sum_{j=0}^{k} {k \choose j} (-1)^{k-j+1} \tau_{f}^{-1} \ln(1 + (\beta_{t} I/N_{t})) \tau_{f}(S-j)$
symptomatic	(0, -1, 1)	$E\eta$
(infectious)		•
imported infections	(0,0,1)	ψ_t
recovery	(0,0,-1)	$I\gamma$

74 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 parameters (q_i) , mean transmission rate (β) , three initial conditions $(S_{(t=0)}, E_{(t=0)}, I_{(t=0)})$, the number of imported infections (ψ) , reporting fraction (ρ) , one parameter accounting for process noise (σ) , and one parameter accounting for measurement noise (τ) . 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 (μ_t) was multiplied by 0.3 to account for the reported 70% vaccination coverage (15).
- 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
- maximimum likelihood estimates (MLEs), in theory.
- We used the IF2 algorithm (16) implemented in the R (17) package pomp (18, 19) to conduct the MIF procedure. To initialize MIF, we generated 5000 parameter sets using Latin Hypercube Sampling over large ranges of the parameter values. We then performed two rounds of MIF, each for 100 iterations, with 10000 particles, and geometric cooling. For the first round of MIF we 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.

We used profiling to estimate 95% confidence intervals for four parameters of interest: mean transmission rate (β) , the initial fraction of susceptible individuals $(S_{(t=0)})$, the number of imported infections (ψ) , and reporting fraction (ρ) . 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 Carlo-adjusted profile algorithm described by (20).

Model simulations

We used the MLE parameter sets to make one-step-ahead predictions, to estimate any trend in transmission 104 rate through time, and to test the ability of early warning signals to anticipate the critical transition at 105 $R_E(t) = 1$. To make one-step-ahead predictions, we used particle filtering with 50000 particles and retained the mean and standard deviation of all latent across all particles before they were filtered at each time step. To estimate a trend in tranmission rate, we allowed mean transmission to take a random walk throughout the 108 filter and retained the filtered distribution of particles for mean transmission and $R_E(t)$. 100 We manipulated the initial size of the susceptible pool to simulate an increase from low $R_E(t)$ to high $R_E(t)$. Doing so allows us to test whether EWS can distinguish between windows of time when $R_E(t)$ is far from a 111 critical transition and when $R_E(t)$ is near a critical transition. We reduced the initial fraction of susceptible 112 individuals by multplying the MLE for $S_{(t=0)}$ by six discounting factors: 1e-4, 0.1, 0.2, 0.3, 0.4, and 0.5. 113

individuals by multplying the MLE for $S_{(t=0)}$ by six discounting factors: 1e-4, 0.1, 0.2, 0.3, 0.4, and 0.5.

These discounting factors represent situations of susceptible depletion after outbreaks of various size. We then simulated the model forward for thirty years using mean birth and death rates for the entire country.

Thirty years was long enough for yearly average $R_E(t)$ to reach 1 for each city. Because the model is stochastic, we repeated these simulations 500 times for each city-susceptible discount combination.

Next, we split each simulated time series into null and test intervals. First, across all simulations for a city-susceptible discount combination, we found the simulation year in which yearly average $R_E(t)$ reached 1, and excluded years past that time. We split the remaining time series into two windows of equal length. The null interval is the first window, where $R_E(t)$ is increasing but far from 1. The test interval is the second window, where $R_E(t)$ is increasing and approaching 1. We did this for each city and for each level of susceptible depletion. We calculated EWS over null and test intervals separately, as described next.

24 Early warning signals

We considered ten candidate early warning signals. We used the spaero::get_stats() function (21) in R (17) to calculate EWS according to the formulas in Table 2. All EWS except the coefficient of variation are expected to increase as $R_E(t)$ approaches 1 (12). However, we still include the coefficient of variation because it is often used EWS in other applications.

For each simulation of each city-susceptible discount combination, we calculated each EWS for the time

series of expected cases in the null and test intervals. This yields a distribution of EWS over the 500 null and test intervals. The bandwidth for EWS calculation varied because we set b to be one-half the length of the interval. That is, we do not use a moving window and if, for example, the length of the interval is 520 weeks (10 years), then $b = 0.5 \times 520 = 260$.

We assessed the performance of each EWS using the Area Under the Curve (AUC) statistic. Specifically, we use AUC to calculate the amount of overlap between the distributions of each EWS from the null and test intervals. Higher values of AUC indicate a greater degree of separation and thus better performance of a particular EWS in terms of classifying whether $R_E(t)$ is approaching a critical transition. We used the pROC: auc() function (22) in R to calculate AUC values.

