Bayesian workflow for disease transmission modelling in Stan

Swiss Epidemiology Winter School 2021

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Preface

- · Objective: fit transmission models in Stan
- Based on Grinsztajn et al., 2020 (link v2 upcoming)
- Prerequisites:
 - · introduction to infectious disease modelling
 - · general understanding of Bayesian inference
 - basic programming with R and Stan 2.21
- All material is available on https://github.com/jriou/bayesian_workflow



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Outline

Introduction

Basics of Bayesian inference

Bayesian modelling with Stan

Fitting a simple SIR model

Simulation-based model assessment

The Bayesian workflow

Scaling-up ODE-based models

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Objectives



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Introduction

Models of disease transmission:

- Interpretability: mechanistic, phenomenological
- Granularity: agent-based, population-based
- Framework: deterministic, stochastic
- Data-generating mechanisms: incubation, contagion, immunity...



Models of disease transmission:

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Mechanistic + population-based + deterministic

ightarrow ordinary differential equations (ODE)-based compartmental model



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Introduction

ODE-based compartmental model:

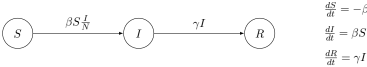
- Divide the population into homogeneous groups (compartments)
- Define the flows between compartments with ODEs
- Define initial conditions (at t₀)
- Solve for the time-dependent volume in each compartment



ODE-based compartmental model:

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- Define the flows between compartments with ODEs
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- Solve for the time-dependent volume in each compartment

The susceptible-infectious-recovered (SIR) model:

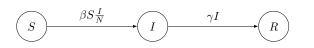


$$\frac{dS}{dt} = -\beta S \frac{I}{N}$$

$$\frac{dI}{dt} = \beta S \frac{I}{N} - \gamma I$$

$$\frac{dR}{dt} = \gamma I$$





$$\frac{dS}{dt} = -\beta S \frac{I}{N}$$

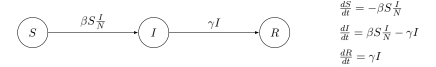
$$\frac{dI}{dt} = \beta S \frac{I}{N} - \gamma I$$

$$\frac{dR}{dt} = \gamma I$$

Where:

- S(t) is the number of people susceptible to infection
- I(t) is the number of people infected (i.e. the prevalence)
- R(t) is the number of people recovered (lifelong immunity)
- N is the population size (S(t) + I(t) + R(t) = N for any t)
- β is the infectious contact rate (per day per person)
- γ is the recovery rate (1/infectious period)

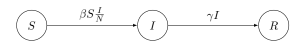




Intuition behind the SIR model:

- I(t)/N is the proportion of infected (and infectious)
- $\beta I(t)/N$ is the daily number of contacts with infectious people
- hence each day, $\beta SI(t)/N$ people become infected (the force of infection)





$$\begin{aligned} \frac{dS}{dt} &= -\beta S \frac{I}{N} \\ \frac{dI}{dt} &= \beta S \frac{I}{N} - \gamma I \\ \frac{dR}{dt} &= \gamma I \end{aligned}$$

Assumptions behind the SIR model:

- homogeneous mixing
- β and γ constant over time
- all infections are observed
- no incubation, exponentially-distributed recovery
- lifelong immunity
- stable population



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Introduction

Simulate in R with package deSolve:

set compartments and differential equations

```
> ## Set model ----
> seir = function(t, x, parms, ...) {
+ with(as.list(c(parms, x)), {
+ dS = - beta*S*I/(S*I+R)
+ dI = beta*S*I/(S*I+R) - gamma*I
+ dR = gamma*I
+ list(c(dS, dI, dR))
+ })
+ }
```



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+ list(c(dS, dI, dR))
+ })
+ }
```

• set (fixed) values for $\beta=0.8$; $\rho=1/7$; $S_0=100,000-50$; $I_0=50$ and $R_0=0$

