Multilevel models in epidemiology

An application to chikungunya and Zika epidemics

Advanced statistical methods for physicists

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Bern, 31 May 2019

Themes

Provide a real-world example of model development in relation to a scientific question:

- Mechanistic model (data-generating processes)
- Multilevel structure in relation to data structure
- Partial pooling of information

mosquitoes

The global invasion of Aedes

Aedes mosquitoes (i)

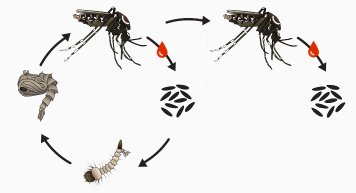


Figure 1: Life cycle of Aedes mosquitoes.

- Blood meal necessary to egg maturation
- Biting behaviour: gonotrophic cycle
- Transmission through saliva: specific vectorial competence

Aedes mosquitoes (ii)

Two species are important to human health:

- Aedes aegypti (tropical and subtropical areas)
- Aedes albopictus (subtropical et temperate areas)



Figure 2: Female adult specimens of Aedes aegypti (left) and Aedes albopictus (right).

The domestication of Aedes aegypti

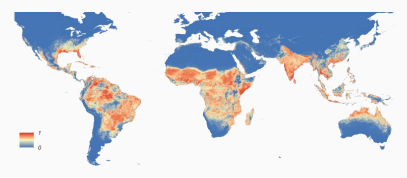


Figure 3: World distribution of Aedes aegypti 1.

- Originally from Africa, extension from the 15th century
- Urban and domestic species: adapted to human settlements²

¹Kraemer et al., eLife (2015); ² Powell et Tabachnick, Memorias do Instituto Oswaldo Cruz (2013)

The invasion of Aedes albopictus

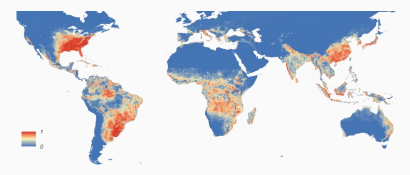


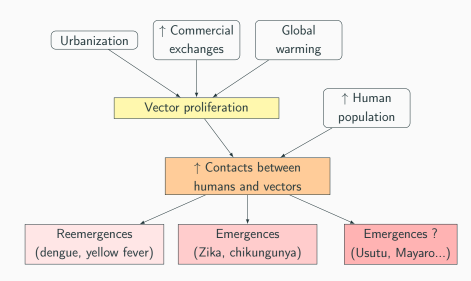
Figure 4: World distribution of Aedes albopictus 1.

- Originally from Asia, extension from the 20th century²
- Invasive species: plasticity, competitive advantages³

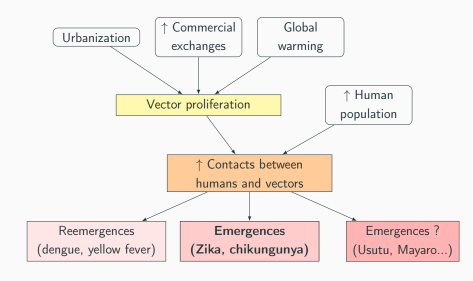
¹Kraemer et al., eLife (2015); ²Reiter, Journal of the American Mosquito Control Assoc. (1998);

³Paupy et al., Microbes and Infection (2009)

Disease emergences



Disease emergences



World propagation of chikungunya (CHIKV)

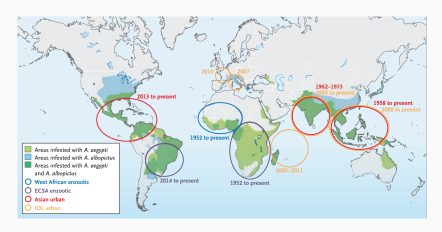


Figure 5: Origin and extension of the chikungunya virus and his vectors¹.

¹Weaver et al., New England Journal of Medicine (2015)

World propagation of Zika (ZIKV)

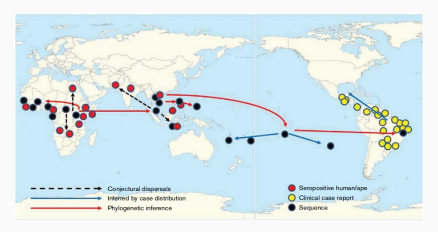


Figure 6: Origin and extension of the Zika virus¹.