Table 2. List of candidate early warning signals and their estimating equations. Note that b denotes the bandwidth. See (12) for details.

EWS	Estimator	Theoretical Correlation with $R_E(t)$
Mean	$\mu_t = \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{X_s}{2b-1}$	Positive
Variance	$\sigma_t^2 = \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{(X_s - \mu_s)^2}{2b-1}$	Positive
Coefficient of variation		Null
Index of dispersion	$CV_t = rac{\sigma_t}{\mu_t}$ $ID_t = rac{\sigma_t^2}{\mu_t}$	Positive
Skewness	$S_t = \frac{1}{\sigma_s^3} \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{(X_s - \mu_s)^3}{2b-1}$	Positive
Kurtosis	$K_t = \frac{1}{\sigma_s^4} \sum_{s=t-(b-1)\delta}^{t+(b-1)\delta} \frac{(X_s - \mu_s)^4}{2b-1}$	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	$\overline{\tau}_t = -\delta/\ln[AC_t(\delta)]$	Positive

139 Results

140 Model fit and simulations

The fitted models adequately reproduce observed dynamics (Figure 1), with in-sample R^2 s from one-step-ahead predictions ranging from 0.54 for Agadez to 0.89 for Maradi (Figure 2). Stochastic simulations of the models displayed dynamics typical of each city (Figure Sx), including the decline in seasonality amplitude as population size decreases (15). Our model for Agadez performs poorly relative to the other cities.

We found no evidence of a directional trend in the rate of tranmission over the course of the time series (Figure Sx). Seasonal fluctuations in all cities cause $R_E(t)$ to go above and below 1 throughout the course of a year (Figure Sx). This results in the observed outbreak cycles in low population cities and the more unpredictable outbreaks in larger cities (Figure 1).

Stochastic simulations from sub-critical ($R_E(t) \ll 1$) to critical dynamics ($R_E(t) = 1$) differed among the four cities (Figure ??). The differences in the rate at which $R_E(t)$ approaches one is a combination of population size, mean transmission rate, and seasonality (Table 3). For example, the dynamics of Niamey reach $R_E(t) = 1$ much faster than the other cities that have smaller populations, lower transmission rates (generally), and lower seasonal amplitude.

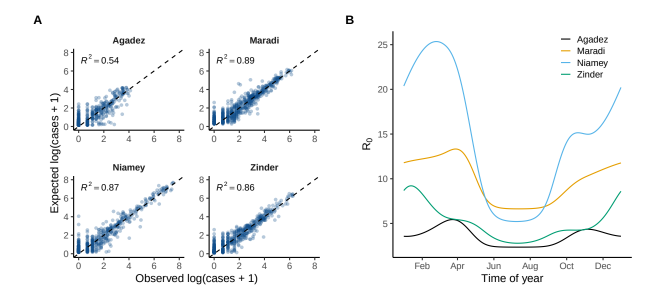


Fig. 2. Accuracy of the fitted SEIR models and estimated seasonality. (A) Comparison of in-sample model predictions and observations for each city. Expected cases are one-step-ahead predictions from the fitted models. The dashed line shows 1:1. Coefficients of determination (R^2) were calculated as the reduction in the sum-of-squared errors from model predictions relative to a null model of the mean number of cases (SI text). (B) The estimated seasonality of the basic reproductive ratio (R_0) for each city. R_0 was calculated as: $\frac{\eta \beta_t \mu}{\nu(\eta + \nu)(\gamma + \nu)}$, where $1/\eta$ is the infectious period, $1/\gamma$ is the recovery period, β_t is the time-specific rate of transmission, μ is the birth rate, and ν is the death rate. Only β_t is estimated by our model. We set $1/\eta = 8$ days, $1/\gamma = 5$ days, $\mu = \nu = 0.05$ for calculating R_0 as shown in this figure.

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

55 Performance of EWS

162

163

164

165

166

The EWS generally perform as expected, with Niamey being an exception (Figure 3). Skewness, kurtosis, and coefficient of variation performed poorly across all levels of susceptible depletion in all cities except
Niamey. For Niamey, it appears these metrics perform well at the highest levels of susceptible depletion (i.e., low discounting factors). In fact, the high AUC values for Niamey result from exceptionally poor performance: they are all higher in the null interval rather than the test interval, contrary to theoretical expectations (Table 2). Thus, these metrics are unreliable.

For the other cities, variance, mean, index of dispersion, decay time, autocovariance, and autocorrelation all perform equally well (Figure 3). Their performance declines as the amount of susceptible depletion decreases. This is expected because more rapid returns to $R_E(t) = 1$ result in shorter null and test intervals, making estimates of EWS less precise. Likewise, as the total time to reach $R_E(t) = 1$ decreases, the null and test intervals start to behave more similarly. In part, this is because susceptible replenishment is linear in our model. Nonlinear increases in the susceptible pool would lead to different dynamics.

