



The 2017 plague outbreak in Madagascar: Data descriptions and epidemic modelling

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ABSTRACT

From August to November 2017, Madagascar endured an outbreak of plague. A total of 2417 cases of plague were confirmed, causing a death toll of 209. Public health intervention efforts were introduced and successfully stopped the epidemic at the end of November. The plague, however, is endemic in the region and occurs annually, posing the risk of future outbreaks. To understand the plague transmission, we collected real-time data from official reports, described the outbreak's characteristics, and estimated transmission parameters using statistical and mathematical models. The pneumonic plague epidemic curve exhibited multiple peaks, coinciding with sporadic introductions of new bubonic cases. Optimal climate conditions for rat flea to flourish were observed during the epidemic. Estimate of the plague basic reproduction number *during the large wave of the epidemic* was high, ranging from 5 to 7 depending on model assumptions. The incubation and infection periods for bubonic and pneumonic plague were 4.3 and 3.4 days and 3.8 and 2.9 days, respectively. Parameter estimation suggested that even with a small fraction of the population exposed to infected rat fleas (1/10,000) and a small probability of transition from a bubonic case to a secondary pneumonic case (3%), the high human-to-human transmission rate can still generate a large outbreak. Controlling rodent and fleas can prevent new index cases, but managing human-to-human transmission is key to prevent large-scale outbreaks.

1. Introduction

One of the deadliest natural disasters in human history was reported as the Black Death—attributed to the bacterium *Yersinia pestis*—killing about 50–200 million people in the 14th century (Boire et al., 2014). Although plague was naturally widespread in ancient times, plague outbreaks occurred following the deliberate use and propagation of this disease, serving as a bioweapon (Riedel, 2005). Nowadays, plague epidemics continue to pose a threat to humans, reporting continuous annual occurrence in five countries: Madagascar, Tanzania, Vietnam, China, and the USA (Boire et al., 2014; Dennis et al., 1999). This lethal bacterium can derive in several forms of plague maintaining its existence in a cycle involving rodents and their fleas (Centers for Disease Control and Prevention, 2005). While sanitation and public health surveillance have greatly reduced the likelihood of a plague pandemic, isolated plague outbreaks are lethal threats to humankind.

The disease manifests in different clinical forms of plague: bubonic, pneumonic, and septicemic (Boire et al., 2014). Human infection is primarily driven by bubonic plague, as a result of being bitten by infected fleas. Additionally, direct contamination with infective materials can be an alternative transmission route (Boire et al., 2014). Patients

with bubonic plague can develop sudden onset of fever, headache, chills, tender and painful lymph nodes (World Health Organization, 2017a). While plague can be successfully treated with antibiotics, if untreated, the bacteria can disseminate from the lymph nodes into the bloodstream causing secondary septicemic plague. In addition to the symptoms presented in the bubonic plague, patients with septicemic plague undergo abdominal pain and possibly bleeding into the skin and other organs, at the same time, skin and other tissues may turn black and die, especially on fingers, toes, and the nose (Centers for Disease Control and Prevention, 2005). However, the most fulminant form of the disease is driven by the pneumonic plague that is the only form of plague that can spread from person to person by infectious droplets. The incubation period of primary pneumonic plague is shorter than in the other forms of the disease with an average of 4 days (Raymond Gani, 2004). The disease progresses rapidly and is nearly always fatal without prompt antibiotic treatments (Mead, 2017).

Patients with pneumonic plague often do not transmit the disease to anyone, but in the right conditions one can infect many people and cause an outbreak (Mead, 2017; Kool and Weinstein, 2005). This was observed in the last epidemic in Madagascar when a 31-year-old man travelled from the central highlands to the eastern city of Toamasina via

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the capital city, Antananarivo. He died in transit and dozens of his contacts subsequently became ill (World Health Organization, 2017c). Since then cases of suspected plague have been reported from many areas of Madagascar. On 13 September 2017, the Madagascar Ministry of Public Health notified WHO of an outbreak of pneumonic plague (World Health Organization, 2017a). A total of 2417 cases of plague were confirmed of which 77% were pneumonic plague, causing a death toll of 209 (World Health Organization, 2017c). The Government of Madagascar with the support of WHO and partners had focused their efforts on strengthening the identification and treatment of patients and their contacts, increasing control of rodents and fleas, and practising safe and dignified burials (World Health Organization, 2017b). These measures prevented new cases and deaths, however, the disease is endemic and occurs annually in the region and elsewhere (Boire et al., 2014; Dennis et al., 1999), posing the risk of future outbreaks. This paper provides descriptive and numerical analyses of the plague outbreak to facilitate further studies in evaluating the spread of the plague as well as targets for disease control and prevention.

