Revisiting the early stage of the 1720 plague epidemic in Gévaudan

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8 October 2024

Preamble

In this report, we revisit some of the key elements of the early stages of the bubonic Plague epidemic in Gévaudan, which started in 1720 in the village of Correjac, soon followed by the town of La Canourgue. Historical data suggest that the epidemic was initiated by a convict who travelled from Marseille to Saint-Laurent-d'Olt, where he infected Jean Quintin, who then seeded the epidemic which would first affect his village before spreading to the rest of Gévaudan.

We re-analyse this scenario by combining historical data on the dates of deaths of the first few cases, alongside published estimates of the incubation time or infectious period distributions. A branching process model with augmented data is used for jointly estimating the effective reproduction number and the rate of zoonotic introductions in the early stages of the epidemic.

Estimates of key epidemiological features

Delay distributions

Incubation period

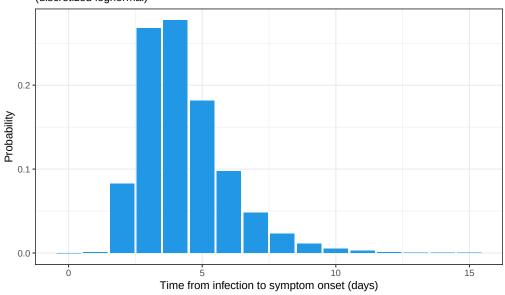
The incubation period is described in multiple papers but a general consensus seems to be around 2-6 days [1–7].

We build a discretized log-normal distribution compatible with these observations:

```
library(distcrete)
library(tidyverse)
incub <- distcrete(</pre>
  "lnorm",
  meanlog = log(3.5),
  sdlog = log(1.5),
  interval = 1,
  w = 1
  )
incub_dat <- tibble(</pre>
  Day = 0:15,
  p = incub $d(0:15)
incub_dat %>%
  ggplot(aes(x = Day, y = p)) +
  geom col(fill = 4) +
  theme bw() +
```

```
labs(
    x = "Time from infection to symptom onset (days)",
    y = "Probability",
    title = "Incubation time distribution - bubonic plague",
    subtitle = "(discretized lognormal)")
```

Incubation time distribution – bubonic plague (discretized lognormal)



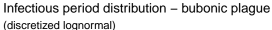
Infectious period

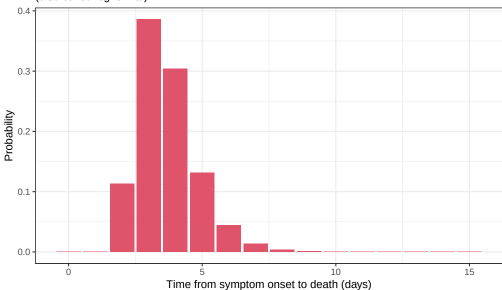
The infectious period distribution is built similarly to match data from the literature, where for historical outbreaks death is reported to take place within 3 to 5 days post-symptom onset [1,6,8,9].

Note that because this form had near 100% CFR, this is also the distribution of the time from onset of symptoms to death.

```
infec <- distcrete(</pre>
  "lnorm",
  meanlog = log(3),
  sdlog = log(1.4),
  interval = 1,
  w = 1
  )
infec_dat <- tibble(</pre>
  Day = 0:15,
  p = infec d(0:15)
  )
infec_dat %>%
  ggplot(aes(x = Day, y = p)) +
  geom_col(fill = 2) +
  theme_bw() +
  labs(
    x = "Time from symptom onset to death (days)",
    y = "Probability",
```

```
title = "Infectious period distribution - bubonic plague",
subtitle = "(discretized lognormal)")
```





We note that the distribution is well in line with the mean duration of fatal, untreated bubonic plague of 3.6 days [1]:

```
## mean of 100 000 values from the distribution:
mean(infec$r(1e5))
```

[1] 3.67142

Patient zero: the convict from Marseille

The theory of the convict walking all the way from Marseille with an infected bundle of wool to eventually infect Jean Quintin in Saint Laurent d'Olt is suspicious: the man would have had to not get infected during his entire trip, to then infect JQ after a fairly brief encounter.

The trip by foot via current roads is 273 km, and would have likely been longer using roads at the time. Especially since patrols were restricting movement in the area, and would have demanded some extra time to be avoided.

We will explore 3 scenarios, with varying durations for the trip, which we posit was at least about 300 km:

- 7 days (very optimistic, > 40 km per day)
- 12 days (~ 25km per day)
- 20 days (~ 15 km per day)

We do not know what the daily rate of infection λ from the wool bundle was, but we can derive a likelihood profile for each scenario, using:

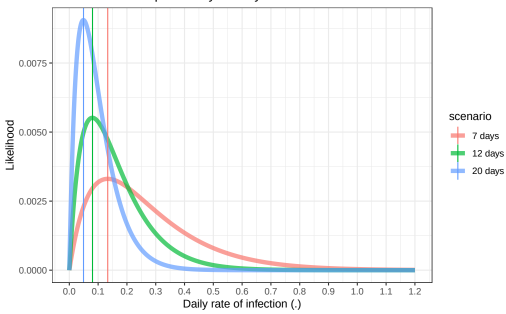
$$\mathcal{L}(\lambda) = p(T|\lambda) = (e^{-\lambda})^T (1 - e^{-\lambda})$$

where T is the number of days the trip took, and $(1 - e^{-\lambda})$ is the daily probability of infection from the infected bundle.