```
> N_0 = 100000

> I_0 = 50

> inits = c(

+ S = N_0 - I_0,

> pars = c(beta = 0.8,

+ gamma = 1/7

+ )
```



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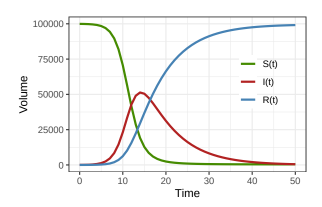
Introduction

• solve the ODE system numerically (Runge-Kutta 4th order) to obtain unique solutions for S(t), I(t) and R(t)

$$f(\beta, \gamma, S_0, I_0, R_0) = \{S(t), I(t), R(t)\}$$

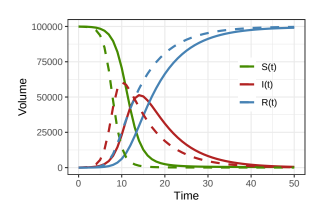


with
$$\beta = 0.8$$
; $\rho = 1/7$; $S_0 = 100000 - 50$; $I_0 = 50$ and $R_0 = 0$



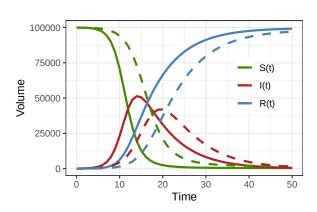


with $\beta = 1.1$ instead of 0.8, we get



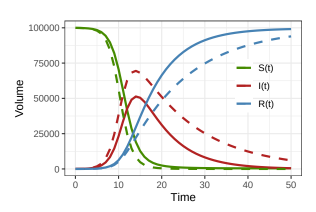


with $\beta = 0.6$ instead of 0.8, we get



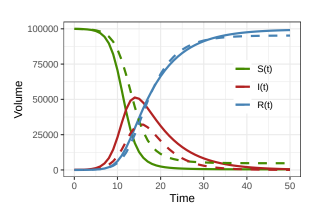


with $\gamma = 1/14$ instead of 1/7, we get



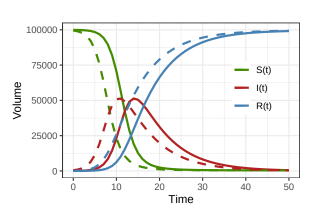


with $\gamma = 1/4$ instead of 1/7, we get



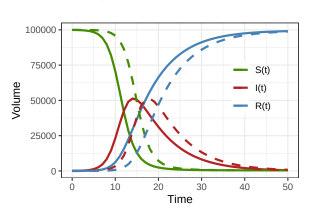


with I(0) = 500 instead of 50, we get



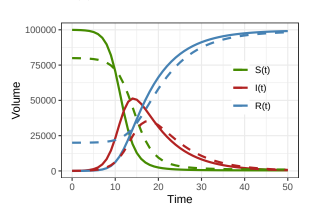


with I(0) = 5 instead of 50, we get





with R(0) = 20,000 instead of 0, we get





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Introduction

Compartmental models have many uses:

- formalize and put numerical values on general concepts (herd immunity, vaccination threshold...)
- get mechanistic insight about an epidemic (transmissibility levels, drivers of transmission)

$$\mathcal{R}_0 = \frac{\beta}{\gamma}$$

produce precise forecasts (based on mechanisms)



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Introduction

Compartmental models have many uses:

- formalize and put numerical values on general concepts (herd immunity, vaccination threshold...)
- get mechanistic insight about an epidemic (transmissibility levels, drivers of transmission)

$$\mathcal{R}_0 = \frac{\beta}{\gamma}$$

- produce precise forecasts (based on mechanisms)
- ightarrow all these uses are based on numerical values for β , ρ and the initial conditions and their uncertainty



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Introduction

Enters Bayesian inference:

- infer parameter values by integrating information from data and from domain knowledge
- more efficient for complex models (high dimensionality)
- rigorously quantify and propagate uncertainty in parameter estimates and forecast



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- more efficient for complex models (high dimensionality)
- rigorously quantify and propagate uncertainty in parameter estimates and forecast

→ Markov Chain Monte Carlo (MCMC) methods and Stan





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Bayesian inference

General principle of Bayesian inference:

- (1) specify a complete Bayesian model
 - consider data $y=\{y_1,...,y_n\}$ that are noisy measurement of an unknown quantity (parameter) θ
 - encode the data-generating processes in an observation model (likelihood)

$$p(y|\theta) = \prod_n^N \mathsf{normal}(y_n|\theta,1)$$

- encode domain knowledge in a prior distribution

$$p(\theta) = \mathsf{normal}(0, 1)$$



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Bayesian inference

(2) reverse the question:

"given a model and data, what are plausible parameter values?"

- the answer comes from Bayes' rule:

$$p(\theta \mid y) \propto p(\theta)p(y \mid \theta)$$

- where $p(\theta \mid y)$ is the posterior distribution, i.e. the set of plausible parameter values given data and model



Bayesian inference

In practice, we rarely have an analytic expression for $p(\theta \mid y)$ and rely on inference algorithms to learn about the posterior distribution

Monte Carlo methods

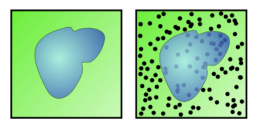


Figure: Monte Carlo approach to determine the area of a lake using random artillery fire.



Bayesian inference

 Markov chain Monte Carlo (MCMC) samplers apply this principle to probability distributions

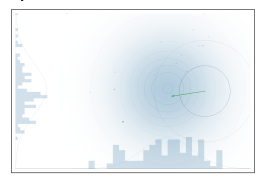


Figure: An MCMC sampler exploring a bivariate normal distribution (https://chi-feng.github.io/mcmc-demo/app.html).



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Bayesian modelling with Stan

Stan is a probabilistic programming framework for Bayesian inference

- it is designed to let the user focus on modelling while inference happens under the hood
- object-oriented language (based on C++) that supports many operations, probability densities and ODE solvers
- extremely efficient MCMC algorithm (Hamiltonian Monte Carlo)
- diagnostic tools to evaluate the inference
- interfaces in R (package rstan), python, julia...



Bayesian modelling with Stan

Programming in Stan is structured in blocks:

the data block defines known variables and constants.

```
data {
   int N;
   real y[N];
}
```

the parameters block defines unknown variables (parameters)

```
parameters {
  real theta;
}
```

 the model block defines the prior distribution(s) and the observation model(s)

```
model {
  theta ~ normal(0,1);
  y ~ normal(theta,1);
}
```

save in model_linear.stan



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Bayesian modelling with Stan

We then explore the target with Stan's MCMC sampler:

load rstan package

```
## Setup ----
library(rstan)
options(mc.cores = parallel::detectCores())
```

• simulate N=50 data points with $\theta=0.7$

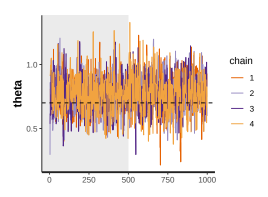
```
## Simulate data ----
N = 50
theta = 0.7
y = rnorm(N,theta,1)
input_data = list(N=N,y=y)
```

run MCMC sampling



Bayesian modelling with Stan

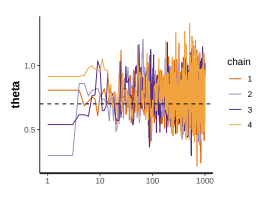
We use multiple chains that should converge after warm-up





Bayesian modelling with Stan

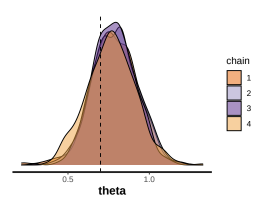
We use multiple chains that should converge after warm-up





Bayesian modelling with Stan

The post-warm-up samples of θ approximate its posterior distribution





Bayesian modelling with Stan

We run basic diagnosis tools: divergences, tree depth, energy

```
> check_hmc_diagnostics(fit)

Divergences:
0 of 2000 iterations ended with a divergence.

