

Reassessing global historical \mathcal{R}_0 estimates of canine rabies

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Abstract

Rabies spread by domestic dogs continues to cause tens of thousands of human deaths every year in low- and middle-income countries. Despite this heavy mortality burden, rabies is often neglected, perhaps because it has been eliminated from high-income countries through mass dog vaccination. [MR: Maybe discuss neglect elsewhere; there are issues.](#) Estimates of the intrinsic reproductive number (\mathcal{R}_0) of canine rabies from a wide range of times and locations are low (values <2), with narrow confidence intervals. This narrow range of estimates across a wide range of dog densities and environments is surprising. [MR: Can we assume readers of this journal know about \$\mathcal{R}_0\$?](#) We combined incidence data from historical outbreaks of canine rabies from around the world (1917-2003) with high-quality contact-tracing data from Tanzania (2002-present) to investigate initial growth rates (r_0), generation intervals distributions (G) and reproductive numbers (\mathcal{R}_0). We updated earlier work by: choosing outbreak windows algorithmically; fitting r_0 using a statistical method that accounts for decreases through time; and propagating uncertainty from both r_0 and G when

estimating \mathcal{R}_0 . Our \mathcal{R}_0 estimates are larger than previous estimates, with wider confidence intervals. Our novel hybrid approach for estimating \mathcal{R}_0 and its uncertainty is applicable to other disease systems where researchers estimate \mathcal{R}_0 by combining estimates of r_0 and G .

Introduction

Canine rabies, primarily spread by domestic dogs, is a vaccine-preventable disease that continues to cause tens of thousands of human deaths every year in low- and middle-income countries (LMICs) (Taylor et al., 2017; Minghui et al., 2018). Canine rabies has been effectively eliminated from high-income countries by mass dog vaccination (Rupprecht et al., 2008). Despite the effectiveness of vaccinating dogs, rabies continues to cause many human deaths and large economic losses in LMICs due to the limited implementation of rabies control strategies (Hampson et al., 2015). The past two decades have seen an increase in rabies control efforts — including dog vaccination campaigns and improvements in surveillance (Kwoba et al., 2019; Mtema et al., 2016; Gibson et al., 2018; Mazeri et al., 2018; Wallace et al., 2015). [MR: Discuss COVID effects briefly?](#) More recently, the World Health Organization (WHO) and partners (OIE, FAO, GARC) joined forces to support LMICs in eliminating human deaths from dog-mediated rabies by 2030 (Minghui et al., 2018; Abela-Ridder et al., 2016). Mass dog vaccination campaigns have begun in some LMICs and are being scaled up (Castillo-Neyra et al., 2019; Evans et al., 2019). An understanding of rabies epidemiology — in particular, reliable estimates of the basic reproductive number (\mathcal{R}_0), a quantitative measure of disease spread that is often used to guide vaccination strategies — could inform rabies control efforts.

\mathcal{R}_0 is defined as the expected number of secondary cases generated from each primary case in a fully susceptible population (Macdonald, 1952). Estimates of \mathcal{R}_0

using various methods (i.e., direct estimates from infection histories, epidemic tree reconstruction, and epidemic curve methods) based on historical outbreaks of rabies have generally been surprisingly low, typically between 1 and 2 with narrow confidence intervals (Hampson et al., 2009; Kurosawa et al., 2017; Kitala et al., 2002). It is surprising that a wide variety of regions and time periods with different densities and population structures of domestic dogs would have such similar values of \mathcal{R}_0 . MR: I think the idea that different densities and population structures imply different R_0 might need to be articulated more explicitly here. Limit to single use of "surprise" in the sentence that does a better job setting up the expectation. In contrast to diseases with large \mathcal{R}_0 (e.g. measles with $\mathcal{R}_0 > 10$ (Guerra et al., 2017)), \mathcal{R}_0 estimates for rabies imply that control through vaccination should be relatively easy (compared to e.g., rinderpest with $\mathcal{R}_0 \approx 4$ (Mariner et al., 2005)). MR: @JD suggested use only rinderpest (omit e.g. measles) With such a low \mathcal{R}_0 one might expect rabies to fade out from behavioural control measures combined with stochastic fluctuations, even in the absence of vaccination. MR: move to before statements of surprise, as it sets the expectation

Here we revisit and explore why rabies, with apparently low \mathcal{R}_0 , nonetheless persists in many countries around the world. Such persistence suggests that rabies's potential for spread, and therefore the difficulty of rabies control, may have been underestimated. MR: Maybe this is the intent, but I experience cognitive dissonance reading these two sentences... I think "apparent" is potentially confusing because even though $R_0 \approx 1$ is what has been estimated, it is apparently not right? In this paper, we will combine information derived from epidemic curves with a high-resolution contact tracing dataset that provides large number of observed generation intervals (which is rare for infectious disease studies) to estimate \mathcal{R}_0 . This reassessment can improve the estimation of \mathcal{R}_0 and understanding of disease control more generally. MR: I think the "more generally" part deserves a bit more preamble/evidence...