2. Materials and methods

Outbreak data – cumulative cases. Data were manually inputted from separate reports of WHO (World Health Organization, 2017c), including the cumulative total numbers of clinical cases (confirmed, probable, and suspected). The data and codes can be found at the following link <https://github.com/systemsmedicine/plague2017>. Data can be updated by sending merge requests to this repository.

Outbreak data – by disease forms. Data were digitized from the figure reported from WHO (World Health Organization, 2017c), including the incidences classified by the three forms of the plague disease: pneumonic, bubonic, and septicemic. The data and the digitized figure can also be found in the same repository.

Temperature and precipitation data. Data were requested from the National Centers for Environmental Information (Order #1133340 Custom GHCN-Daily CSV).

Descriptive analyses. With the aim of facilitating modelling works, we described dynamics and patterns of variables that have previously shown to be relevant to plague outbreaks, including temperature and precipitation (Kreppel et al., 2014).

Statistical estimate of the reproduction number. We estimated the reproduction number (R0) of *Yersinia pestis* using data of pneumonic cases (excluding the bubonic cases) during the second (large) wave of the epidemic, i.e., visually defined from 22/09/17 onwards. The serial interval of plague was assumed gamma distributed with shape and scale parameters as 5.4 and 0.9, respectively (Nishiura et al., 2006). We reported the R0 estimates using several methods for comparison purposes: exponential growth (EG) (Wallinga, 2007), maximum likelihood (ML) (White et al., 2009), sequential Bayesian estimation (SB) (Bettencourt and Ribeiro, 2008), binomial assumption (Nishiura, 2010), and sub-exponential growth model (Viboud et al., 2016; Chowell et al., 2016).

Plague transmission model (PTM). We proposed a plague transmission model using a modified SEIR (Susceptible-Exposed-Infectious-Removed) model with two additional incorporated components reflecting the seasonal infection from infected rat fleas and imperfect sigmoidal effects of public health interventions. A schematic illustration of the PTM is shown in Fig. 1. The model equations are as follows:

$$\frac{dS_b}{dt} = -S_b \alpha f_{irf} f_{itv,f} - \beta S_b \frac{I_p}{N} f_{itv,h} \quad (1)$$

$$\frac{dS_p}{dt} = -\beta S_p \frac{I_p}{N} f_{itv,h} \quad (2)$$

$$\frac{dE_b}{dt} = S_b \alpha f_{irf} f_{itv,f} - \gamma_b E_b \quad (3)$$

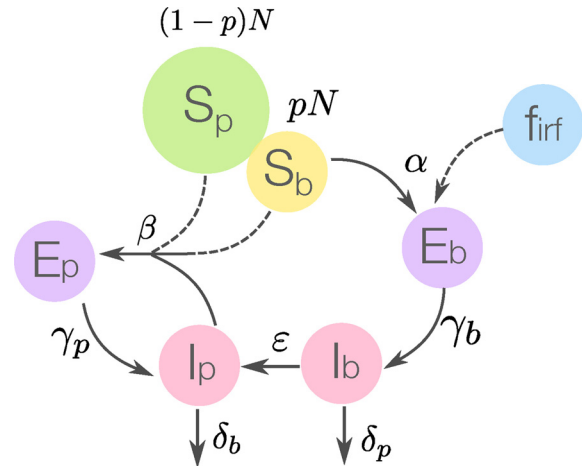


Fig. 1. Schematic of plague transmission model (PTM). Assuming a small proportion (p) of the population is exposed to the risk of being bitten by infected rat fleas. The flea density (f_{irf}) is approximated with a sinusoidal function fitted on Madagascar temperature (see more details in the text of Section 2). Dashed lines indicate parameters which public health interventions would affect.