Tree depth:
0 of 2000 iterations saturated the maximum tree depth of 10.

Energy:
E-BFMI indicated no pathological behavior.
```



Bayesian modelling with Stan

Printing the object gives:

- diagnostics: effective sample size, Gelman-Rubin \hat{R}
- inference: full posterior distribution of θ

```
> print(fit)
Inference for Stan model: model_linear.
4 chains, each with iter=1000; warmup=500; thin=1;
post-warmup draws per chain=500, total post-warmup draws=2000.

mean se_mean sd 2.5% 25% 50% 75% 97.5% n_eff Rhat
theta 0.78 0.01 0.14 0.51 0.69 0.78 0.87 1.05 760 1
lp__ -18.30 0.03 0.73 -20.28 -18.44 -18.03 -17.86 -17.81 799 1

Samples were drawn using NUTS(diag_e) at Thu Nov 12 19:15:33 2020.
For each parameter, n_eff is a crude measure of effective sample size,
and Rhat is the potential scale reduction factor on split chains (at
convergence, Rhat=1).
```



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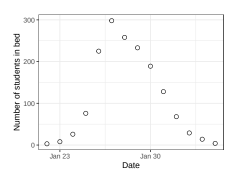
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Example data: outbreak of influenza A (H1N1) at a British boarding school in 1978 (available in R package outbreaks)

- 763 students, 512 had symptoms
- daily number of students in bed over 14 days (prevalence data)





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Fitting a simple SIR

Specifying the model:

- prevalence data: \mathbb{I}_t with $t \in \{1, \dots, 14\}$
- parameters to estimate: $\theta = \{\beta, \gamma, \phi\}$
- parameters that will remain fixed: $\{S_0 = 762, I_0 = 1, R_0 = 0\}$
- map data \mathbb{I}_t to SIR model output I(t) using an observation model with an appropriate probability distribution:

$$\Pr(\mathbb{I}| heta) = \prod_{t=1}^{14} \mathsf{neg ext{-}bin}(\mathbb{I}_t|I(t),\phi)$$

prior distributions

$$\Pr(\beta) = \text{exponential}(1)$$

 $\Pr(1/\gamma) = \text{normal}(2, 0.5)$
 $\Pr(1/\phi) = \text{exponential}(5)$



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Fitting a simple SIR

We define the ODE system in the function block

```
real[] sir(real t, real[] y, real[] theta, real[] x_r, int[] x_i) {
 real S = y[1];
 real I = y[2];
 real R = y[3];
 real N = x_i[1];
 real beta = theta[1]:
 real gamma = theta[2];
  real dS dt = -beta * I * S / N:
 real dI_dt = beta * I * S / N - gamma * I;
  real dR_dt = gamma * I;
 return {dS_dt, dI_dt, dR_dt};
```

▲ Be careful of the signature and formats!



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Fitting a simple SIR

We declare the data variables in the data block

```
data {
  int<lower=1> T;
  real y0[3];
  real t0;
  real ts[T];
  int N;
  int cases[T];
}
```



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Fitting a simple SIR

We declare the data variables in the data block

```
data {
   int<lower=1> T;
   real y0[3];
   real t0;
   real ts[T];
   int N;
   int cases[T];
}
```

and define additional data variables in transformed data

```
transformed data {
  real x_r[0];
  int x_i[1];
  x_i[1]=N;
}
```



Similarly, parameters are declared in the parameters block

```
parameters {
  real<lower=0> beta;
  real<lower=0> recovery_time;
  real<lower=0> phi_inv;
}
```

f A It sometimes makes more sense to transform some parameters (e.g., recovery rate γ and overdispersion ϕ) to improve interpretability



In transformed parameters, we define additional parameters and solve the ODE system $\,$

```
transformed parameters{
  real y[T,3];
  real phi = 1. / phi_inv;
  real gamma = 1. / recovery_time;
  real theta[2];
  theta[1] = beta;
  theta[2] = gamma;

  y = integrate_ode_rk45(sir, y0, t0, ts, theta, x_r, x_i);
}
```



```
y = integrate_ode_rk45(sir, y0, t0, ts, theta, x_r, x_i);
```

Two crucial points:

- be careful about the formats and signatures
 - the ODE output y is an array of size T×3 (number of time steps and number of compartments)
 - sir is the name of the function defined in the function block
 - y0 is an array of size 3 defined in the data block
 - ts is an array of size T defined in the data block
 - theta is an array of size 2 storing the parameters
 - x_r is defined as empty in transformed data, but can be used to store fixed real values
 - x_i is an array of size 1 storing the population size N (can also be used to store fixed integer values)



```
y = integrate_ode_rk45(sir, y0, t0, ts, theta, x_r, x_i);
```

Two crucial points:

- two ODE solvers are available:
 - integrate_ode_rk45 uses the Runge-Kutta method (quicker but non-adapted to stiff systems)
 - integrate_ode_bdf uses the backward differentiation method (slower but adapted to stiff systems)



In the model block, we write the priors and the observation model

```
model {
    // priors
    beta ~ exponential(1);
    recovery_time ~ normal(2,0.5);
    phi_inv ~ exponential(5);

    // observation model
    cases ~ neg_binomial_2(col(to_matrix(y),2), phi);
}
```

A It's important that the chosen distributions correspond with the boundaries set in the parameters block (<lower=0>)

 \triangle col(to_to_matrix(y)) extracts the 2nd column of y



Last, we add a generated quantities block that does not influence sampling and can be used for "post-processing":

- reproduction number $\mathcal{R}_0 = \beta/\gamma$
- model predictions of prevalence from the negative binomial