maybe this is a bit late to get to the general overall, although this may depend where you're publishing and the hopes that this can spur rabies vaccination campaigns

Materials and Methods

\mathcal{R}_0 is often estimated by combining two other epidemiological quantities: the initial growth rate of an epidemic (r_0) and the generation interval (G) distribution, where a G is defined as the time between successive infections along a transmission chain. [MR: does statement of "often" demand a citation?](#) The growth rate r_0 is often estimated by fitting a growth rate to time series data from the early stages of epidemics. G is an individual-level quantity that measures the time between an individual getting infected to infecting another individual. The generation interval distribution is the natural way to link r_0 and \mathcal{R}_0 (Wallinga and Lipsitch, 2006; Champredon and Dushoff, 2015). During an outbreak in a fully susceptible population, \mathcal{R}_0 can be calculated from r_0 and the G distribution by the Euler-Lotka equation (Wallinga and Lipsitch, 2006)

$$\mathcal{R}_0 = \frac{1}{\sum_{t=1}^{\infty} G(t)e^{-rt}}, \quad (1)$$

where t is time, and $G(t)$ is the generation interval distribution. This formula is convenient to calculate point estimates of \mathcal{R}_0 ; however, propagating uncertainty from the estimates of r_0 and the G distribution can be difficult. [MR: Replace "can be difficult" with something like "hasn't been done often" or "isn't part of the standard epi protocol" or something more empirical?](#)

Initial growth rate

Disease incidence typically increases approximately exponentially during the early stages of an epidemic. The initial growth rate r_0 is often estimated by fitting exponential curves from near the beginning to near the peak of an epidemic. However,

growth rates estimated from an exponential model can be biased downward, overconfident, and sensitive to the choice of fitting windows (Ma et al., 2014). Here we used the logistic curves provide more robust estimates of r_0 (Ma et al., 2014; Chowell, 2017). MR: This seems like such a big piece of the difference between KH and WZLi estimates; I'm surprised it only gets mentioned here in M&M but not highlighted in abstract/methods. Especially since KH is an author, seems fine to be more firm in claiming logistic is more appropriate and likely to resolve some of the "paradox"

MR: do you need to set up the need to break longer TSs into windows more generally? Or is this so basic it doesn't require any verbiage? We select our fitting window consistently for each outbreak as follows: ML: Feed me with the new method of window selection.

Observed Generation intervals

Transmission events are generally hard to observe for most diseases. In an earlier, influential paper, MR: is "influential" helpful here? Useful to narrow the scope to rabies? estimated generation intervals were constructed by summing two quantities: a latent period (the time from infection to infectiousness), and a wait time (time from infectiousness to transmission) (Hampson et al., 2009). Since clinical signs and infectiousness appear at nearly the same time in rabies, the incubation period (the time from infection to clinical signs) is routinely used as a proxy for the latent period. In the Hampson et al. analysis, latent (really, incubation) periods and infectious periods were randomly and independently resampled from empirically observed distributions (Hampson et al., 2009), and then waiting times sampled uniformly from the selected infection periods.

However, this approach for constructing G values (i.e., summing independently resampled values of incubation and infectious periods) does not account for the possibility of multiple transmissions from the same individual, nor does it account for

correlations between time distributions and biting behaviour. Figure 1 illustrates the generation intervals of a single transmission event from a rabid animal (comprising a single incubation period plus a waiting time) and multiple transmission events from a rabid animal (comprising a single incubation period and three waiting times). For diseases like rabies, where transmissions links (and generation intervals) are observable, multiple transmissions and possible correlation structures are all accounted for within the observation processes [\[\[BB: unclear\]\]](#).