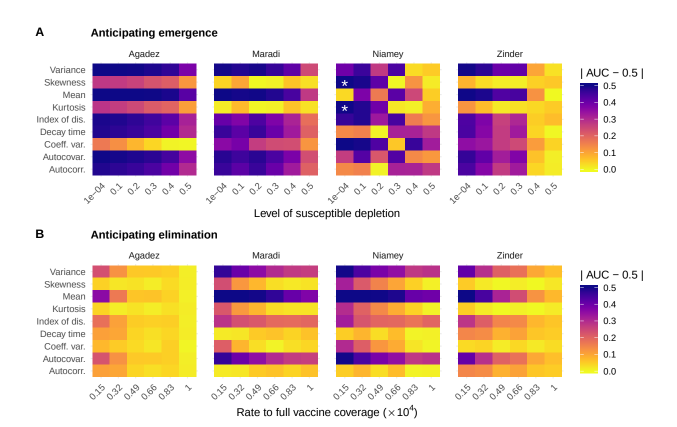


Fig. 3. Performance of early warning signals (EWS). calculated over two windows (far from and near $R_E(t)=1$) from the time series of 500 simulated dynamics, shown as a heatmap of AUC values minus 0.5. The two windows were defined as equally-sized windows over the course of the time series up to $R_E(t)=1$ (red lines in Figure ??). 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.

Discussion

Acknowledgments

This research was funded by the National Institute of General Medical Sciences of the National Institutes of Health (Award Number U01GM110744). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. This work was done on the Olympus High Performance Compute Cluster located at the Pittsburgh Supercomputing Center at Carnegie Mellon University, which is supported by National Institute of General Medical Sciences Modeling Infectious Disease Agent Study (MIDAS) Informatics Services Group grant 1U24GM110707.

References

- 1. B. A. Han, J. M. Drake, EMBO reports, e201642534 (2016).
- 2. J. M. Drake, S. I. Hay, Tropical Medicine and Infectious Disease. 2, 20 (2017).
- 3. S. M. O'Regan, J. M. Drake, *Theoretical Ecology*. **6**, 333–357 (2013).
- 4. J. M. Heffernan, R. J. Smith, L. M. Wahl, Journal of the Royal Society Interface. 2, 281–293 (2005).
- 5. M. Scheffer et al., Nature. 461, 53–59 (2009).
- 6. M. Scheffer et al., Science. 338, 344–348 (2012).
- 7. E. H. van Nes, M. Scheffer, *The American Naturalist*. **169**, 738–747 (2007).
- 8. S. R. Carpenter, W. A. Brock, *Ecology Letters*. **9**, 311–318 (2006).
- 9. C. J. Dibble, E. B. O'Dea, A. W. Park, J. M. Drake, *Journal of the Royal Society Interface*. **13**, 20160540 (2016).
- 10. S. M. O'Regan, J. W. Lillie, J. M. Drake, *Theoretical Ecology*, **9**, 269–286 (2016).
- 11. T. S. Brett, J. M. Drake, P. Rohani, Journal of the Royal Society Interface. 14, 20170115 (2017).
- 12. T. S. Brett et al., PLoS Computational Biology. 14, e1006204 (2018).
- 13. P. B. Miller, E. B. O'Dea, P. Rohani, J. M. Drake, *Theoretical Biology and Medical Modelling*. **14**, 17 (2017).
- 14. C. Bretó, E. L. Ionides, Stochastic Processes and their Applications. 121, 2571–2591 (2011).
- 15. M. J. Ferrari et al., Nature. 451, 679-684 (2008).
- 16. E. L. Ionides, D. Nguyen, Y. Atchadé, S. Stoev, A. A. King, *Proceedings of the National Academy of Sciences*. **112**, 719–724 (2015).
- 17. R Core Team, R: A language and environment for statistical computing (2017), (available at https://www.r-project.org/).
- 18. A. A. King, D. Nguyen, E. L. Ionides, Journal Of Statistical Software. 69, 1–43 (2016).
- 19. A. A. King *et al.*, pomp: Statistical Inference for Partially Observed Markov Processes (R package, version 1.18) (2018).
- 20. E. L. Ionides, C. Breto, J. Park, R. A. Smith, A. A. King, Journal of the Royal Society Interface. 14,

20170126 (2017).

- 21. E. B. O'Dea, spaero: Software for Project AERO (R package version 0.3.0) (2018).
- 22. X. Robin et al., BMC Bioinformatics. 12, 77 (2011).