¹Gatherer et Kohl, *Journal of General Virology* (2016)

Comparing Zika and chikungunya

epidemics

Strategy

Successive waves of chikungunya and Zika epidemics:

- each circulating for the first time
- in the same areas
- within a short timespan
- ⇒ Comparison of epidemics of different viruses:
 - in the same populations (immunologically naive)
 - in the same environments (vectors)
 - observed by the same surveillance systems

Data

Time-series of incidence data for 18 outbreaks of ZIKV and CHIKV:

- weekly number of reported cases CHIKV or ZIKV
- between 2013 and 2016
- in 9 islands with similar surveillance systems

```
oad("zikachik.Rdata")
zikachikSrecord... SREGION SREGION ID SISLAND SISLAND ID SISLAND ABB SVIRUS SVIRUS ID SDATE
                                                                                                  SWEEK SYEAR SN DAY SN WEEK SNCASES
             185 FP
                                  0 AUSTRA...
                                                      6 AUS
                                                                                     0 2014-11-09
                                                                                                     45 2014
                                                                                                                                       7000 91954
                                  @ AUSTRA...
                                                                     CHTKV
                                                                                                     46 2014
             187 FP
                                   6 AUSTRA..
                                                      6 AUS
                                                                                    0 2014-11-23
                                                                                                     47 2014
                                  0 AUSTRA...
                                                                                    0 2014-11-30
                                                                                                     48 2014
             189 FP
                                  @ AUSTRA...
                                                                     CHTKV
                                                                                                     49 2014
                                                                                                                                       7860 91954
                                   0 AUSTRA...
                                                      6 AUS
                                                                                    0 2014-12-14
                                                                                                     50 2014
             191 FP
                                  0 AUSTRA...
                                                                                    0 2014-12-21
                                                                                                     51 2014
             192 FP
                                   @ AUSTRA.
                                                                     CHTKV
                                                                                     0 2014-12-28
                                                                                                     52 2014
             193 FP
                                  0 AUSTRA...
                                                      6 AUS
                                                                                    0 2015-01-04
                                                                                                      1 2015
                                                                                                                                    65 7880 91954
                                  A ALISTRA
                                                                                     0 2015-01-11
                                                                                                      2 2015
                                                                                                                                    74 7060 91954
```

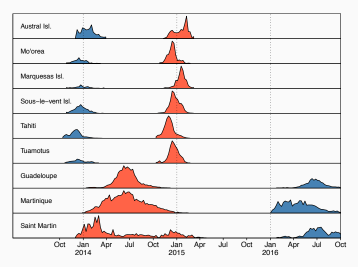


Figure 7: Profiles of CHIKV (red) and ZIKV (blue) incidence in nine territories during 2013-2016.

Data-generating processes: observation

How is the number of reported cases on week t (O_t) generated?

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- ullet true number of infections by CHIKV or ZIKV (I_t)
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- sufficient to lead to a consultation with a physician
- recognized by the physician
- reported by the physician to the surveillance authorities

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At the observation level:

$$O_t \sim \mathsf{Binom}(I_t, \rho)$$

 \Rightarrow Parameter ρ : probability of reporting

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- (if we ignore importations)
- all new cases can be linked back to cases in the last few weeks
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At the transmission level:

$$I_t \sim \mathsf{Binom}\left(S_t, eta \middle[rac{1}{N} \sum_{n=1}^5 w_{t,n} I_{t-n} \middle] \right)$$

- \Rightarrow Parameter β : number of secondary cases by primary case (\mathcal{R}_0)
- \Rightarrow Exposure: depends on the serial interval w_t

Definition

<u>Serial interval</u>: the time between the disease onset of a primary case and one of its secondary cases¹

Reconstruction using the full transmission cycle

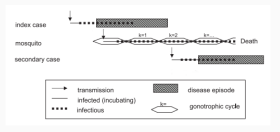


Figure 8: Framework for calculating the distribution of the serial interval².