$$\frac{dE_p}{dt} = \beta \left(S_b + S_p \right) \frac{I_p}{N} f_{itv,h} - \gamma_p E_p \quad (4)$$

$$\frac{dI_b}{dt} = \gamma_b E_b - \epsilon I_b - \delta_b I_b \quad (5)$$

$$\frac{dI_p}{dt} = \gamma_p E_p + \epsilon I_b - \delta_p I_p \quad (6)$$

where S , E , I describe Susceptible, Exposed, and Infectious and the subscripts b and p denote bubonic and pneumonic form, respectively. The model assumes that in a population of size N , only a small part $S_b = pN$ is exposed to infected rat fleas. The infected bubonic cases become infectious with a proportion ϵ progressing to the pneumonic stage (Kool and Weinstein, 2005). The overall transmission rates of the two modes flea-to-human and human-to-human are denoted by α and β , respectively.

It has been shown that climatic conditions favour the survival and reproduction of fleas (Eisen and Gage, 2011; Chotellersak et al., 2015; Shrewsbury, 2005). Fleas density fluctuates by season temperatures, and changes in their host (rats) also depend on the season which relates to the breeding patterns of rats (Varlik, 2015). To this end, we assumed that the density of infected rat fleas can be described as a sinusoidal function $f_{irf} = A + B \sin(2\pi/12t) + C \cos(2\pi/12t)$ (Aron and May, 1982) following the temperature fluctuation; here the average temperature of Madagascar in the period 1960–2008 (Kreppel et al., 2014) was used. As such, the overall flea-to-human transmission parameter α implicitly incorporates a scaling factor from temperature to rat fleas density.

We assumed interventions would reduce flea-to-human and human-to-human transmission rates via rodent and flea control, via active case finding and the identification and prophylaxis of contacts. The interventions effect is assumed *imperfect* and has a logistic form as $f_{itv,h} = 1 - 1/[1 + \theta + \exp(\tau_{h,f} - t)]$, where $\tau > 0$ denotes the time at which the interventions reached the half maximum effect in controlling human-to-human (τ_h) and flea-to-human (τ_f) transmission. The reduction from a perfect intervention is defined by $\theta \geq 0$, where $\theta = 0$ means a perfect scenario. The infected cases are assumed to recover and die with the total rate of removal from the infected pool being δ_b and δ_p for bubonic and pneumonic cases, respectively.

We fitted the PTM to the daily data of pneumonic and bubonic cases during the large wave of the epidemic curve which was visually defined from 22/09/17 onwards. Model parameters were estimated using the

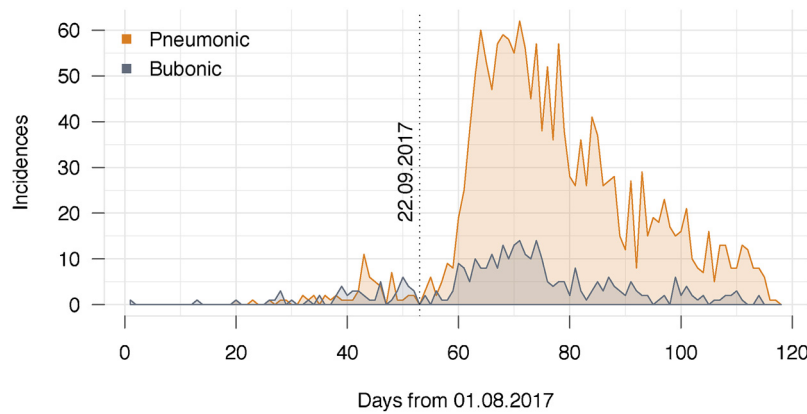


Fig. 2. Plague dynamics August–October 2017. Reported incidences of the two forms of plague diseases during the 2017 outbreak. The data were digitized from WHO report's figure (World Health Organization, 2017a).

global optimisation algorithm Differential Evolution (Mullen et al., 2011). We derived also R_0 using the next generation matrix (Diekmann et al., 2009). Simulations and estimations were written in R using packages base (R Core Team, 2017), deSolve (Soetaert et al., 2010), and R_0 (Obadia et al., 2012) and in Python. Stochastic simulations were performed using a tau-leaping algorithm with a fixed time-step of ca. 15 min.