Let us look at these profiles, ensuring we re-standardise both densities to 1 to make them comparable, as they rely on datasets of different sizes:

```
like_trip <- function(lambda, T) {</pre>
  p \leftarrow 1 - exp(-lambda)
  (1 - p)^T * p
lambda res <- tibble(</pre>
  val = seq(0, 1.2, length = 1000),
  "7 days" = like_trip(val, 7),
  "12 days" = like_trip(val, 12),
  "20 days" = like_trip(val, 20)
) %>%
  mutate_at(
    vars(contains("days")),
    function(x) x / sum(x)
  )
lambda_res_long <- lambda_res %>%
  pivot_longer(-1, names_to = "scenario", values_to = "likelihood") %>%
  mutate(scenario = factor(scenario, levels = paste(c(7, 12, 20), "days")))
lambda_res_mle <- lambda_res_long %>%
  group_by(scenario) %>%
  summarise(MLE = val[which.max(likelihood)])
lambda_res_long %>%
  ggplot(aes(x = val, y = likelihood, color = scenario)) +
  geom_line(linewidth = 2, alpha = 0.7) +
  geom_vline(data = lambda_res_mle, aes(xintercept = MLE, color = scenario)) +
  theme_bw() +
  labs(
    x = "Daily rate of infection (\U03BB)",
    y = "Likelihood",
    title = "Estimation of the probability of daily infection"
  scale_x_continuous(n.breaks = 20)
```

Estimation of the probability of daily infection



lambda_res_mle

```
## # A tibble: 3 x 2
## scenario MLE
## <fct> <dbl>
## 1 7 days 0.133
## 2 12 days 0.0805
## 3 20 days 0.0492
```

We find some more likely values than others, and this leans towards fairly low rates ranging from 0.0492 to 0.1333. We calculate the corresponding probabilities that these events occurred using the MLE:

```
lambda_res_mle <- lambda_res_mle %>%
  mutate(
    days = as.integer(sub(" days", "", scenario)),
    proba = like_trip(lambda = MLE, T = days)
    )
lambda_res_mle
```

```
## # A tibble: 3 x 4
##
     scenario
                 MLE
                      days
                           proba
##
     <fct>
               <dbl> <int>
                            <dbl>
## 1 7 days
              0.133
                         7 0.0491
## 2 12 days 0.0805
                        12 0.0294
                        20 0.0179
## 3 20 days 0.0492
```

The resulting probabilities are quite low, ranging from 1.8% chances for a trip of 20 days, to 4.9% for 7 days.

To assess the overall plausibility of these scenari, questions remain: how many such trips where made by people carrying the infection from Marseille at the time? And were such trips indeed possible in the first place, given the containment measures in place?

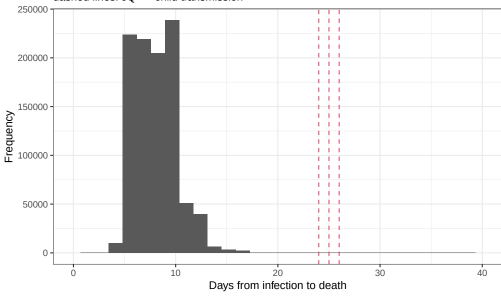
The first case after Jean

calculate possible delays from infection to death

Jean Quintin showed symptoms on the 23rd November, and died 3 days later on the 26th. The first case in his household died on the 18th December. We can assess how likely this delay is by looking at the delay distributions for Bubonic plague, integrating over the possible dates of infection (24th, 25th, 26th November), and convolving the incubation period, and the duration of the symptomatic period.

```
child_delay <- as.integer(as.Date("1720-12-18") - as.Date("1720-11-24") + 0:2)
child_delay
## [1] 24 25 26
## convolve using 1e6 draws from distributions
sim_delay_inf_death <- incub$r(1e6) + infec$r(1e6)</pre>
## calculate p-values
child_delay_pval <- sapply(child_delay, function(x) mean(sim_delay_inf_death >= x))
child_delay_pval
## [1] 1.8e-05 7.0e-06 4.0e-06
mean(child_delay_pval)
## [1] 9.666667e-06
qplot(sim_delay_inf_death) +
  theme_bw() +
  geom_vline(xintercept = child_delay, color = 2, linetype = 2) +
 xlim(0, 40) +
  labs(
   title = "Simulated delays from infection to death - bubonic plague",
   subtitle = "dashed lines: JQ -> child transmission",
   x = "Days from infection to death",
   y = "Frequency"
```

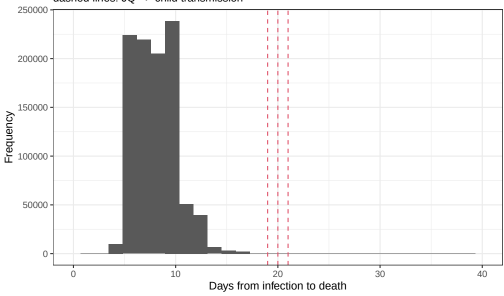
Simulated delays from infection to death – bubonic plague dashed lines: JQ –> child transmission