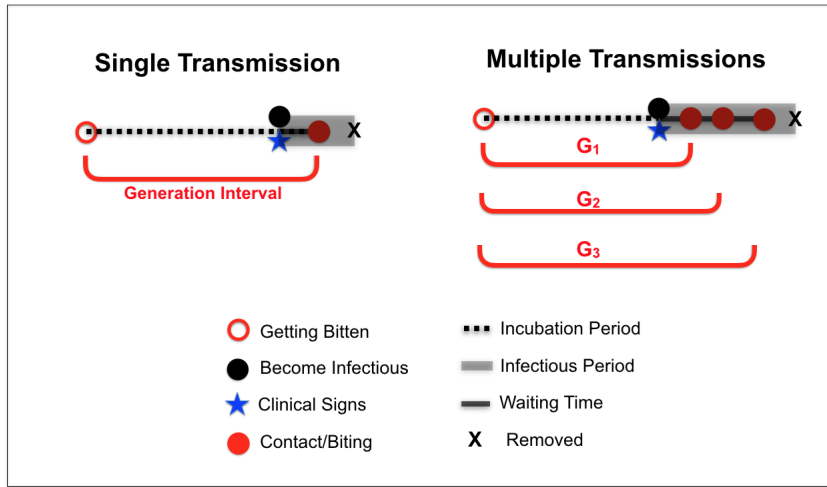


Figure 1: **Decomposing generation intervals.** Generation intervals start when a focal animal acquires infection (open red circle) and end after a period of viral replication (dashed line) when an animal shows clinical signs (blue star), becomes infectious (solid black circle) and infects another animal — in rabies the onset of clinical signs and of becoming infectious are closely synchronized. Once the infectious period (grey block) starts, there is a wait time (solid black line) until a susceptible host (solid red circles) is bitten. The infectious period ends with the death of the focal host (black X). The generation interval is the interval between getting infected and infecting a new case (red interval between open and solid circles). (right) If a single biter transmits multiple times, the wait times are generally different, but the incubation period is the same for each transmission event.

Correcting for vaccination

In a population where some animals are not susceptible, calculations based on estimates of r_0 and the G distribution (1) estimate the *realized* average number of cases per case, also known as the effective reproductive number \mathcal{R}_e . In the case of rabies, vaccination is the only known cause of immunity (case fatality in dogs is believed to be 100%). For a given population with ν vaccination proportion, the estimated \mathcal{R}_0 with vaccination correction is the following:

$$\mathcal{R}_0 = \frac{\mathcal{R}_e}{(1 - \nu)}. \quad (2)$$

Data and material

We used data from December 2002 – November 2022, from an ongoing contact tracing project in Tanzania (Hampson et al., 2008, 2009). Since 2002, there are 8636 domestic dog recorded events (i.e., domestic dogs bitten by an animal), and 3552 suspected rabid dogs in the Serengeti, Tanzania. Transmission events were documented through retrospective interviews with witnesses, applying diagnostic epidemiological and clinical criteria from the six-step method (Tepsumethanon et al., 2005). Each dog was given a unique identifier and date of the bite and clinical signs were recorded if applicable and available. 2132 of dog transmissions were from unidentified domestic animals or wildlife. We restricted our analysis in this paper to domestic dog transmissions (i.e., dog to dog), and obtained 293 directly observed generation intervals (i.e. both biter and secondary case have time bitten recorded). There were four observed dogs with multiple exposures (i.e., bitten by different identified biters), generating extra generation intervals, but it is unclear which transmission event transmitted rabies to these dogs. For simplicity, we omitted these four dogs and their generation intervals from our analysis.

Fitting and Propagating Parameter Uncertainties

To propagate uncertainties for both r_0 and G , we used a hybrid approach. We first fitted logistic models, with negative binomial observation error, to incidence data to estimate r_0 implemented in the R package “epigrowthfit”. **ML: cite** We then compute a sample of 300 $\hat{\mathcal{R}}_0$ values using equation (2); for each value of $\hat{\mathcal{R}}_0$, we first draw a value of \hat{r}_0 from a normal distribution from the estimates of the logistic fit and 100 G from the empirical contact tracing data. **ML: fix me, use R to fill in the sample numbers.** We then matched bootstrap samples of G to the r_0 samples to produce a range of estimates for \mathcal{R}_0 . This hybrid bootstrap approach incorporates both sources of uncertainties from r_0 and G (frequentist empirical bootstrap) **MR: this parenthetical seems strange - (why isnt there one for little r as well?)** when calculating \mathcal{R}_0 estimates. Finally, we take the 25, 50, 97.5% percentiles of the distribution of \mathcal{R}_0 estimates for each rabies outbreak **MR: I want 25th, 50th, 97.5th. Also, do you want to justify these three quantiles?.**