3. Results

3.1. Descriptive analyses

During August, bubonic cases appeared sporadically with almost no records of the pneumonic form (Fig. 2). An increase in number of pneumonic cases was not necessarily preceded by an increase in the number of bubonic cases (Fig. 2). It seemed to be that the epidemic curves include several waves of incidence overlapping each other.

Fig. 3 shows that plague incidences emerged every day in the weeks though in some weeks fewer cases were reported during the weekends. No distinctive time lag was observed between the appearances of bubonic and pneumonic cases (Fig. 3). The incidences were negligible during the period when the Famadihana tradition was presumably practised. Precipitation measure exhibited no pattern before or during the outbreak but generally showed a dry climatic condition. Average temperature increased and reached a higher level (above 23 °C) around the same time as the outbreak.

Fig. 4 shows that the temperature would typically remain at a level favouring the rat fleas (20–25 °C (Shrewsbury, 2005)) in the upcoming months and until May 2018.

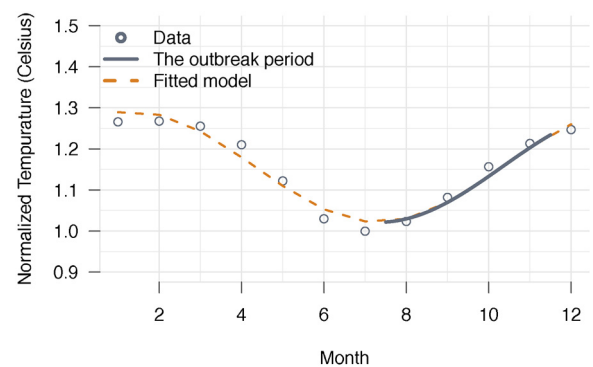


Fig. 4. Fitted sinusoidal of Madagascar temperature. Fitted sinusoidal of Madagascar temperature (see Section 2) with $A = 1.15$, $B = 0.08$, $C = 0.1$. The temperature is normalised by the lowest value of July.

3.2. Models and parameter estimates of plague epidemic

Fig. 5 shows that the model and the estimated parameters (Table 1) capture well the dynamics of both pneumonic and bubonic data. Stochastic transitions could lead to larger or smaller waves of the dynamics. There are epidemic trajectories that resemble the intermittent and small epidemic wave during August. The parameters suggest that only a small fraction exposed to infected rat fleas is enough to generate the observed epidemic. Table 1 shows that several statistical estimation methods gave a similar value for plague's basic reproduction number which is approximately 7. We also estimated R_0 using the PTM.