```
generated quantities {
  real R0 = beta/gamma;
  real pred_cases[T];
  pred_cases = neg_binomial_2_rng(col(to_matrix(y),2), phi);
}
```



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Fitting a simple SIR

In summary:

- functions: define the ODE system (A signature and formats)
- data: declare data variables that will be provided
- tranformed data: additional quantities that can be computed internally or from data variables
- parameters: declare parameters (boundaries)
- transformed parameters: quantities that can be computed internally or from data or parameters variables, including the ODE output (A signature and format)
- model: priors and observation model
- generated quantities: additional quantities that can be computed without influencing the sampling



As before, we conduct the inference from R with the package rstan:

```
# prevalence data
cases = influenza england 1978 school$in bed
N = 763
n days = 14
t0 = 0
t = 1:n_days
i0 = 1
s0 = N - i0
r0 = 0
y0 = c(s0, i0, r0)
input_data = list(T = n_days, y0 = y0, t0 = t0, ts = t, N <u>= N, cases = cases)</u>
```

A data is put in a list with names matching the data block in Stan



Hit the inference button!

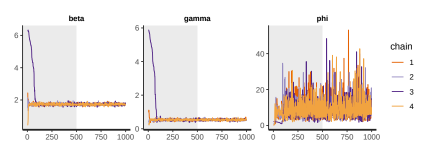


Run basic diagnosis tools:

```
> check_hmc_diagnostics(fit)
Divergences:
0 of 2000 iterations ended with a divergence.
Tree depth:
0 of 2000 iterations saturated the maximum tree depth of 10.
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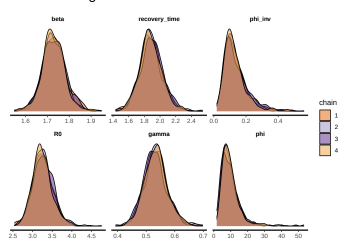


Examine trace plots:



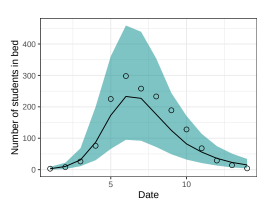


Examine chain mixing:





Posterior predictive checking (always show!):





Print the results:

```
print(fit,pars=c("beta","gamma","phi","R0","recovery time"))
Inference for Stan model: sir negbin.
4 chains, each with iter=1000; warmup=500; thin=1;
post-warmup draws per chain=500, total post-warmup draws=2000.
             mean se mean
                            sd 2.5% 25% 50% 75% 97.5% n eff Rhat
                     0.00 0.06 1.61 1.70 1.73 1.76 1.85 1049
beta
             1.73
            0.53  0.00  0.04  0.44  0.50  0.53  0.56  0.63  1382
gamma
phi
             9.61 0.22 6.13 2.94 5.72 8.31 11.80 23.40 743
RO
             3.27
                     0.01 0.29 2.79 3.09 3.25 3.42 3.96 1403
recovery time 1.89
                     0.00 0.16 1.59 1.79 1.88 1.98 2.25 1410
Samples were drawn using NUTS(diag e) at Thu Nov 12 19:10:02 2020.
For each parameter, n_eff is a crude measure of effective sample size,
and Rhat is the potential scale reduction factor on split chains (at
convergence, Rhat=1).
```



Conclusions:

- we estimate \mathcal{R}_0 to 3.3 (95% credible interval: 2.8 to 4.0)
- this corresponds to the direct estimation from the final size of the epidemic q=512/763=0.67

$$\mathcal{R}_0 = 1/(1-q) = 3.03$$

- based on many assumptions:
 - common to all SIRs (homogeneous mixing, no incubation...)
 - prior distributions (especially on the recovery period)
 - complete ascertainment, no asymptomatics
 - no initial immunity



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Simulation-based model assessment

In practice, the situation is often less clear than in the boarding school example:

- incomplete data, insufficient domain knowledge
- uncertainty on necessary model features

Fake data can be used to probe the model and better understand its behaviour:

- prior predictive checking
- simulation study



Simulation-based model assessment

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Prior predictive checking consists in simulating data from the priors:

- visualize priors (especially after transformation)
- this shows the range of data compatible with the model
- it helps understand the adequacy of the chosen priors, as it is often easier to elicit expert knowledge on measureable quantities of interest rather than abstract parameter values



Simulation-based model assessment

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- visualize priors (especially after transformation)
- this shows the range of data compatible with the model
- it helps understand the adequacy of the chosen priors, as it is often easier to elicit expert knowledge on measureable quantities of interest rather than abstract parameter values
- remove (or switch off) the likelihood from the model block

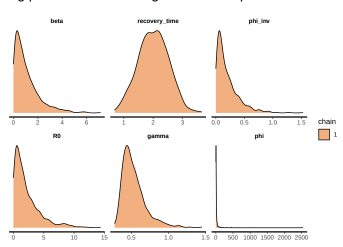
```
// observation model
if(switch_likelihood==1) cases ~ neg_binomial_2(col(to_matrix(y),2), phi);
```



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Simulation-based model assessment

Simulating priors in the boarding school example:

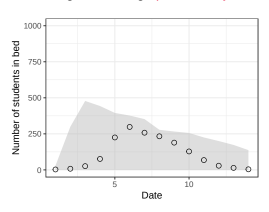




Simulation-based model assessment

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Prior predictive checking: simulating epidemic trajectories

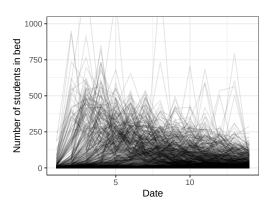




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Simulation-based model assessment