Results

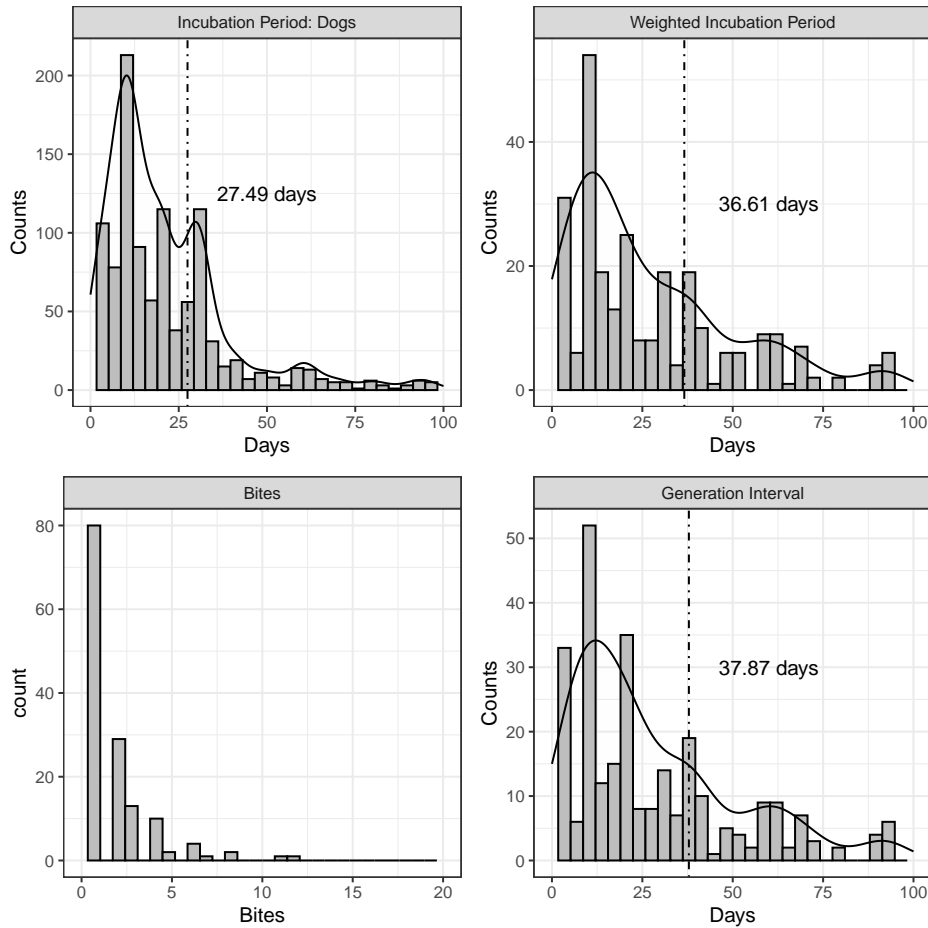


Figure 2: **Empirical distributions from contact tracing data.** MR: I like it when the first sentence in the caption states the figure's take-home point Top left is the distribution of observed incubation periods. Top right is the distribution of infectious periods weighted by each dog's biting frequency (biting frequency shown bottom left). The weighted distribution corresponds to the contribution of incubation periods to generation intervals, which are shown bottom right. Dash-dotted lines show the means of each time-interval distribution. MR: maybe add letters for each panel?, Oh, Ben said the same thing :P [[BB: one less sig digit in mean values? Lower left ("bites" panel) seems like maybe it should be of the same style as type = h for base plots (i.e. geom_segment from 0 to number of counts, with a point at the top?) Why is the top right facet label "incubation period: dogs" ? Can we have letter labels on the subplots so we can refer to a, b, c, d, rather than top left/top right/etc.??]

Figure 2 shows the empirical distributions of the observed incubation periods, rabid dog biting frequency, and generation intervals from contact tracing data. The mean observed incubation period is 27.5 days ($n = 1109$ dogs), the mean biting frequency is 1.65 bites per rabid dog, and the weighted mean incubation period is 36.6

days ($n = 143$ biting dogs). The mean observed generation interval is 37.9 days ($n = 143$ primary infections resulting in 293 secondary cases), which is significantly larger than the mean generation interval constructed from independently summing incubation periods and wait times (24.9 days (Hampson et al., 2009)). The weighted incubation period distribution more closely resembles the generation interval distribution than the incubation period of all dogs.