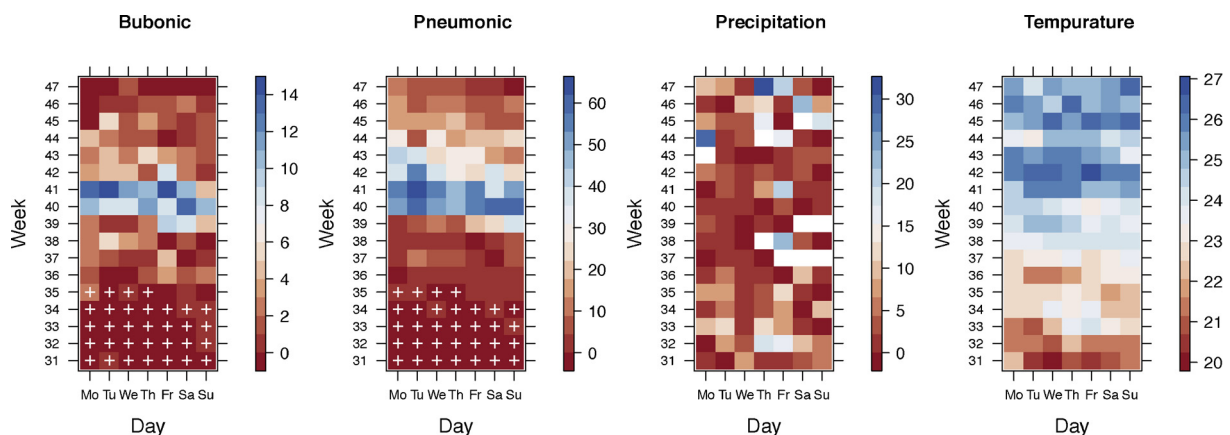


Fig. 3. Incidences and climate variables. Reported incidences and average temperature and precipitation classifying by week and day of the weeks. The “+” signs indicate a supposed time period when the Famadihana tradition is practised.

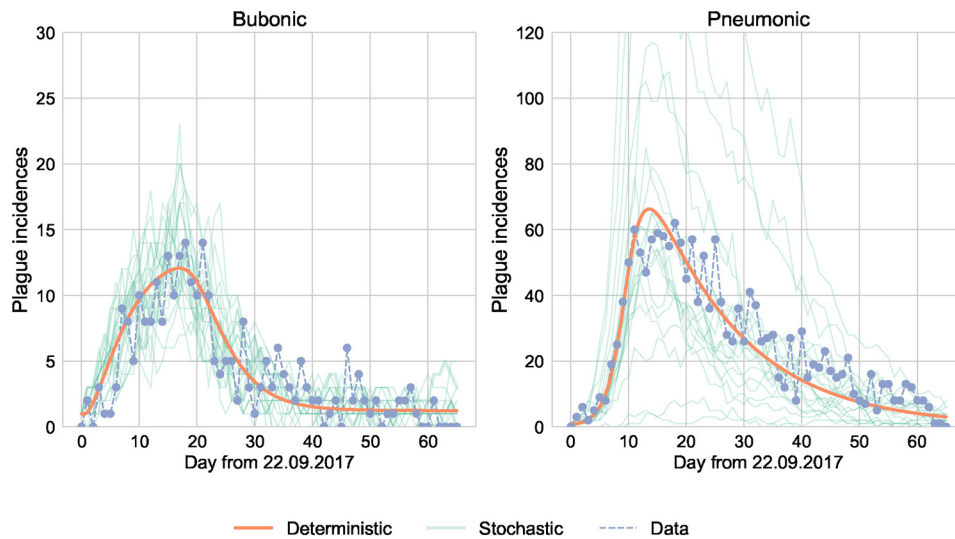


Fig. 5. Deterministic and stochastic simulations of plague epidemics. The parameters were estimated with the global optimisation algorithm Differential Evolution. Stochastic simulations were done with tau-leaping algorithm.

Evaluating the Jacobian of the PTM at the disease-free equilibrium yielded a threshold parameter R^*

$$R^* = \frac{\beta f_{itv,h}(\bar{s}_b + \bar{s}_p)}{\delta_p} \tag{7}$$

such that $R^* \geq 1$ results in $\lambda_6 \geq 0$, where λ_6 is the largest eigenvalue of the Jacobian, all the others being non-positive (Appendix A). This result agrees with the estimate obtained from the next-generation-matrix (NGM) (Diekmann et al., 2009), for which we find the leading eigenvalue at $t = 0$ to be $R_0 = \beta/\delta_p = R^*(0)$. In other words, at the beginning ($S_b + S_p = N$) when there are no interventions ($f_{itv,h} = 1$), an infected subject has a transmission rate β spent an average $1/\delta_p$ days and thus generate $\beta \times \frac{1}{\delta_p} = 6.54$ secondary cases.

Table 1
Transmission parameters estimated from PTM^a and statistical models.