¹Svensson et al, *Math. biosciences* (2007); ²Boëlle et al, *Vector-borne and zoonotic diseases* (2007)

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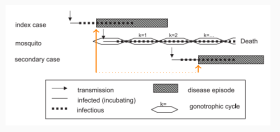


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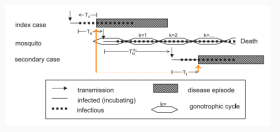


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Using published data on each stage (including dependence to local temperature \mathcal{T}) we obtain:

$$T_{SI} = -T_V + T_B + T_M(T) + T_I$$

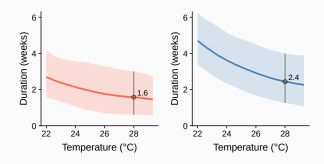


Figure 9: Distribution of the serial interval for CHIKV (red) and ZIKV (blue) according to temperature.

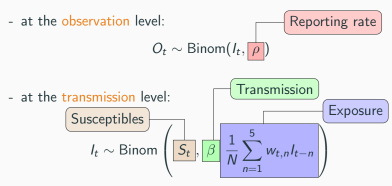
One disease, one island: model

Let O_t be the observed incidence in one epidemic on week t:



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- at the transmission level: Transmission Exposure $I_t \sim \text{Binom}\left(\frac{S_t}{N}, \frac{\beta}{N} \sum_{n=1}^5 w_{t,n} I_{t-n}\right)$

$$O_t \sim \mathsf{Neg\text{-}Binom}\left(S_t \frac{\beta}{N} \sum_{n=1}^5 w_{t,n} \frac{O_{t-n}}{\rho}, \frac{\phi}{\phi}\right)$$

Overdispersion

In a separate .stan file:

Data block:

```
data {
   int<lower=1> W; // number of records
   int<lower=0> O_t[W]; // number of reported cases at time t
   real<lower=0> Ostar_t[W]; // exposure at time t
   int<lower=0> sumO_t[W]; // cumulative number of reported cases at time t
   int<lower=0> pop; // island population
}
```

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Parameters block:

```
parameters {
  real<lower=0> beta;
    real<lower=0,upper=1> rho;
  | real<lower=0> phi;
}
```

• Transformed parameters block:

```
transformed parameters {
  real<lower=0> lp[W];
  real<lower=0> sampledisp[W];
  for(i in 1:W) {
    lp[i] = ( 1 - sum0_t[i] / (rho * pop)) * beta * Ostar_t[i] ;
    sampledisp[i] = lp[i]/phi;
  }
}
```

NB:
$$S_t \frac{\beta}{N} \sum_{n=1}^{5} w_{t,n} \frac{O_{t-n}}{\rho} = \left(1 - \frac{\sum_t O_t}{\rho N}\right) \beta O_t^*$$

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Model block:

```
model {
  beta ~ exponential(0.1);
  rho ~ beta(1,1);
  phi ~ cauchy(0,2.5);
   // likelihood
   target += neg_binomial_2_lpmf(0_t|lp,sampledisp);
}
```

Control Stan from R with library(rstan):

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Results: posterior distributions of β , ρ and ϕ

```
> print(S_GUAD,pars=c("beta","rho","phi"))
Inference for Stan model: TSIR_one_island.
4 chains, each with iter=2000; warmup=1000; thin=1;
post-warmup draws per chain=1000, total post-warmup draws=4000.

mean se_mean sd 2.5% 25% 50% 75% 97.5% n_eff Rhat
beta 1.54 0.00 0.06 1.42 1.50 1.53 1.58 1.66 1632 1
rho 0.33 0.00 0.02 0.30 0.32 0.33 0.34 0.38 1539 1
phi 51.04 0.24 11.47 33.31 42.74 49.42 57.74 77.75 2258 1

Samples were drawn using NUTS(diag_e) at Tue May 28 13:04:45 2019.
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).
```

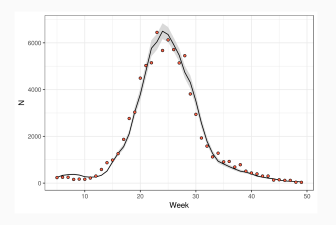


Figure 10: Model fit for the epidemic of Zika virus in Guadeloupe.