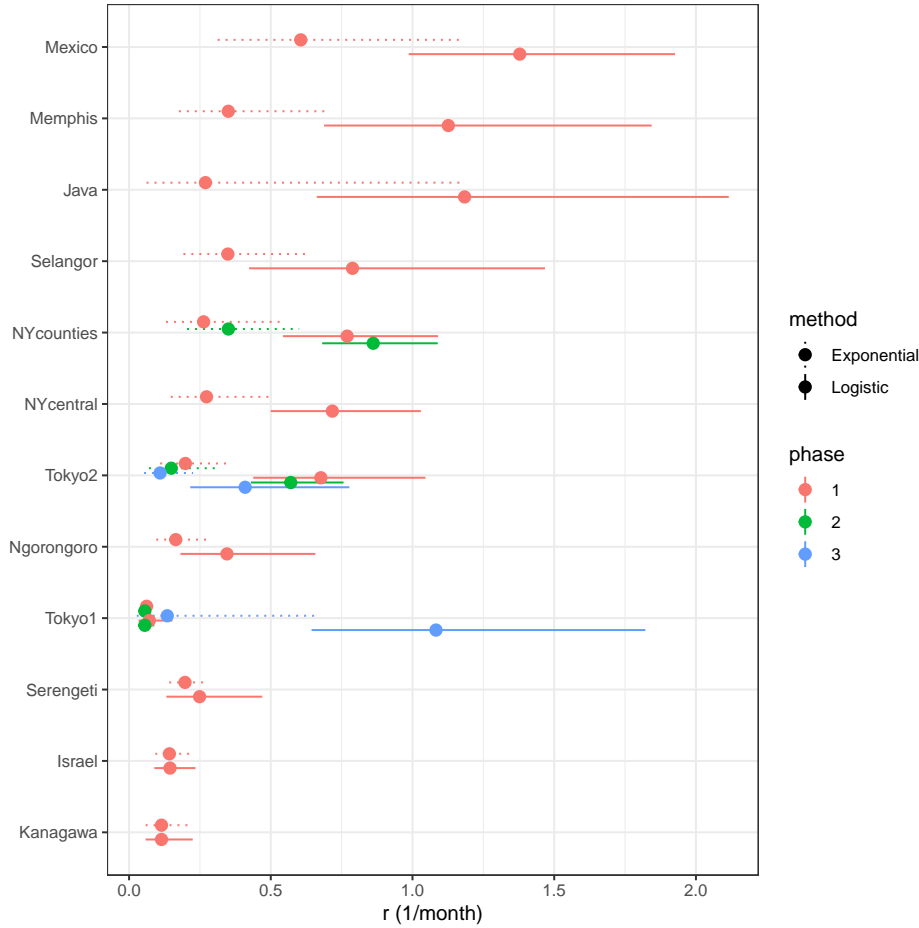


Figure 3: **Growth rate estimates for global historical outbreaks of rabies.** Estimates and 95% confidence intervals of r_0 in global historical outbreaks estimated from exponential (dotted) and logistic (solid) model fits. Different colors represents different phases from the times series data.

We estimated r_0 from historical outbreak data (Figure 3). For a direct compari-

son, we also estimated r_0 from an exponential model. MR: I think it would be helpful to explain more what the comparisons are you're hoping the reader will make' Overall, r_0 estimates from the logistical model are larger with wider confidence intervals compared to r_0 estimates from the exponential model.