Estimate	Range	Meaning
p	1×10^{-4} 0–0.5	fraction exposed to infected rat flea
γ_b	0.23 2–6 days (Centers for Disease Control and Prevention, 2005)	inverse bubonic plague incubation period
γ_p	0.29 1–6 (Kool and Weinstein, 2005; Mead, 2017), 1–12 (Raymond Gani, 2004) days	inverse pneumonic plague incubation period
δ_b	0.26 4–10 days (Nikiforov et al., 2016)	inverse bubonic plague infection period
δ_p	0.34 1–12 (Raymond Gani, 2004), 1–15 (Tieh and Landauer, 1948) days	inverse pneumonic plague infectious period
ϵ	0.03 < 5% (Kool and Weinstein, 2005), 8–10% (Nikiforov et al., 2016)	rate of transition to secondary pneumonic
α	1.9×10^{-3} 0–10	(scaled) flea-to-human transmission rate
β	2.23 0–10	human-to-human transmission rate
θ	0.11 ≥ 0	reduction in intervention effects
τ_h	8.89 0–43 ^b	time at which interventions reached half its maximum effect on human-to-human
τ_f	17.93 0–43	— in flea-to-human

R0	Estimate ^c	Methods
	6.54	Next generation matrix (Diekmann et al., 2009) ($R_0 = \beta/\delta_p$)
	6.9[5.2–9.3]	Exponential Growth (Wallinga, 2007)
	6.9[5.7–8.3]	Maximum likelihood (Forsberg White and Pagano, 2008)
	6.9[0.8–11.7]	Sequential Bayes estimation (Bettencourt and Ribeiro, 2008) at $t = 0$
	6.9[5.2–9.7]	Binomial assumption (Nishiura, 2010)
	7.1, 5.1	Sub-exponential growth (Viboud et al., 2016; Chowell et al., 2016) at 1 week, at epidemic peak.

^a Differential Evolution algorithm was run minimizing the least absolute differences of both the bubonic and pneumonic data weighted by their range. The total population used for simulations is $N = 25, 570, 895$ (United Nations, 2017). Initial conditions of the model is $\{Np, N(1 - p), 0, 0, 1, 1\}$. For parameters have been previously estimated elsewhere with a large variance, their range was used here as boundary constrains for the optimisation.

^b Maximum simulation time.

^c Number in brackets are 95% confident interval.

However, while it has been noted for the last Ebola outbreak (Althaus, 2014), data sharing practices are still poor and WHO practices of reporting (separate PDF files) are putting constraints on modelling works. In the 2014 Ebola outbreak, for example, most of the modelling studies were done rather late in the process (Chretien et al., 2015). The solution can be as simple as putting a unique Excel file and update it, or better establishing a central website for all WHO outbreak reports, or alternatively, a data hub to encourage user-contributed reports (a demo of such interface can be viewed as <https://epidemic.Live>).

From modelling aspects, plague outbreaks can be more challenging because: (1) there is a continuous input of flea-to-human transmission (Fig. 2); this implies the observed epidemic curve can be a mixture of multiple waves of infected cases generated from different index cases. Thus, epidemic evaluations could risk over- or under-estimating the consequences, e.g. the reproduction number or the end time of epidemics; (2) there is a known seasonal pattern of the plague epidemic (Kreppel et al., 2014) for which a direct measure of the rat flea population does not exist; (3) the flea-to-human transmission as well as the transition from bubonic cases to pneumonic cases seemed stochastically driven (Fig. 2) and could be highly affected by interventions.

Here, the epidemic curves showed plague incidences appearing sporadically during August (Fig. 2). But then a large increase in pneumonic cases was observed, preceded by only a few bubonic cases. Estimates of the plague reproduction number showed that a high estimate is needed to capture this fast-growing phase of the epidemic; the estimate doubled the previous estimates ranging 2.8–3.5 (Nishiura et al., 2006). It can be speculated that superspreaders were likely to exist in order to generate the large number of cases in a short time as observed in the pneumonic epidemic curve. This was a common pattern as described in plague outbreaks in Madagascar 2017 (World Health Organization, 2017c) and in the US 1919, 1924, and 1980 (Kool and Weinstein, 2005). Considering the potential mix of epidemic waves and outbreak locations in the used data, our estimate of R_0 could be over-estimated. However, the smaller estimates of R_0 for this outbreak ranging 1.1–1.4 (Tsuzuki et al., 2017; Nishiura et al., 2012) could be due to the inclusion of the long sporadic incidences occurred during August which can misjudge the transmission ability of pneumonic plague in optimal conditions from 22.09.2017 onwards (Fig. 2).

