# Environmentally Transmitted Pathogens

Models - Part deux :

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The tetanus model of Cvjetanović The model of Capasso for ETP

A model for zoonotic transmission of waterborne disease

The first schistosomiasis model of Woolhouse

The third schistosomiasis model of Woolhouse – Heterogeneous contacts

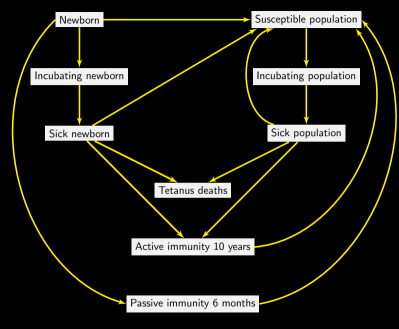
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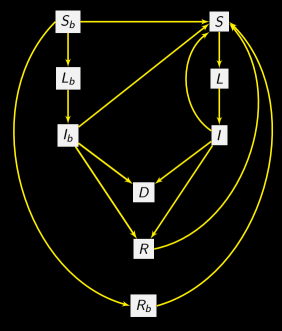
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# The discrete-time tetanus model (notation mine)

where

$$T = S + L_b + L + I_b + I + R + R_b$$
 and  $\delta_T = \frac{\Delta D}{T}$  (1j)

### Parameter assumptions – Tetanus

- **Incubation period** Mean duration 6 days for newborn and 8 days for general population  $\Rightarrow$  daily rate of exit  $\varepsilon_b = 0.1667$  and  $\varepsilon = 0.125$
- Period of sickness Mean duration 3 days for newborn and 14 days for general population  $\Rightarrow$  daily rate of exit  $\gamma_b = 0.3333$  per sick newborn and  $\gamma = 0.0714$  for sick general in general population
- Mortality from tetanus Untreated tetanus cases, fatality rate 90% for newborn  $S_b$  and 40% for general population. Treated: 80% for newborn and 30% general population
- Immunity Tetanus cases do not lead to immunity to reinfection. But as a general rule, recovered people are vaccinated. Convalescents and general population effectively immunised by complete course of vaccination go to R for average 10 years, daily rate of exit is  $\nu = 0.000274$  per person.
- ▶ Immunity of newborns Newborn to women vaccinated during pregnancy are temporarily protected by maternal antibodies and pass through  $R_b$  for a mean duration of 6 months. Daily rate of exit  $\nu_b = 0.005479$  per immunised newborn

## Parameter assumptions – Demography

Live birth rate 35 per 1,000 population and annual crude death rate 15 per 1,000 population (annual rate of growth 2%)  $\Rightarrow$  daily birth and death rates b = 0.00009889 and d = 0.0000411 per person, respectively

## Parameter assumptions – Force of infection

No H2H transmission ⇒ incidence proportional to number of susceptible individuals and force of infection, which quantifies combined effect of all variables involved in infection process:

- degree of soil contamination with Clostridium tetani
- climate
- frequency of lesions
- proportion of rural population
- socioeconomic conditions
- level of medical care for the wounded and during deliveries

Force of infection acting on newborn  $(\lambda_b)$  and susceptible population  $(\lambda)$  fixed at 3 different levels adequate for reproducing the following stable annual incidence rates of tetanus cases in the community

- For newborn, 200 cases, 400 cases and 600 cases per 100,000 newborn
- For general population (without newborn), 9, 18 and 27 cases

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### Recall the base model of Capasso

$$E' = c_H H - d_E E$$

$$H' = g(E) - \gamma_H H$$

$$c_H H$$

$$g(E)$$

$$g(E)$$
Human population
$$Environment$$
(2a)

 $1/\gamma_H$  mean infectious period,  $1/d_E$  mean lifetime of the agent in the environment,  $c_H$  growth rate of the agent due to the human population, g(E) incidence of the agent on human population