Prior predictive checking: simulating epidemic trajectories





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Simulation-based model assessment

Prior predictive checking brings insight about non-obvious features:

- even if the priors seem weakly informative, there is actually not a lot of leeway
- highly constrained model:
 - if β is high, the epidemic will stop rapidly by lack of susceptibles
 - if β is small, the epidemic will be small
- the negative binomial might lead to problems in extreme situations, e.g. more cases (>1000) than the overall number of students



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Simulation-based model assessment

A simulation study consists in two steps:

- simulate data with specified parameter values
- measure the capacity of the model to recover the chosen parameter values



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Simulation-based model assessment

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- simulate data with specified parameter values
- measure the capacity of the model to recover the chosen parameter values

Many advantages:

- check for bugs and coding mistakes
- · check for identifiability issues
- compare different versions of a model
- understand in what situations a model works or not

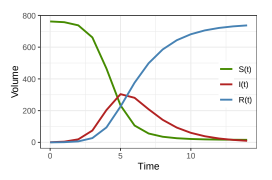


Simulation-based model assessment

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Let's go back to the simple SIR example from the beginning:

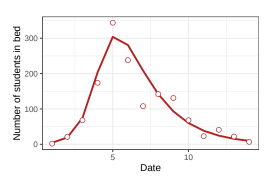
- set $\beta=2$ and $\gamma=0.5$ (so that $\mathcal{R}_0=4$)
- simulate in a susceptible population of size N=763 with $I_0=1$





Simulation-based model assessment

Add noise with a negative binomial distribution with $\phi=15$

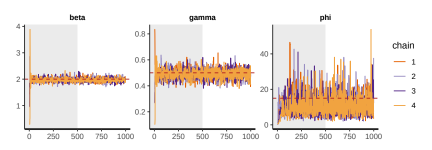




Simulation-based model assessment

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Fit the same model as for the boarding school example

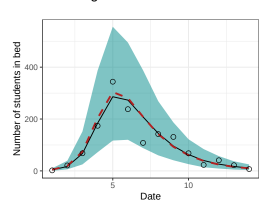




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Simulation-based model assessment

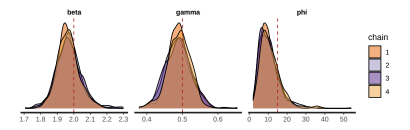
Posterior predictive checking





Simulation-based model assessment

Compare the posterior distributions of the parameters with the "truth"



- no identifiability issue
- β and γ are well estimated, but ϕ is not
- try with other values to understand when does the model break



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Outline

Introduction

Basics of Bayesian inference

Bayesian modelling with Stan

Fitting a simple SIR model

Simulation-based model assessment

The Bayesian workflow

Scaling-up ODE-based models

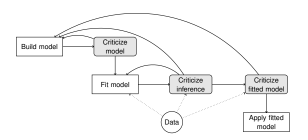
Extensions

Objectives



The Bayesian workflow

Bayesian workflow: model development as an iterative process





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The Bayesian workflow

- (1) Build a complete model:
 - encode the data-generating processes with observation models
 - · encode domain knowledge with prior distributions
- (2) Criticize the model with fake data
 - prior predictive checking: is the model adequate?
 - simulation study: does the model work as intended?
- (3) Fit the model to real data
- (4) Criticize the inference
 - chain mixing (trace plots, posteriors by chain, Rhat)
 - basic diagnosis tools (divergences, energy, n_eff)
- (5) Ciriticize fitted model
 - · posterior predictive checking
- (6) Application (inference, forecast...)



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Scaling up ODE-based models

Inference with ODE-based models is computationally intensive

Be attentive to the model structure:

 HMC requires to compute the gradient of the log joint density multiple times for each iteration

$$\nabla_{\theta} \log \Pr(y, \theta)$$

- each block is treated differently
 - transformed data and generated quantities are evaluated once per iteration
 - parameters, transformed parameters and model are evaluated multiple times for each iteration
 - \rightarrow put everything that does not influence the inference (e.g. \mathcal{R}_0 or predicted values) in generated quantities



Scaling up ODE-based models

Limiting the load of the ODE solver:

```
y = integrate_ode_rk45(sir, y0, t0, ts, theta, x_r, x_i);
```

- the computational cost of solving the ODEs scales with N+NK
 - *N*: the number of compartments
 - K: the number of parameters in y0 and theta
- \rightarrow remove unnecessary compartments (e.g. R(t))
- reparametrize initial conditions



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Scaling up ODE-based models

Picking the right ODE solver:

- two are available:
 - integrate_ode_rk45 uses the Runge-Kutta method (quicker but non-adapted to stiff systems)
 - integrate_ode_bdf uses the backward differentiation method (slower but adapted to stiff systems)
- there is no formal definition of stiffness
- intuitively, it occurs when the time step of the integrator needs to be very small to keep the solution stable (e.g. large variations in magnitude in time)
- \rightarrow start with rk45, move to bdf if there are problems (folk theorem of statistical computing)



Scaling up ODE-based models

Tuning the ODE solver:

additional options in the solver function

- rel_tol is the relative tolerance
- abs_tol is the absolute tolerance
- max_steps is the maximum number of steps
- can be adjusted depending on the level of precision needed

A be cause of the tolerance, the ODE solver may sometimes give negative values when too close to zero, causing issues

 \rightarrow this can be solved by adding 5-10 times the absolute tolerance to the ODE output



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Extensions

In practice, the boarding school example quickly reaches its limits:

- epidemics are not often observed in such a controlled environment (under-ascertainment)
- epidemics are not always left uncontrolled
- data generally consist of daily counts of new cases (incidence)
 rather than counts of currently sick people (prevalence)
- most infectious diseases have an incubation period (SEIR instead of SIR)
- transmission is generally not homogeneous in the full population (stratification by age, sex...)
- ...