It is very unlikely that the first secondary case was directly infected by Jean Quintin. There remains the possibility that infected surfaces (clothes, bedsheets) were infected by Y. pestis, which has been shown to be able to last up to 5 days [10]. We can replicate the same analysis subtracting these 5 days from the overal delay:

```
qplot(sim_delay_inf_death) +
  theme_bw() +
  geom_vline(xintercept = child_delay - 5, color = 2, linetype = 2) +
  xlim(0, 40) +
  labs(
    title = "Simulated delays from infection to death - bubonic plague",
    subtitle = "dashed lines: JQ -> child transmission",
    x = "Days from infection to death",
    y = "Frequency"
  )
```

Simulated delays from infection to death – bubonic plague dashed lines: JQ –> child transmission



```
child_delay_pval_alt <- sapply(child_delay - 5, function(x) mean(sim_delay_inf_death >= x))
child_delay_pval_alt
```

```
## [1] 0.000399 0.000205 0.000110
mean(child_delay_pval_alt)
```

```
## [1] 0.000238
```

It seems we can rule out the hypothesis of a direct transmission of bubonic plague. Possible aternatives are:

- it was a bubonic plague, with a long duration of illness in the kid; might be unlikely seeing that patients seem to die very quickly TBC
- the kid was **not** infected by his dad, but from a zoonotic source

Modelling the early epidemic

Epidemic curve

Here we analyse the linelist of the first few cases in Corréjac, followed by La Canourgue.

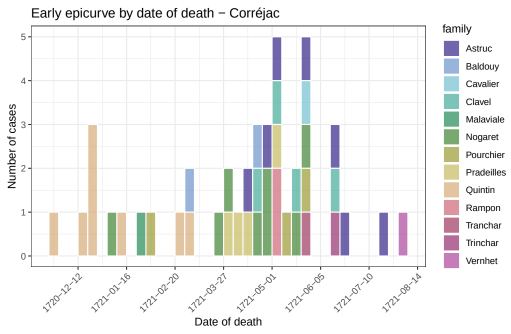
```
file_path <- here::here("data", "early_linelist.ods")</pre>
x <- rio::import(file_path, sheet = "linelist") %>%
  tibble() %>%
  mutate(date_of_death = as.Date(date_of_death, format = "%d/%m/%Y"))
X
## # A tibble: 52 x 4
##
      name
                           date_of_death location family
##
      <chr>
                                                   <chr>
                           <date>
                                          <chr>
##
   1 QUINTIN Jean
                           1720-11-26
                                          Corréjac Quintin
                                         Corréjac Quintin
   2 QUINTIN Ambroise
##
                           1720-12-18
   3 CASSANHES Marguerite 1720-12-25
                                         Corréjac Quintin
##
## 4 QUINTIN enfant 1
                           1720-12-25
                                         Corréjac Quintin
##
  5 QUINTIN enfant 2
                           1720-12-25
                                         Corréjac Quintin
##
  6 NOGARET Pierre
                           1721-01-04
                                         Corréjac Nogaret
   7 DEROUCH Jean
##
                           1721-01-09
                                         Corréjac Quintin
## 8 MALAVIALE Marguerite 1721-01-28
                                          Corréjac Malaviale
## 9 POURCHIER Antoine
                           1721-02-02
                                          Corréjac Pourchier
## 10 DEROUCH Jean fils
                           1721-02-22
                                          Corréjac Quintin
```

We build a simple epidemic curve for Corréjac, and for both locations together:

i 42 more rows

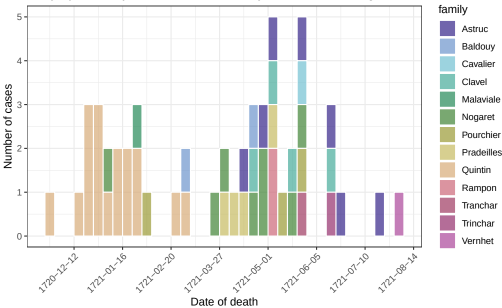
```
library(incidence2)

x %>%
  filter(location == "Corréjac") %>%
  incidence("date_of_death", groups = "family", interval = 7) %>%
  plot(fill = "family", colour_palette = muted, angle = 45, border = "white") +
  labs(
    title = "Early epicurve by date of death - Corréjac",
    x = "Date of death",
    y = "Number of cases"
  )
```



```
x %>%
incidence("date_of_death", groups = "family", interval = 7) %>%
plot(fill = "family", colour_palette = muted, angle = 45, border = "white") +
labs(
   title = "Early epicurve by date of death - Corréjac and La Canourgue",
   x = "Date of death",
   y = "Number of cases"
)
```

Early epicurve by date of death - Corréjac and La Canourgue



There is one key question here: do we assume Corréjac and La Canourgue were disconnected at the time, or do we treat these as a single location.