Now for 9 islands

Degrees of pooling:

• independent β_i and ρ_i for each island: no pooling

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- independent β_i and ρ_i for each island: no pooling
- the same β and ρ for all islands: complete pooling
- correlated β_i and ρ_i for each island: partial pooling = multilevel or hierarchical

For the epidemic in island i, we have:

• a transmission parameter β_i which depends on hyperparameters μ_{β} and σ_{β} :

$$\ln \beta_{i,j=0} \sim \mathcal{N}(\mu_{\beta}, \sigma_{\beta}^2)$$

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• a transmission parameter β_i which depends on hyperparameters μ_{β} and σ_{β} :

$$\ln \beta_{i,j=0} \sim \mathcal{N}(\mu_{\beta}, \sigma_{\beta}^2)$$

• a reporting parameter ρ_i which depends on hyperparameters μ_{ρ} and σ_{ρ} :

$$\ln \frac{\rho_{i,j=0}}{1-\rho_{i,j=0}} \sim \mathcal{N}(\mu_{\rho}, \sigma_{\rho}^2)$$

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• a transmission parameter β_i which depends on hyperparameters μ_{β} and σ_{β} :

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• a reporting parameter ρ_i which depends on hyperparameters μ_{ρ} and σ_{ρ} :

$$\ln \frac{\rho_{i,j=0}}{1-\rho_{i,j=0}} \sim \mathcal{N}(\mu_{\rho}, \sigma_{\rho}^2)$$

 \Rightarrow We now also estimate μ_{β} , μ_{ρ} , σ_{β} and σ_{ρ}

We now model together the epidemics of ZIKV (j = 1) and CHIKV (j = 0) assuming proportionality:

• on the transmission parameters of ZIKV and CHIKV:

$$\beta_{i,j=1} = \eta \times \beta_{i,j=0}$$

 on the reporting parameters of ZIKV and CHIKV (on the logit scale):

$$\frac{\rho_{i,j=1}}{1 - \rho_{i,j=1}} = \omega \times \frac{\rho_{i,j=0}}{1 - \rho_{i,j=0}}$$

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 \Rightarrow We now also estimate η and ω

The model fit is acceptable for CHIKV:

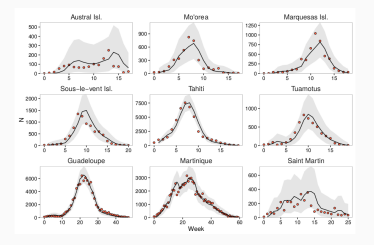


Figure 11: Model fit for the CHIKV epidemics.

And for ZIKV:

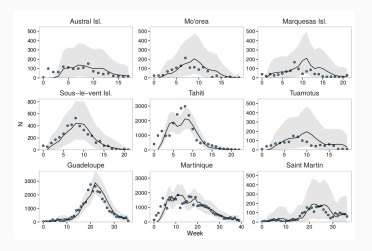


Figure 12: Model fit for the ZIKV epidemics.

The main results are:

- a similar transmissibility of CHIKV and ZIKV within an area
 - η |data = 1.04 [0.97 1.13]
- a lower reporting rate for ZIKV
 - $\omega | \text{data} = 0.37 [0.34 0.40]$

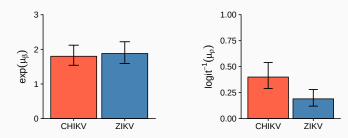


Figure 13: Posterior estimates of μ_{β} and μ_{ρ} for CHIKV and ZIKV.

We also find heterogeneity between areas:

- σ_{β}^2 data > 0, lower β in the French West Indies
- σ_{ρ}^2 data > 0, higher ρ in small islands and in Martinique

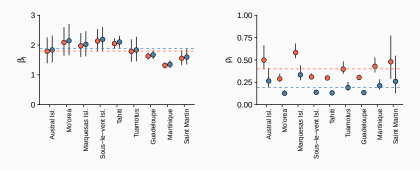


Figure 14: Island-specific posterior estimates of β and ρ for CHIKV (red) and ZIKV (blue).

Conclusion

Remember about:

- Data-generating mechanisms
- Plot model predictions (fit)
- Multi-level structure following data structure (partial pooling)