Extensions

Example with SARS-CoV-2 from Hauser et al. (2020):

PLOS MEDICINE

⑥ OPEN ACCESS № PEER-REVIEWED RESEARCH ARTICLE

Estimation of SARS-CoV-2 mortality during the early stages of an epidemic: A modeling study in Hubei, China, and six regions in Europe

Anthony Hauser, Michel J. Counotte, Charles C. Margossian, Garyfallos Konstantinoudis, Nicola Low, Christian L. Althaus, Julien Riou 🖸

Published: July 28, 2020 • https://doi.org/10.1371/journal.pmed.1003189



Background

The case fatality ratio (CFR) is computed as the number of deaths divided by the number of reported cases at time t.

Estimated in real time, the CFR is a misleading indicator of mortality due to SARS-CoV-2 because of two opposing biases:

- preferential ascertainment of severe cases: severe cases are both more likely to die and more likely to be diagnosed and reported
- → overestimates mortality
 - right-censoring of deaths: there is a long delay between infection and deaths, so that part of the cases at time t will die in the future
- → underestimates mortality



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Background

This limits the interpretability of the CFR:

- varies in time (ascending or descending phase)
- varies across countries (depending on surveillance system)
- → April 2020: from 2.4% in Wuhan to 17.8% in Lombardy)
 - the real indicator of interest is the infection fatality ratio, i.e. the total number of deaths that occur among people infected with SARS-CoV-2



Objectives

- (1) Simulate the dynamics of transmission and mortality of SARS-CoV-2 using publicly available surveillance data
- (2) Provide overall and age-stratified estimates of IFR for SARS-CoV-2 infection adjusted for right-censoring and preferential ascertainment

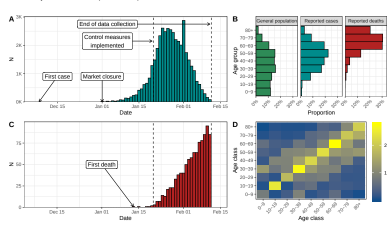
in seven different geographic locations with available data on:

- reported cases by date of disease onset
- deaths linked to SARS-CoV-2 infection by date of death
- age distribution of cases
- age distribution of deaths



Data

In Hubei province (China):



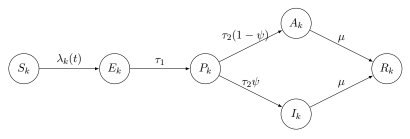


Specific features and natural history of SARS-CoV-2 infection:

- Incubation period of 5 days (SEIR)
- Pre-symptomatic transmission accounting for 44-48% (SEPIR)
- Symptoms in 81% (95%Crl 71%–89%) of cases (SEPIAR)
- Respiratory virus transmitted through contacts (age stratification)
- Effect of control measures (time-dependent force of infection)
- Mortality is delayed by 20.2 ± 11.6 days (fit to mortality data)



Final model:



S_k	Susceptible (for age group k)
E_k	Exposed
P_k	Presymptomatic
A_k	Infected asymptomatic
I_k	Infected symptomatic
R_k	Removed

 $\begin{array}{c|c} \lambda_k(t) & \text{Force of infection (time-dependent)} \\ 1/\tau_1+1/\tau_2 & \text{Incubation period (split in two)} \\ \psi & \text{Proportion of symptomatic} \\ 1/\tau_2 & \text{Presymptomatic infectious period} \\ 1/\mu & \text{Symptomatic infectious period} \end{array}$



Incubation:

 SEIR: adding a compartment E for exposed, i.e. infected but not yet symptomatic



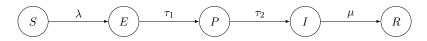
- τ is the inverse of the incubation period
- individuals are infectious from symptom onset (when entering *I*)

$$\lambda = \beta \frac{I}{N}$$



Pre-symptomatic transmission:

• SEPIR: adding a compartment P for pre-symptomatic, i.e. not yet symptomatic but already infectious



- the incubation period is split in two phases with rates au_1 and au_2
- individuals are infectious before symptom onset (entering *P*)

$$\lambda = \beta \frac{P + I}{N}$$

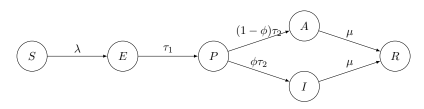
we can introduce reduced transmissibility before symptom onset

$$\lambda = \beta \frac{\kappa P + I}{N}$$



Asymptomatic infections:

• SEPIAR: adding a compartment A for asymptomatic



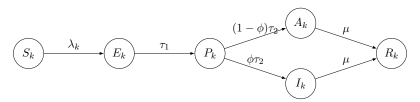
- we introduce the proportion of symptomatics ϕ
- we can introduce reduced transmissibility for asymptomatics

$$\lambda = \beta \frac{\kappa P + I + \kappa A}{N}$$



Age stratification:

- the transmission of respiratory viruses (influenza virus, rhinovirus...) is highly dependent on age
- the mortality of respiratory infections (even more so for SARS-CoV-2) is highly dependent on age
- \rightarrow stratification in nine age groups $k \in \{1, \dots, 9\}$ for (0-9, ..., 80+)





Characterizing the force of infection:

• in the simple SIR, the force of infection is defined as the rate at which susceptible individuals acquire infection

$$\lambda = \beta \frac{I}{N}$$

• the transmission rate β can be split in a contact rate c times a probability of transmission upon contact β , so that

$$\lambda = \beta c \frac{I}{N}$$

f A the same symbol eta is used for the transmission rate and the probability of transmission upon contact depending on context



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Model development

time-dependent force of infection using a forcing function

$$\lambda = f(t)\beta c \frac{I}{N}$$

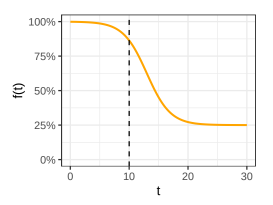
- to model the effect of control measures, we want a downward function that maps onto the interval [0,1], e.g. a logistic function:

$$f(t) = \eta + \frac{1 - \eta}{1 + \exp(\xi(t - t_c - \nu))}$$

- η is the relative reduction in transmission after control measures
- ξ is the slope of implementation of the control measures
- ν is the delay until the control measures are 50% effective (in days after t_c , the date of introduction of control measures).