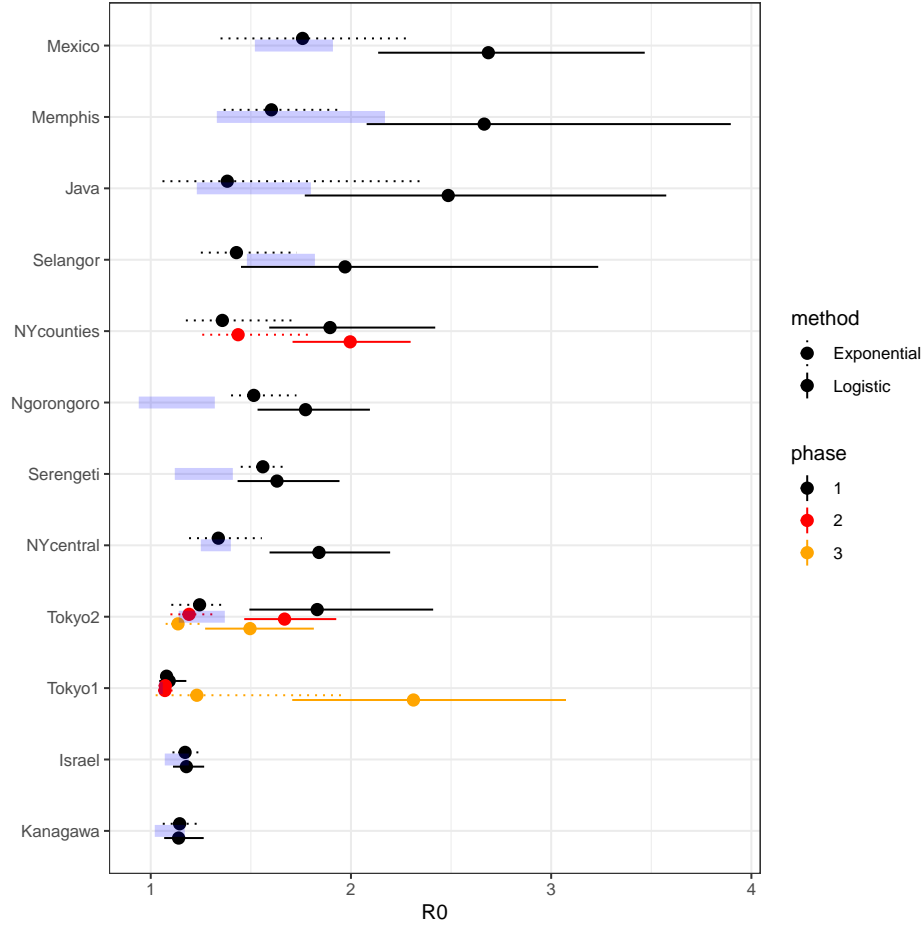


Figure 4: **Reproductive number estimates for global historical outbreaks of rabies**

Previous estimates of \mathcal{R}_0 are shown in blue highlights; \mathcal{R}_0 estimates and confidence interval (95% quantiles from the estimated \mathcal{R}_0 sample) from our hybrid approach using exponential (dotted lines) and logistic (solid lines). \mathcal{R}_0 are corrected for vaccination coverage. ML: todo: add vaccination coverage like KH's paper in the location name.

We combined our estimates of r_0 from the logistic model with the empirical G

from our detailed Tanzanian data to produce \mathcal{R}_0 estimates. Of the listed historical outbreaks, four occurred in locations with prior rabies vaccination coverage: Memphis, US (1947, 10% vaccine coverage); Serengeti, Tanzania (2003, 20% coverage); Ngorongoro, Tanzania (2003, 20% coverage); and Sultan Hamad, Kenya (1992, 24% coverage). Our estimates of \mathcal{R}_0 ranged from 1.08 to 2.66 **ML: check and use program**, with upper confidence intervals greater than 2 for most locations. The hybrid approach provides larger values of \mathcal{R}_0 and wider confidence intervals than previous \mathcal{R}_0 estimates after propagating uncertainty from both r_0 and generation interval distributions (see Figure ??).

MR: you set up the idea that R_0 should vary across populations, but I don't think very clearly yet (density seems like a good one, since the idea behind R_0 is that it includes both the contact rate and the transmission rate). But you use a single region to estimate generation intervals... can you state a bit about why this seems like a fine assumption (i.e., something like you expect similar variation in dogs within and across populations – even though the other paper is all about how important the dog-to-dog variation in latent period is for understanding dynamics?) Is this line of thinking citeable?