A transmission model for bubonic plague

A branching process will likely be best at capturing small fluctuations in case incidence over time, whilst neglecting the impact of the depletion of susceptible individuals. However we need to allow for a background rate of zoonotic introduction in addition to the classical person-to-person transmission [11]. As such, we will be using a Hawkes process [12] to model the case incidence over time, using augmented data to make up for the lack of precise information on dates of symptom onset and infection.

Notations

Data and augmented data

- i = 1, ..., n: index of individuals
- t = 1, ..., T: time index
- d_i : date of death of case i (data)
- O_i : date of symptom onset of case i (augmented data)
- I_i : date of infection of case i (augmented data)
- Y_t : incidence of new infections at time t (augmented data, derived from I_i)

Distributions

• \mathcal{F} : probability mass function (pmf) of the infectious/symptomatic period distribution

- \mathcal{G} : pmf of the incubation period distribution
- \mathcal{H} : pmf of the generation time distribution, obtained from the convolution $\mathcal{F} * \mathcal{G} = \mathcal{H}$

Parameters

- λ_z : the rate of zoonotic introduction, assumed constant over time
- R: the effective reproduction number, assumed constant over time

The model

We use a classical Bayesian framework where the posterior distribution is defined for parameters θ and data (x) as:

$$p(\theta|x) \propto p(x|\theta)p(\theta)$$
 (1)

where $p(x|\theta)$ is the likelihood function and $p(\theta)$ the prior distributions.

The likelihood can be written as:

$$p(x|\theta) = p(d, O, I|\lambda_z, R)$$
(2)

$$= p(d|O)p(O|I)p(I|\lambda_z, R)$$
(3)

$$= \left(\prod_{i} p(d_i|O_i) \prod_{i} p(O_i|I_i)\right) p(I|\lambda_z, R) \tag{4}$$

$$= \left(\prod_{i} \mathcal{F}(d_{i} - O_{i})\mathcal{G}(O_{i} - I_{i})\right) p(I|\lambda_{z}, R)$$
(5)

The calculation of $p(I|\lambda_z, R)$ is defined by the Hawkes process for the incidence Y:

$$p(I|\lambda_z, R) = p(Y|\lambda_z, R) \tag{6}$$

$$= p(Y_t | \lambda_z, R)$$

$$= \prod_t p(Y_t | Y_1, ..., Y_{t-1}, \lambda_z, R)$$
(6)
(7)

The incidence Y_t is governed by:

$$Y_t \sim \mathcal{P}(\lambda_t)$$
 (8)

where $\mathcal{P}(.)$ is the Poisson distribution, and with:

$$\lambda_t = \lambda_z + \sum_{s=1}^{t-1} RY_s \mathcal{H}(t-s) \tag{9}$$

Finally, we assume independent priors for λ_z and R such that:

$$p(\theta) = p(\lambda_z, R) = p(\lambda_z)p(R) \tag{10}$$

Estimation process

We can sample from the posterior distribution using the Metropolis algorithm with augmented data with the following process:

- 1. draw augmented data O using $d_i O_i \sim \mathcal{F}$
- 2. draw augmented data I using $O_i I_i \sim \mathcal{G}$

- 3. propose (using symmetric proposal distributions) new values for θ^* and accept/reject these values with probability: $max(1, \frac{p(\theta^*|x)}{p(\theta|x)})$, where θ represents the previous parameter state; in practice, separate movements are used for λ_z and R, using Normal proposal distributions with standard deviations manually tailored to reach about 30-40% acceptance rates
- 4. go back to 1 until desired number of iterations reached

Distributions

Generation time

The generation time distribution \mathcal{H} is estimated using the following procedure:

- 1. Sample a large number n of incubation periods X
- 2. Sample a large number n of infectious period durations Y
- 3. Sample n delays from onset to infections Z, uniformly distributed between 0 and Y
- 4. Derive the empirical distribution of \mathcal{H} from X+Z

Here the estimates are based on 10 million draws:

```
## RNG for delay from onset to infection
onset_to_inf_r <- function(n) {</pre>
  floor(runif(n, 0, infecred{s}r(n) + 1))
## PMF for delays of 0, 1, ...
gentime_dat <- incub$r(1e7) + onset_to_inf_r(1e7)</pre>
range(gentime_dat)
## [1] 1 35
gentime_freq <- sapply(</pre>
  seq(0, max(gentime_dat), by = 1L),
  function(i) sum(i == gentime_dat)
gentime_pmf <- gentime_freq</pre>
gentime pmf <- gentime pmf / sum(gentime pmf) # superfluous
## emulate density function
gentime_d <- function(x, log = FALSE) {</pre>
  ## make sure we don't get out of bounds
  x <- as.integer(x)</pre>
  max_x <- max(gentime_dat)</pre>
  x[x < 0] <- 0
  x[x > max_x] \leftarrow 0
  out <- gentime_pmf[x+1]</pre>
  if (log) {
    out <- log(out)
  }
  out
}
## emulate rng function
gentime r <- function(n) {</pre>
  sample(gentime_dat, size = n, replace = TRUE)
}
```