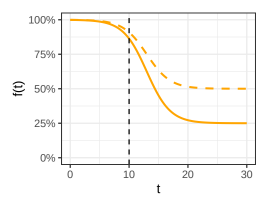


with
$$t_c=10$$
; $\eta=0.25$; $\nu=3$ and $\xi=0.5$



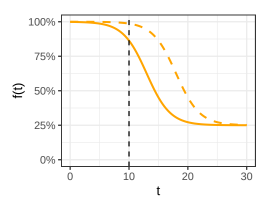


with $\eta=0.5$ instead of 0.25, we get



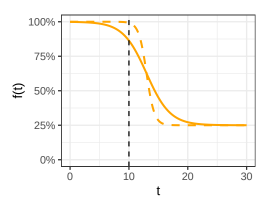


with $\nu=8$ instead of 3, we get





with $\xi = 1.5$ instead of 0.5, we get





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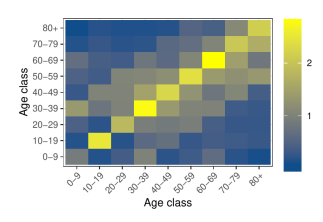
we account for behaviour differences across age groups:

$$\lambda_k(t) = f(t)\beta \sum_{l=1}^{9} \mathbb{F}_{k,l} \frac{I_l}{N_l}$$

- one force of infection for each age group k
- includes a specific contact rate between age group k and each age group l (corresponding to one cell of the contact matrix $\mathbb{F}_{k,l}$)
- includes the prevalence in age group $l \; (I_l/N_l)$



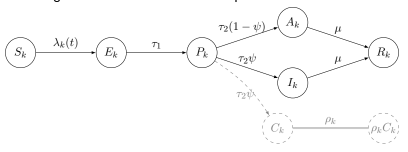
Age-specific contact matrix in China





Model fit to reported cases:

• obtaining incidence from the ODE output:



- dummy compartment $C_k(t)$ records the cumulative incidence of symptomatic infections for each age group k

$$\frac{dC_k}{dt} = \tau_2 \psi P_k$$



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Model development

new symptomatic infections by day of symptom onset by age:

$$\Delta C_{k,t} = C_k(t) - C_k(t-1)$$

• new reported infections per day of symptom onset, introducing the age-specific ascertainment proportion ρ_k :

$$A_t = \sum_{k=0}^{9} \rho_k \Delta C_{k,t}^I$$

• the age distribution of all reported cases up to $t_{\sf max}$:

$$B_k = \frac{\rho_k C_k^I(t_{\text{max}})}{\sum_k^9 \rho_k C_k^I(t_{\text{max}})}$$



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Model development

• A_t can be mapped to reported incidence data ${\mathbb A}$ using a negative binomial likelihood:

$$\Pr(heta|\mathbb{A}) = \prod_{t=t_1}^{t_{\mathsf{max}}} \mathsf{Neg\text{-Bin}}(\mathbb{A}_t|A_t,\phi_1)$$

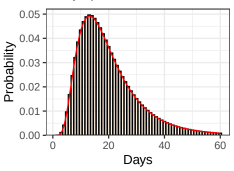
• B_k can be mapped to the age distribution of reported cases $\mathbb B$ using a multinomial likelihood:

$$\Pr(\theta|\mathbb{B}) = \mathsf{Multinomial}(\mathbb{B}_1, \dots, \mathbb{B}_9 | B_1, \dots, B_9)$$



Model fit to deaths:

- mortality is considered outside of the system of ODEs, using an age-specific mortality parameter ε_k (probability of death given symptomatic infection)
- we account for the delay with a discretized log-normal distribution of time from symptom onset to death I of length 60





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Model development

• deaths in age group k at time t ($1 \le t \le t_{\max} + 60$) among people infected up to t_{\max} :

$$M_{k,t} = \varepsilon_k \sum_{d}^{60} \Delta C_{k,t-d} \mathbb{I}_d$$

 deaths summed over age groups, assuming that all deaths are reported:

$$M_t = \sum_{k=0}^{9} M_{k,t}$$

the age distribution of all deaths occurring up to t_{max}:

$$D_k = \frac{\sum_{t=1}^{t_{\text{max}}} M_{k,t}}{\sum_{t=1}^{t_{\text{max}}} M_t}$$



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Model development

• M_t can be mapped to daily death data \mathbb{C} using a negative binomial likelihood:

$$\Pr(heta|\mathbb{C}) = \prod_{t=t_1}^{t_{\sf max}} \mathsf{Neg\text{-Bin}}(\mathbb{C}_t|M_t,\phi_2)$$

• D_k can be mapped to the age distribution of deaths \mathbb{D} using a multinomial likelihood:

$$\Pr(\theta|\mathbb{D}) = \mathsf{Multinomial}(\mathbb{D}_1, \dots, \mathbb{D}_9|D_1, \dots, D_9)$$

This leads to the following joint likelihood:

$$\Pr(\theta|\mathbb{A}, \mathbb{B}, \mathbb{C}, \mathbb{D}) = \Pr(\theta|\mathbb{A}) \cdot \Pr(\theta|\mathbb{B}) \cdot \Pr(\theta|\mathbb{C}) \cdot \Pr(\theta|\mathbb{D})$$

with
$$\theta = \{\beta, \eta, \xi, \nu, \psi, \pi, \rho_k, \varepsilon_k, \phi_1, \phi_2\}.$$