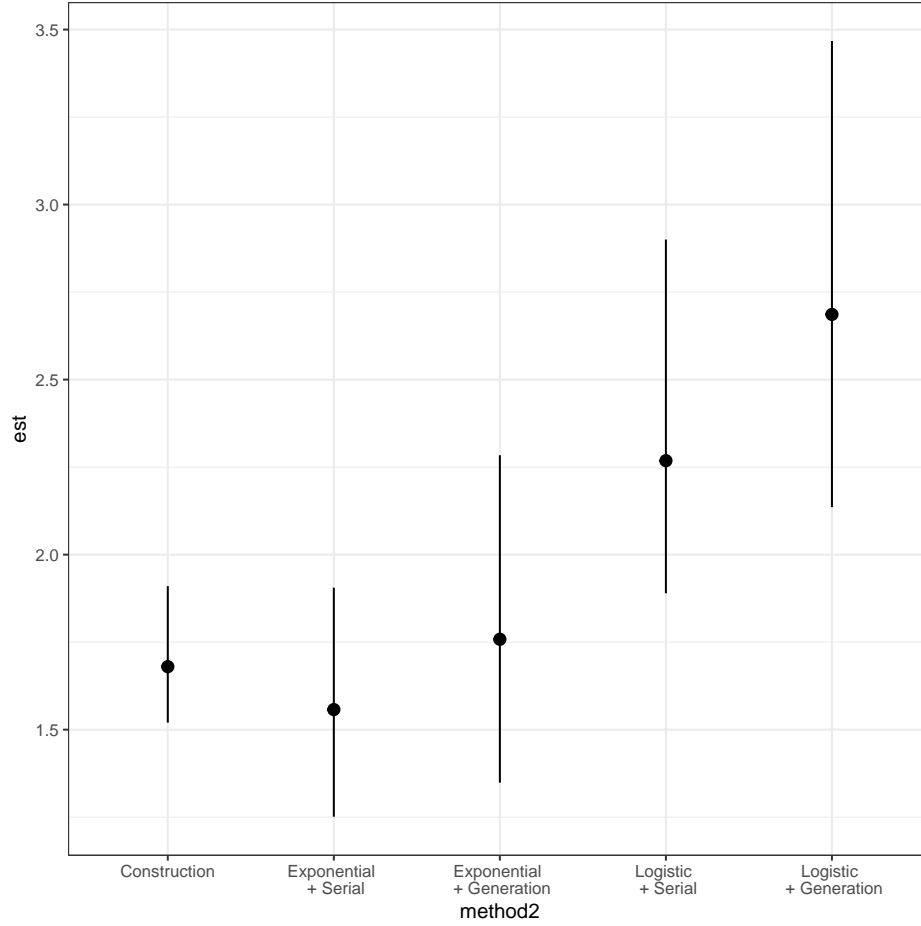


Figure 5: **Effects of r_0 , corrected G on the estimates of \mathcal{R}_0 in Mexico outbreak.** [[BB: more complete caption? Make this horizontal to match the other figs? Extend R0-axis to have a lower limit at $\mathcal{R}_0 = 1$?]] MR: what ben said :-)

Lastly, we compare the effects of different estimation techniques of r_0 and G on estimates of \mathcal{R}_0 (Figure 5). For illustrative purposes, we used the 1987 outbreak in Mexico where there was no vaccination; the supplementary material shows similar plots for the other historic outbreaks. Propagating uncertainty from both r_0 and G generally leads to wider confidence intervals compared to previous \mathcal{R}_0 estimates in Hampson et al. (2015). \mathcal{R}_0 estimate increases when we estimate r_0 via the logistic model, or G instead of the constructed approach in ? . The overall effect of \mathcal{R}_0 increases even more when included both corrections (i.e., in r_0 and G) with even larger uncertainty.

Discussion

MR: I like it when the first sentence of the discussion states the most important result. Here, I think you could lead with a condensed and refined version of "We resolved the mystery of rabies persistence despite the fact that R_0 estimates in historical outbreaks have been consistently low, showing that substantially higher basic reproduction numbers are compatible with historical outbreak data. Here, we reanalyzed historical rabies epidemics with improved model assumptions and uncertainty propagation, showing that historical estimates of R_0 were downward biased and overconfident". I think the place to funnel back out the broader implications is a bit later, so you don't need to start with " R_0 is commonly used..."

The basic reproductive number \mathcal{R}_0 is commonly used to summarize the risk of infectious disease and to inform control measures. Here, we used a relatively simple approach to estimate \mathcal{R}_0 by combining initial growth rate (r_0) estimates from incidence data and generation intervals from contact tracing data. We improved on earlier work by correcting for slowdown in growth when estimating r_0 , and by developing an approach to propagate uncertainty from both r_0 and G , resulting in higher \mathcal{R}_0 estimates with wider confidence intervals.

Re-analysis of these data also allowed us to identify an overlooked fact about rabies generation intervals: observed generation intervals are longer on average than constructed ones [\[\[BB: than intervals constructed by naively adding ...?\]\]](#), because of within-individual correlations in time distributions and biting behaviour. The unexpected importance of these correlations could have implications for G -based studies of other infectious diseases. [MR: sharpen this sentence](#) Further investigation of how these correlations affect the overall dynamics of rabies is warranted. [MR: this paragraph seems out of place; the next paragraph follows the previous one in discussing logistic vs. exponential growth. I kind of think more of this belongs in the intro; otherwise it's not as clear why you make the choice to use the logistic/trust those](#)

estimates more'

Estimates of \mathcal{R}_0 are strongly affected by estimates of the growth rate during the initial phase of the epidemic. The logistic model gives a better approximation of the initial phase of the epidemic resulting in a larger estimate of r_0 compared to the exponential model (Ma et al., 2014). Our estimates of r_0 account for observation error (measurements may not perfectly match reality), but not for process error (the fundamental stochasticity of the system itself). Thus, there may be more uncertainty in r_0 than we estimate (King et al., 2015), but this is not always true in practice (Li et al., 2018). [\[\[BB: don't really understand the second half of this sentence?\]\]](#)

Nevertheless, our estimates suggest that rabies \mathcal{R}_0 may be larger, and more uncertain, than previously thought. This finding may explain some of the formerly unexplained variations in the success of rabies-control programs (e.g., low levels of coverage (30–50%) have been successful in some settings while high coverage 75% was not enough to control rabies in others (Eng et al., 1993)). While our primary goal was to understand why estimates of rabies \mathcal{R}_0 were small with narrow confidence intervals, our analysis also revealed an interesting biological process in rabies generation interval. [\[\[BB: which is ... ?\]\]](#) \mathcal{R}_0 is typically used as a first approximation for interventions such as vaccination to determine herd immunity thresholds. However, the main heterogeneity in contacts and correlations with incubation periods observed through the generation interval may suggest that simple \mathcal{R}_0 estimation methods may be inadequate and should be used with caution. [\[\[BB: refine this sentence?\]\]](#) A mechanistic fitting framework will be useful to study these patterns in more detail.