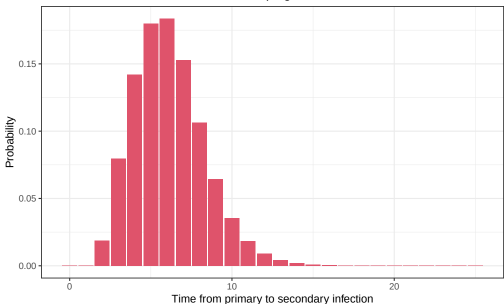
```
gentime <- list(d = gentime_d, r = gentime_r)

gt_dat <- tibble(
   Day = 0:25,
   p = gentime$d(0:25)
)

gt_dat %>%

ggplot(aes(x = Day, y = p)) +
   geom_col(fill = 2) +
   theme_bw() +
   labs(
    x = "Time from primary to secondary infection",
    y = "Probability",
    title = "Generation time distribution - bubonic plague"
   )
```

Generation time distribution - bubonic plague



Priors

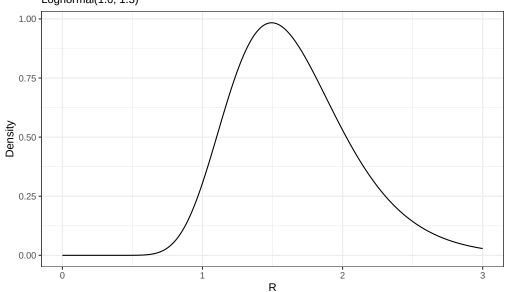
We use a prior for R derived from [8] as a lognormal distribution with mean 1.6 and standard deviation 1.3:

```
prior_R <- function(x, log = FALSE) dlnorm(x, log(1.6), log(1.3), log = log)
prior_R_dat <- tibble(
    x = seq(0, 3, length.out = 1000),
    d = prior_R(seq(0, 3, length.out = 1000))
)

ggplot(prior_R_dat, aes(x = x, y = d)) +
    geom_line() +
    theme_bw() +
    labs(
        x = "R",
        y = "Density",
        title = "Prior for R",</pre>
```

```
subtitle = "Lognormal(1.6, 1.3)"
)
```

Prior for R Lognormal(1.6, 1.3)



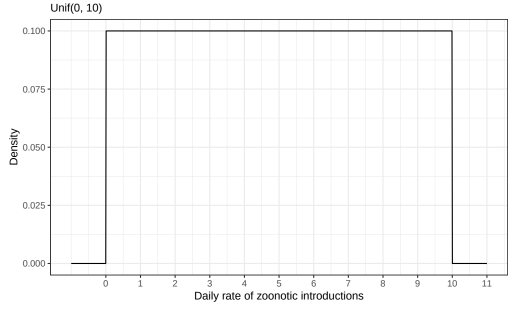
Rates of zoonotic introductions are usually harder to estimate, so we use a flat, uninformative priors for λ_z , uniformly distributed from 0 to 10:

```
prior_zoo <- function(x, log = FALSE) dunif(x, 0, 10, log = log)

prior_zoo_dat <- tibble(
    x = seq(-1, 11, length.out = 1000),
    d = prior_zoo(seq(-1, 11, length.out = 1000))
)

ggplot(prior_zoo_dat, aes(x = x, y = d)) +
    geom_line() +
    theme_bw() +
    labs(
        x = "Daily rate of zoonotic introductions",
        y = "Density",
        title = "Prior for zoonotic introductions",
        subtitle = "Unif(0, 10)"
    ) +
    scale_x_continuous(breaks = 0:11)</pre>
```