#### Incidence function

$$g(E) = h(E)N\beta p \tag{3}$$

where

- h(E) probability for an exposed susceptible to get the infection
- N total human population
- $\triangleright$   $\beta$  fraction of susceptible individuals in N
- p fraction exposed to contaminated environment per unit time ("probability per unit time to have a "snack" of contaminated food")

Typically, we would assume p and  $\beta$  independent of E and H and h to be saturating. We take a Holling type II functional response

$$h(E) = h_{max} \frac{E}{h_{half} + E} \tag{4}$$

Let

$$\mathcal{R}_0 = \frac{g'_+(0)c_H}{d_F\gamma_H} \tag{5}$$

#### Theorem 1

- $\blacktriangleright$  If  $0 < \mathcal{R}_0 < 1$ , then (2) admits only the trivial equilibrium in the positive orthant, which is GAS
- ▶ If  $\mathcal{R}_0 > 1$ , then two EP exist: (0,0), which is unstable, and  $z^* = (E^*, H^*)$  with  $E^{\star}, H^{\star} > 0$ , GAS in  $\mathbb{R}^2_{\perp} \setminus \{0, 0\}$

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## Simulating (in R) – Incidence function

```
h = function(E, params) {
    # Use Michaelis Menten (Holling type II) growth
    OUT = params$g_max * E / (params$g_half+E)
    return(OUT)
}
g = function(E, params) {
    OUT = params$N * params$beta * params$p * h(E,params)
    return(OUT)
}
```

## The right hand side

```
rhs_Capasso_ODE = function(t, x, params) {
   with(as.list(c(x, params)), {
     dE = c_H*H-d_E*E
     dH = g(E, params)-gamma_H*H
     list(c(dE, dH))
   })
}
```

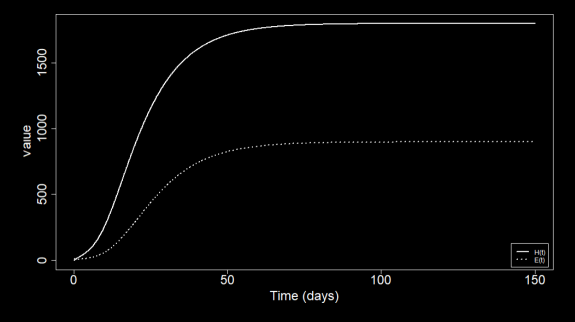
### **Setting parameters**

```
params = list()
params$N = 1000  # Total population
params$gamma_H = 1/10 # Infectious period
params$d_E = 1/5  # Lifetime agent
params$c_H = 0.1  # Flow from humans
params$beta = 0.2 # Fraction susceptible
params$p = 0.1  # Probability of having "snack"
params$g_max = 10
params$g_half = 100
params$t_f = 150
```

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## Running and plotting (base)

```
IC \leftarrow c(E = 10, H = 0)
tspan = seq(from = 0, to = params$t_f, by = 0.1)
sol_ODE = ode(v = IC,
               func = rhs_Capasso_ODE.
               times = tspan,
               parms = params)
plot(sol_ODE[,"time"], sol_ODE[,"H"],
      type = "1", 1 \text{wd} = 2.
      xlab = "Time_(days)", vlab = "Value")
lines(sol ODE[,"time"], sol ODE[,"E"],
      lwd = 2, ltv = 3)
legend("bottomright", legend = c("H(t)", "E(t)"),
        1 \text{wd} = c(2,2), 1 \text{ty} = c(1,3), inset = 0.01)
```



## Adding a periodic component

Assume p in (3) takes the form

$$p(t) = p(t + \omega) > 0, \quad t \in \mathbb{R}$$
 (6)

i.e., p has period  $\omega$ . So we now consider the incidence

$$g(t,E) = p(t)h(E) \tag{7}$$

with h having the properties prescribed earlier. Letting

$$p_{min} := \min_{0 \le t \le \omega} p(t), \quad p_{max} := \max_{0 \le t \le \omega} p(t)$$
 (8)

then we require that

$$\lim_{z \to \infty} \frac{g(z)}{z} < \frac{d_E \gamma_H}{c_H p_{max}} \tag{9}$$