Nonetheless, approaches adjusting for the propagated outbreak data are needed to further understand its effect on parameter estimates. For example, how do the data of a non-synchronised and combined epidemic curve affect the reproduction number? Implementations and calculations using current methods raised also an issue that has been discussed elsewhere (Forsberg White and Pagano, 2008), that is estimating the reproduction number using serial interval could yield very high value during the first days of the epidemic growing phase as the denominator of the estimator is extremely small during this period. Overestimate of R_0 values have also been observed in populations with heterogeneous contact pattern population (Llensa et al., 2014).

Appendix A. Supporting information

The eigenvalues of the Jacobian of the PTM evaluated at the disease-free state $(S_b, S_p, E_b, E_p, I_b, I_p)/N = (\bar{s}_b, \bar{s}_p, 0, 0, 0, 0)$ have the form

$$\lambda_1 = 0 \quad (8)$$

$$\lambda_2 = -\alpha f_{ivf} f_{ivf} \quad (9)$$

$$\lambda_3 = -\gamma_b \quad (10)$$

$$\lambda_4 = -(\delta_b + \varepsilon) \quad (11)$$

$$\lambda_5 = -\frac{1}{2} \left(\gamma_p + \delta_p \right) \left[\left(1 + \frac{4\gamma_p [\beta f_{iv,h} (\bar{s}_b + \bar{s}_p) - \delta_p]}{(\gamma_p + \delta_p)^2} \right)^{1/2} + 1 \right] \quad (12)$$

It seemed that sporadic inputs of bubonic cases brought new index cases to the human-to-human transmission network constantly with an estimate of transition to secondary pneumonic is of 3% (Fig. 2). It followed that the pneumonic epidemic curve exhibited multiple peaks and waves. As the transmission rate can be high, this observation suggests that vector control measures are central to prevent potential next waves of the epidemic. This might be practical as the fraction of the population exposed to infected rat fleas was low (Table 1). On the other hand, the estimate of human-to-human transmission needs to be large in order to capture the data. This prompts that intervention efforts, in cases of exhaustible resources, should prioritise stopping human-to-human transmission route. In this case, while the bubonic cases would continue to appear they would not be able to generate large size outbreaks. This approach is practical as with the usual public health intervention of early detection of the incidences would not only stop the human-to-human transmission but also provide a better chance for new bubonic cases to be treated early.

Experiments have shown that an optimal climate for rat fleas to flourish is a dry climate with temperatures of 20–25 °C (Shrewsbury, 2005). These conditions were observed during the outbreak: a generally dry weather with the optimal temperature coincided with the period of high epidemic activity. As further shown in Fig. 4, these conditions would typically remain the same until May 2018, see further in Kreppel et al. (2014). This again stresses the role of vector control in preventing the next waves. It is worth noting that the global changes in climate conditions could pose a risk of irregular changes in vector biting rate and reproduction (World Health Organization, in press), requiring affected regions to be vigilant in plague control and prevention. For the modelling aspect, studies on the dynamics of rat fleas are needed to help further parameterise the plague transmission model.

In this paper, we collected and described relevant data of the 2017 plague epidemic in Madagascar. We proposed a working mathematical model for evaluating and predicting epidemic consequences and what-if scenarios. We discussed potential drawbacks in modelling propagated epidemic data. We hope that the results would contribute informative insights for public health officers and provide a framework for further understanding the dynamics of plague outbreaks.

Author's contributions

VKN, EAH-V: conceptualization and data curation. VKN: methodology, analysis, investigation, and visualisation. CP-R: analysis, stochastic simulations implementation, and visualisation. EAH-V: funding acquisition, supervision. All authors: review & editing.

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$$\lambda_6 = \frac{1}{2} \left(\gamma_p + \delta_p \right) \left[\left(1 + \frac{4\gamma_p [\beta_{inv,h}^c (\bar{s}_b + \bar{s}_p) - \delta_p]}{(\gamma_p + \delta_p)^2} \right)^{1/2} - 1 \right] \quad (13)$$

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