Prior for zoonotic introductions



Implementation

The following code implements the model.

```
## Function to generate augmented data
## This will return a list with all the data needed for likelihood calculation.
## Note that dates of deaths are converted to integers, with 0 the earliest death.
make_aug_data <- function(date_death = x$date_of_death,</pre>
                            r_incub = incub$r,
                            r_infec = infec$r
                            ) {
  if (any(is.na(date_death))) {
    msg <- "Some dates of death are missing"</pre>
    stop(msg)
  }
  n <- length(date_death)</pre>
  date_death <- as.integer(date_death - min(date_death))</pre>
  date_onset <- date_death - r_infec(n)</pre>
  date_infection <- date_onset - r_incub(n)</pre>
  out <- data.frame(</pre>
    infection = date_infection,
    onset = date_onset,
    death = date_death
  )
  ## Use this to have 0 as the earliest date; otherwise we do get some negative
  ## dates:
  ## data.frame(lapply(a, function(x) x - min(a)))
  out
}
```

```
## Function to calculate the log-likelihood
## Oparam data: a data.frame with 3 columns as returned by make_aug_data()
## @param params: a list with two items: zoo, and R, in this order
## @param d_incub: a PMF function for the incubation period
## @param d_infec: a PMF function for the infectious period
## Oparam d_gentime: a PMF function for the generation time
compute_loglike <- function(data,</pre>
                             params,
                             d_incub = incub$d,
                             d_infec = infec$d,
                             d_gentime = gentime$d) {
  ### There are 3 components to the likelihood: ^
  ###
  ### 1. delay from onset to death (infectious period)
  ### 2. delay from infection to onset (incubation time)
  ### 3. incidence of infections from Hawkes process
  ### Component 1
  p_1 <- sum(d_infec(data$death - data$onset, log = TRUE))</pre>
  ### Component 2
 p_2 <- sum(d_incub(data$onset - data$infection, log = TRUE))</pre>
  ### Component 3
  first_day <- min(data$infection)</pre>
  last_day <- max(data$infection)</pre>
  \#\#\# calculate incidence for the whole time period, do no miss the zeros
  incid <- sapply(</pre>
    seq(first_day, last_day, by = 1),
    function(i) sum(data$infection == i)
  #### get relative FOIs - check that w indeed start at 1 in EpiEstim
  lambdas p2p <- EpiEstim::overall infectivity(incid, d gentime(0:100))
  lambdas <- params$zoo + (lambdas_p2p * params$R)</pre>
 p_3 <- sum(na.omit(dpois(incid, lambdas, log = TRUE)))</pre>
 p_1 + p_2 + p_3
## Function to calculate log-priors
compute_priors <- function(params,</pre>
                            d_zoo = prior_zoo,
                            d_R = prior_R) {
 d_zoo(params$zoo, log = TRUE) + d_R(params$R, log = TRUE)
## Compute log-posterior
```

```
compute_post <- function(data,</pre>
                          d_incub = incub$d,
                          d_infec = infec$d,
                          d_gentime = gentime$d,
                          d_zoo = prior_zoo,
                          d_R = prior_R
  compute_loglike(data, params, d_incub, d_infec, d_gentime) +
    compute_priors(params, d_zoo, d_R)
}
## Function implementing the whole MCMC procedure
## @param n_iter the number of iterations of the MCMC; defaults to 1000
## @param sd_zoo the standard deviation of the normal proposal distribution for
   the rate of zoonotic introductions
## @coaram sd_R the standard deviation of the normal proposal distribution for
## the effective reproduction number
## Cparam ini_zoo the initial value of the daily rate of zoonotic introduction
## @param ini_R the initial value of the effective reproduction number
estimate_params <- function(date_death = x$date_of_death,</pre>
                             n_{iter} = 1e3,
                             d_incub = incub$d,
                             d_infec = infec$d,
                             d gentime = gentime$d,
                             d_zoo = prior_zoo,
                             d_R = prior_R,
                             r_incub = incub$r,
                             r_infec = infec$r,
                             sd_zoo = 0.1,
                             sd_R = 0.4,
                             ini_zoo = runif(1, 0, 1),
                             ini_R = runif(1, 0, 3)) {
  ### Initialize MCMC
  params <- list(zoo = ini_zoo, R = ini_R)</pre>
  aug_data <- make_aug_data(date_death, r_incub, r_infec)</pre>
  ini_post <- compute_post(aug_data,</pre>
                            params,
                            d_incub,
                            d infec,
                            d_gentime,
                            d_zoo,
                            d R)
  new_params <- current_params <- params</pre>
  ### Build output structure
  mcmc <- list(</pre>
   step = seq_len(n_iter),
    post = double(n_iter),
    zoo = double(n_iter),
```

```
R = double(n_iter)
accept_zoo <- 0</pre>
accept_R <- 0
for (i in seq_len(n_iter)) {
  #### make new augmented data
  aug_data <- make_aug_data(date_death, r_incub, r_infec)</pre>
  current_post <- compute_post(aug_data,</pre>
                                   current_params,
                                   d_incub,
                                   d_infec,
                                  d_gentime,
                                   d_zoo,
                                   d_R)
  #### propose new zoo
  new_params$zoo <- current_params$zoo + rnorm(1, sd = sd_zoo)</pre>
  #### accept/reject zoo
  new_post <- compute_post(aug_data,</pre>
                              new_params,
                              d_incub,
                              d infec,
                              d_gentime,
                              d zoo,
                              d R)
  p_accept_zoo <- exp(new_post - current_post)</pre>
  if (runif(1) <= p_accept_zoo) { # accept move</pre>
    current_params$zoo <- new_params$zoo</pre>
    current_post <- new_post</pre>
    accept_zoo <- accept_zoo + 1</pre>
  } else { # reject move
    new_params$zoo <- current_params$zoo</pre>
  #### propose new R
  new_params$R <- current_params$R + rnorm(1, sd = sd_R)</pre>
  #### accept/reject R
  new_post <- compute_post(aug_data,</pre>
                              new_params,
                              d incub,
                              d_infec,
                              d_gentime,
                              d_zoo,
                              d_R)
  p_accept_R <- exp(new_post - current_post)</pre>
  if (runif(1) <= p_accept_R) { # accept move</pre>
    current_params$R <- new_params$R</pre>
    current_post <- new_post</pre>
    accept_R <- accept_R + 1</pre>
```

```
} else { # reject move
      new_params$R <- current_params$R</pre>
    ### store info from this iteration
    mcmc$post[i] <- current_post</pre>
    mcmc$zoo[i] <- current_params$zoo</pre>
    mcmc$R[i] <- current_params$R</pre>
  }
  list(
    mcmc = data.frame(mcmc),
    accept_zoo = accept_zoo / n_iter,
    accept_R = accept_R / n_iter
  )
}
We run the chains for a few iterations to check all runs smoothly:
system.time(res <- estimate_params(n_iter = 10))</pre>
##
      user system elapsed
##
      1.44
              0.08
head(res)
## $mcmc
##
      step
                post
                             Z00
## 1
         1 -392.9124 0.37955265 2.332001
## 2
         2 -356.0220 0.21550427 2.125770
## 3
         3 -339.4880 0.06956936 2.125770
         4 -351.9763 0.03599563 2.138901
## 4
        5 -344.0236 0.03599563 2.138901
## 5
         6 -365.6880 0.03599563 2.138901
## 6
## 7
         7 -346.6676 0.03599563 2.171586
        8 -339.9714 0.03599563 2.171586
## 8
## 9
        9 -335.0354 0.03599563 2.171586
## 10
        10 -341.7831 0.04498218 2.171586
##
## $accept zoo
## [1] 0.4
##
## $accept_R
## [1] 0.4
tail(res)
## $mcmc
                 post
##
      step
                             Z00
## 1
         1 -392.9124 0.37955265 2.332001
## 2
         2 -356.0220 0.21550427 2.125770
         3 -339.4880 0.06956936 2.125770
## 3
## 4
         4 -351.9763 0.03599563 2.138901
## 5
         5 -344.0236 0.03599563 2.138901
```

```
## 6
         6 -365.6880 0.03599563 2.138901
## 7
         7 -346.6676 0.03599563 2.171586
## 8
         8 -339.9714 0.03599563 2.171586
## 9
         9 -335.0354 0.03599563 2.171586
## 10
        10 -341.7831 0.04498218 2.171586
##
## $accept zoo
## [1] 0.4
##
## $accept_R
## [1] 0.4
```