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Let

$$\mathcal{R}_0^{min} = \frac{c_H p_{min} h'_+(0)}{d_E \gamma_H}, \quad \mathcal{R}_0^{max} = \frac{c_H p_{max} h'_+(0)}{d_E \gamma_H}$$
(10)

#### Theorem 2

- If  $0 < \mathcal{R}_0^{max} < 1$ , then (2) with incidence (7) always goes to extinction
- If  $\mathcal{R}_0^{min} > 1$ , then a unique nontrivial periodic endemic state exists for (2) with incidence (7)

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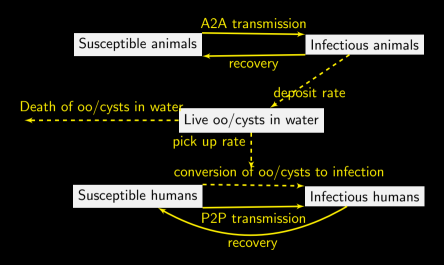
A model for zoonotic transmission of waterborne disease

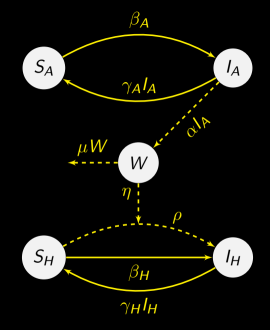
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### Zoonotic transmission of waterborne disease

Waters, Hamilton, Sidhu, Sidhu, Dunbar. Zoonotic transmission of waterborne disease: a mathematical model. Bull Math Biol (2016) Used for instance to model Giardia transmission from possums to humans





### The full model

$$S'_{A} = -\beta_{A}S_{A}I_{A} + \gamma_{A}I_{A}$$
 (11a)  

$$I'_{A} = \beta_{A}S_{A}I_{A} - \gamma_{A}I_{A}$$
 (11b)  

$$W' = \alpha I_{A} - \eta W(S_{H} + I_{H}) - \mu W$$
 (11c)  

$$S'_{H} = -\rho \eta WS_{H} - \beta_{H}S_{H}I_{H} + \gamma_{H}I_{H}$$
 (11d)  

$$I'_{H} = \rho \eta WS_{H} + \beta_{H}S_{H}I_{H} - \gamma_{H}I_{H}$$
 (11e)

Considered with  $N_A = S_A + I_A$  and  $N_H = S_H + I_H$  constant

## Simplified model

Because  $N_A$  and  $N_H$  are constant, (11) can be simplified:

$$I'_{A} = \beta_{A} N_{A} I_{A} - \gamma_{A} I_{A} - \beta_{A} I_{A}^{2}$$

$$W' = \alpha I_{A} - \eta W N_{H} - \mu W$$
(12a)

$$I'_{H} = \rho \eta W(N_{H} - I_{H}) + \beta_{H} N_{H} I_{H} - \gamma_{H} I_{H} - \beta_{H} I_{H}^{2}$$
 (12c)

Three EP: DFE (0,0,0); endemic disease in humans because of H2H transmission; endemic in both H and A because of W

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Three EP: DFE (0,0,0); endemic disease in humans because of H2H transmission; endemic in both H and A because of W

Let

$$\mathcal{R}_{0A} = \frac{\beta_A}{\gamma_A} N_A$$
 and  $\mathcal{R}_{0H} = \frac{\beta_H}{\gamma_H} N_H$  (13)

- ▶ DFE LAS if  $\mathcal{R}_{0A} < 1$  and  $\mathcal{R}_{0H} < 1$ , unstable if  $\mathcal{R}_{0A} > 1$  or  $\mathcal{R}_{0H} > 1$
- ▶ If  $\mathcal{R}_{0H} > 1$  and  $\mathcal{R}_{0A} < 1$ , (12) goes to EP with endemicity only in humans
- ▶ Endemic EP with both A and H requires  $\mathcal{R}_{0A} > 1$  and  $\mathcal{R}_{0H} < 1$