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Model development

Last bits:

- there is an identifiability issue with ρ
- \rightarrow fix ρ_9 (for 80+) to 100%, assuming that all symptomatic infections among very high risk persons will be reported
 - some remaining unknowns (data correction in China, role of children, lower ρ_9 ...)
- → sensitivity analyses



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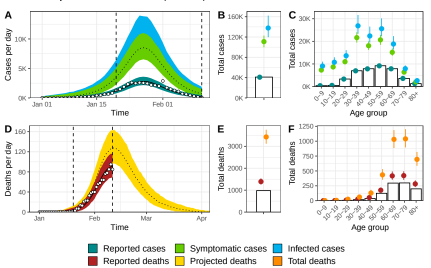
Inference

You know the drill:

- set priors
- prior predictive check
- ullet sampling (on high performance computing cluster \sim 2h)
- basic diagnostic tests
- examine trace plots and chains
- posterior predictive check

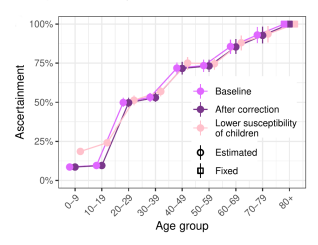


Posterior predictive check (Hubei):



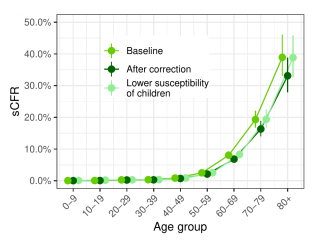


Ascertainment (posteriors of ρ_k):



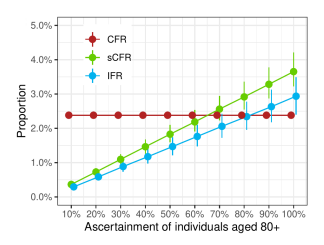


Mortality among symptomatics or sCFR (posteriors of ε_k):





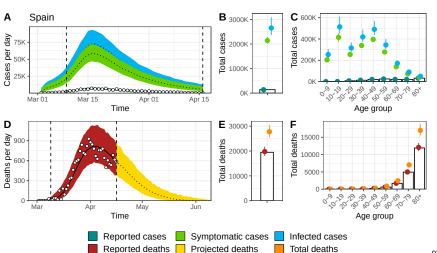
Effect on the assumption on ρ_9 on IFR estimate:





Results in all regions

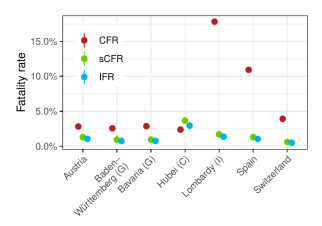
Posterior predictive check (Spain):





Results in all regions

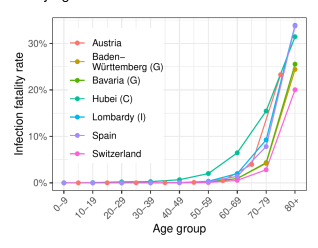
IFR estimates (compared to CFR and sCFR)





Results in all regions

IFR estimates by age





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Conclusions

Location (limit date)	Estimated attack rate	CFR	sCFR	IFR
Hubei, China (11 February)	0.2% (0.2-0.3)	2.0%	3.1% (2.7-3.5)	2.5% (2.1-2.9)
Austria (14 April)	0.8% (0.6-0.9)	2.8%	1.3% (1.1-1.6)	1.1% (0.8-1.3)
Baden-Württemberg,	1.9% (1.7-2.2)	2.6%	0.9% (0.6-1.6)	0.7% (0.5-1.3)
Germany (16 April)				
Bavaria,	2.0% (1.7-2.3)	2.9%	0.9% (0.7-1.3)	0.8% (0.5-1.1)
Germany (16 April)				
Lombardy, Italy (25 April)	11.5% (10.1-13.4)	17.8%	1.7% (1.5-2.0)	1.4% (1.1-1.6)
Spain (16 April)	5.7% (5.0-6.6)	10.9%	1.3% (1.2-1.5)	1.0% (0.9-1.2)
Switzerland (23 April)	3.6% (2.9-4.5)	3.9%	0.6% (0.5-0.8)	0.5% (0.4-0.6)

- IFR estimates adjusted for under-ascertainment and right-censoring are more similar across countries than CFR
- still some degree of heterogeneity
- clear increase of mortality with age



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Conclusions

General comments:

- knowing the data-generating mechanisms to avoid misinterpretation
- this knowledge can be used to build a model to adjust for known biases
- estimates of IFR obtained early in the epidemic, mostly confirmed in later seroprevalence studies



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Acknowledgements & ressources

General Bayesian inference & Stan:

- Andrew Gelman et al. Bayesian Data Analysis
 http://www.stat.columbia.edu/~gelman/book/
- Stan forums https://discourse.mc-stan.org/
- Michael Betancourt, Introduction to Stan
 https://betanalpha.github.io/assets/case_studies/
 stan_intro.html
- Andrew Gelman et al., Bayesian workflow https://arxiv.org/abs/2011.01808
- Richard McElreath, Statistical rethinking https://youtu.be/4WVelCswXo4



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Acknowledgements & ressources

Stan and ODEs or infectious disease models:

- Daniel Lee, ODEs in Stan https://youtu.be/hJ34_xJhYeY
- Chatzilena et al., Contemporary statistical inference for ID models

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https://www.sciencedirect.com/science/article/pii/S1755436519300325
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- Grinsztajn et al., Bayesian workflow for disease transmission modeling in Stan

https://arxiv.org/abs/2006.02985