Resuts on all data

Parameters are estimated for all data, using 8 separate chains run in parallel (works only on linux), with a burn-in of 100 iterations. For simplicity, chains have been saved in a separate RDS file.

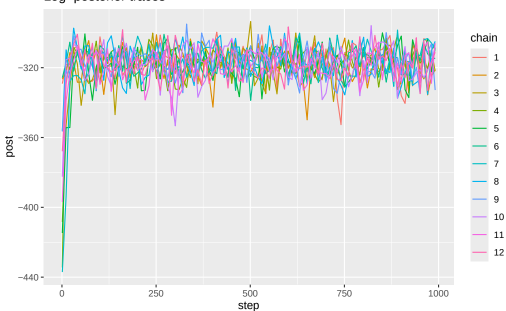
```
## library(parallel)
## res <- mclapply(1:12, function(i)
##
     cbind.data.frame(estimate_params(n_iter = 1000), chain = i),
##
     mc.cores = 12
##
     )
## saveRDS(res, file = "res 1000iter 12chains all data.rds")
res <- readRDS("res_1000iter_12chains_all_data.rds")</pre>
## Some diagnostics
library(coda)
lapply(res, function(e) effectiveSize(mcmc(e))) %>%
  bind_rows()
## # A tibble: 12 x 7
##
      mcmc.step mcmc.post mcmc.zoo mcmc.R accept_zoo accept_R chain
          <dbl>
                                                 <dbl>
                                                          <dbl> <dbl>
##
                    <dbl>
                              <dbl>
                                     <dbl>
```

```
##
               0
                     1000.
                                 134.
                                        172.
                                                                    0
                                                                          0
    1
                                                         0
##
    2
               0
                      896.
                                263.
                                        182.
                                                         0
                                                                    0
                                                                          0
                                  62.7
                                                         0
##
    3
               0
                      150.
                                        256.
                                                                    0
                                                                          0
##
    4
               0
                      398.
                                120.
                                        158.
                                                         0
                                                                    0
                                                                          0
                                 37.8
##
    5
               0
                       78.4
                                        127.
                                                         0
                                                                    0
                                                                          0
##
    6
               0
                     1000
                                151.
                                        169.
                                                         0
                                                                    0
                                                                          0
   7
                                                         0
##
               0
                      105.
                                 51.6
                                        167.
                                                                    0
                                                                          0
##
   8
               0
                      778.
                                191.
                                        181.
                                                         0
                                                                    0
                                                                          0
##
    9
               0
                      611.
                                175.
                                         90.7
                                                         0
                                                                    0
                                                                          0
                                157.
## 10
               0
                      769.
                                        164.
                                                         0
                                                                    0
                                                                          0
## 11
               0
                      164.
                                 51.4 147.
                                                         0
                                                                    0
                                                                          0
               0
                      847.
                                134.
                                        188.
                                                                          0
## 12
```

```
## Putting chains together and thining
## Given the reported ESS a thining of 1/10 seems reasonable
to_keep <- seq(from = 1, to = 1000, by = 10)
chains <- Reduce(rbind, lapply(1:length(res), function(i) res[[i]][to_keep, ]))
names(chains) <- gsub("mcmc.", "", names(chains))
chains$chain <- factor(chains$chain)</pre>
```

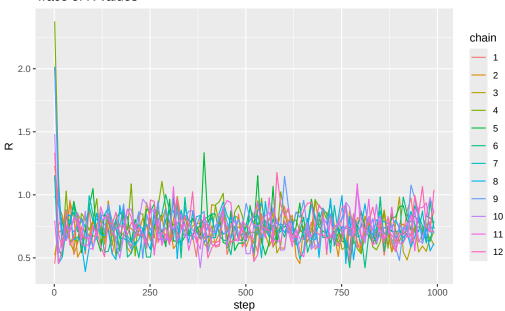
```
## Plots
ggplot(chains) +
  geom_line(aes(x = step, y = post, color = chain)) +
  labs(title = "Log-posterior traces")
```