Note that proof is **not** global

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#### A model of Woolhouse

Woolhouse. On the application of mathematical models of schistosome transmission dynamics. I. Natural transmission. Acta Tropica 49:241-270 (1991)

### The model

Population of H individuals using a body of water containing N snails

 $i_H$  mean number of schistosomes per person and  $i_S$  the proportion of patent infections in snails (prevalence)

$$i_{H}^{\prime} = \alpha Ni_{S} - \gamma i_{H} \tag{14a}$$

$$i_S' = \beta H i_H (1 - i_S) - \mu_2 i_S$$
 (14b)

- $ightharpoonup \alpha$  number of schistosomes produced per person per infected snail per unit time
- $ightharpoonup 1/\gamma$  average life expectancy of a schistosome
- $\triangleright$  1/ $\mu_2$  average life expectancy of an infected snail
- $\triangleright$   $\beta$  transmission parameter

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Let the basic reproductive rate for schistosomes be

$$\mathcal{R}_0 = \frac{\alpha N \beta H}{\gamma \mu_2} \tag{15}$$

(14) has two EP

- $(i_H^{\star}, i_S^{\star}) = (0, 0)$ , LAS when  $\mathcal{R}_0 < 1$  and unstable when  $\mathcal{R}_0 > 1$
- $(i_H^{\star}, i_S^{\star}) = \left(\frac{\alpha N}{\gamma} \frac{\mu_2}{\beta H}, 1 \frac{1}{R_0}\right)$ , which only "exists" when  $R_0 > 1$  (and is LAS) then)

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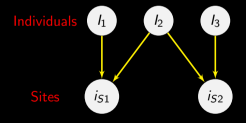
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### Heterogeneities in contact rates

 $l_i$  the number of schistosomes in person  $i=1,\ldots,H$  and  $i_{Si}$  the proportion of patent infected snails in site j = 1, ..., L (L sites each supporting N snails)



 $l_i$  the number of schistosomes in person  $i=1,\ldots,H$  and  $i_{Sj}$  the proportion of patent infected snails in site  $j=1,\ldots,L$  (L sites each supporting N snails)

$$I_i' = \alpha \left( \sum_j \eta_{ij} \mathsf{N} i_{Sj} \right) - \gamma I_i \tag{16a}$$

$$i'_{Sj} = \beta \left( \sum_{i} \eta_{ij} I_i \right) (1 - i_{Sj}) - \mu_2 i_{Sj}$$
 (16b)

 $\eta_{ii}$  rate of water contact by individual i at site j

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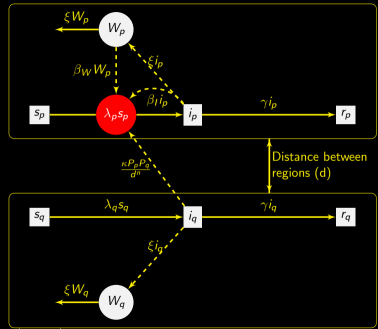
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## Spatial aspects – Cholera in Haiti

Tuite, Tien, Eisenberg, Earn, Ma & Fisman, Cholera Epidemic in Haiti, 2010: Using a Transmission Model to Explain Spatial Spread of Disease and Identify Optimal Control Interventions. Annals of Internal Medicine 154(9) (2011)



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#### Metapopulation model with implicit movement

$$s'_{p} = \mu - \lambda_{p} s_{p} - \mu s_{p}$$

$$i'_{p} = -\gamma i_{p} + \lambda_{p} s_{p} - \mu i_{p}$$

$$r'_{p} = \gamma r_{p} - \mu r_{p}$$

$$w'_{p} = \xi (i_{p} - w_{p})$$
(17a)
$$(17b)$$

$$(17c)$$

with force of infection

$$\lambda_p = \beta_{i_p} i_p + \beta_{W_p} w_p + \sum_{q=1}^{10} \theta_{pq} i_q \tag{17e}$$

Influence of infection prevalence in q on incidence in p is gravity-type

$$\theta_{pq} = \kappa \frac{P_p P_q}{d^n}$$