Log-posterior traces

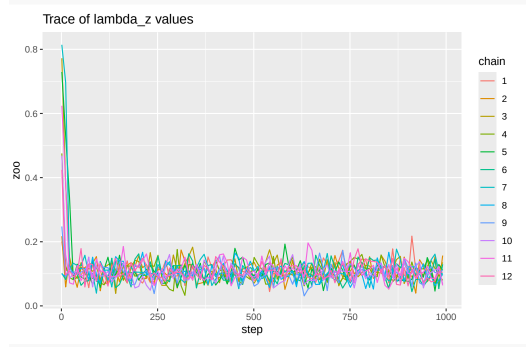


```
## Plots
ggplot(chains) +
  geom_line(aes(x = step, y = R, color = chain)) +
  labs(title = "Trace of R values")
```

Trace of R values

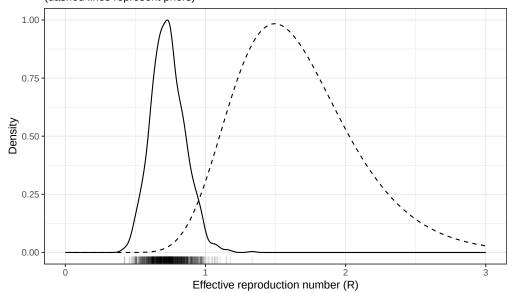


```
ggplot(chains) +
geom_line(aes(x = step, y = zoo, color = chain)) +
labs(title = "Trace of lambda_z values")
```



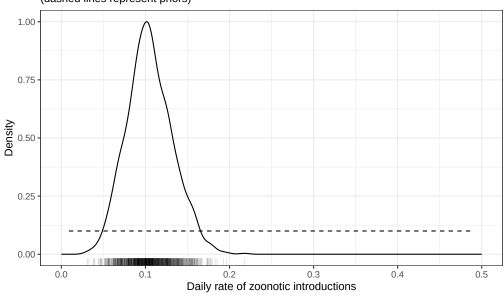
```
chains %>%
  filter(step > 100) %>%
  ggplot(aes(x = R)) +
  stat_density(aes(y = after_stat(scaled)), geom = "line") +
  geom_rug(alpha = .1) +
  geom_line(data = prior_R_dat, aes(x = x, y = d), linetype = 2) +
  theme_bw() +
  labs(
    x = "Effective reproduction number (R)",
    y = "Density",
    title = "Posterior distribution of R",
    subtitle = "(dashed lines represent priors)")
```

Posterior distribution of R (dashed lines represent priors)



```
chains %>%
  filter(step > 100) %>%
  ggplot(aes(x = zoo)) +
  stat_density(aes(y = after_stat(scaled)), geom = "line") +
  geom_rug(alpha = .1) +
  geom_line(data = prior_zoo_dat, aes(x = x, y = d), linetype = 2) +
  theme_bw() +
  labs(
    x = "Daily rate of zoonotic introductions",
    y = "Density",
    title = "Posterior distribution of lambda_z",
    subtitle = "(dashed lines represent priors)") +
  xlim(0, 0.5)
```

Posterior distribution of lambda_z (dashed lines represent priors)



We can report the mean, median and 95% CrI for each parameter:

```
chains_smry <- chains %>%
  filter(step > 100) %>%
  select(R, zoo) %>%
  lapply(function(x) data.frame(
    mean = mean(x),
    median = median(x),
    CrI_low = quantile(x, 0.025),
    CrI_up = quantile(x, 0.975))
    ) %>%
  bind_rows()

rownames(chains_smry) <- c("R", "Zoo")
  chains_smry</pre>
```

```
## mean median CrI_low CrI_up
## R 0.7334454 0.7260987 0.50836176 0.9760880
## Zoo 0.1052512 0.1036642 0.05575248 0.1608936
```

We can also check acceptance rates:

```
chains %>%
  filter(step > 100) %>%
  select(accept_R, accept_zoo) %>%
  summarise_all(mean)
```

```
## accept_R accept_zoo
## 1 0.3160833 0.2944167
```

Results suggest that:

- the data is informative on R and λ_z , with posterior distributions well different from the priors
- R values were most likely below 1, with a mean of 0.73 (95% CrI: 0.51 0.98)
- zoonotic introductions played a substantial role in transmission, with a mean rate of introduction of 0.11~(95%~CrI:~0.06 0.16), corresponding to an average of one introduction every 9.5~days

Conclusions

The statistical analysis of historical data suggests the established scenario of how the Plague epidemic in Gévaudan started may not be accurate. In short:

- It is rather unlikely that an individual would have travel with an infected wood bundle without getting infected the entire trip, to then infect a new person overnight.
- The delays between Jean Quintin's infection and his son's death are not compatible with direct transmission.
- Epidemic modelling suggests that person-to-person transmission was insufficient to sustain the epidemic, and that zoonotic transmission, while low, indeed played a role in the epidemic.

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