[MR: not sure which paper is destined to come out first, or if you're subitting them \(i.e., this one an correlations paper\) together somehow... I think there could be a more explicit linking between the two papers. I think the sentence Ben suggested you refine could be more positive: you advocate estimating \$R_0\$ with uncertainty propagation \(and also why logistic is more appropriate than exponential, and how](#)

general you think that should be. I don't love landing on the mechanistic fitting suggestion; think it would be good to end on strong sentences focused on take-home messages.) Consider moving most of last paragraph up to first in discussion. I think more can be said about how Rabies is a nice model system because latent period and transmission events are so easy to observe, and natural immunity is not thought to exist, but your findings can be applicable in disease systems where the uncertainties in generation interval may be even greater, or something like that.

[[BB: really, really unimportant, but: check for appropriate capitalization of PLOS? (PLoS?)]]

References

- Abela-Ridder, B., L. Knopf, S. Martin, L. Taylor, G. Torres, and K. De Balogh (2016). 2016: the beginning of the end of rabies? *The Lancet Global Health* 4(11), e780–e781.
- Castillo-Neyra, R., A. M. Toledo, C. Arevalo-Nieto, H. MacDonald, M. De la Puente-León, C. Naquira-Velarde, V. A. Paz-Soldan, A. M. Bittenheim, and M. Z. Levy (2019, August). Socio-spatial heterogeneity in participation in mass dog rabies vaccination campaigns, Arequipa, Peru. *PLOS Neglected Tropical Diseases* 13(8), e0007600.
- Champredon, D. and J. Dushoff (2015). Intrinsic and realized generation intervals in infectious-disease transmission. *Proceedings of the Royal Society B: Biological Sciences* 282(1821), 20152026.
- Chowell, G. (2017). Fitting dynamic models to epidemic outbreaks with quantified uncertainty: A primer for parameter uncertainty, identifiability, and forecasts. *Infectious Disease Modelling* 2(3), 379–398.
- Eng, T., D. Fishbein, H. Talamante, D. Hall, G. Chavez, J. Dobbins, F. Muro, J. Bustos, M. De Los Angeles Ricardy, A. Munguia, et al. (1993). Urban epizootic of rabies in Mexico: epidemiology and impact of animal bite injuries. *Bulletin of the World Health Organization* 71(5), 615.
- Evans, M., J. B. Bailey, F. Lohr, W. Opira, M. Migadde, A. Gibson, I. Handel, M. Bronsvoort, R. Mellanby, L. Gamble, et al. (2019). Implementation of high coverage mass rabies vaccination in rural Uganda using predominantly static point methodology. *The Veterinary Journal* 249, 60–66.

- Gibson, A. D., S. Mazeri, F. Lohr, D. Mayer, J. L. B. Bailey, R. M. Wallace, I. G. Handel, K. Shervell, M. Barend, R. J. Mellanby, et al. (2018). One million dog vaccinations recorded on mHealth innovation used to direct teams in numerous rabies control campaigns. *PloS one* 13(7), e0200942.
- Guerra, F. M., S. Bolotin, G. Lim, J. Heffernan, S. L. Deeks, Y. Li, and N. S. Crowcroft (2017). The basic reproduction number (R_0) of measles: a systematic review. *The Lancet Infectious Diseases* 17(12), e420–e428.
- Hampson, K., L. Coudeville, T. Lembo, M. Sambo, A. Kieffer, M. Attlan, J. Barrat, J. D. Blanton, D. J. Briggs, S. Cleaveland, et al. (2015). Estimating the global burden of endemic canine rabies. *PLoS Neglected Tropical Diseases* 9(4), e0003709.
- Hampson, K., A. Dobson, M. Kaare, J. Dushoff, M. Magoto, E. Sindoya, and S. Cleaveland (2008). Rabies exposures, post-exposure prophylaxis and deaths in a region of endemic canine rabies. *PLoS Neglected Tropical Diseases* 2(11), e339.
- Hampson, K., J. Dushoff, S. Cleaveland, D. T. Haydon, M. Kaare, C. Packer, and A. Dobson (2009). Transmission dynamics and prospects for the elimination of canine rabies. *PLoS Biology* 7(3), e1000053.
- King, A. A., M. Domenech de Cellès, F. M. Magpantay, and P. Rohani (2015). Avoidable errors in the modelling of outbreaks of emerging pathogens, with special reference to Ebola. *Proceedings of the Royal Society B: Biological Sciences* 282(1806), 20150347.
- Kitala, P., J. J. McDermott, P. Coleman, and C. Dye (2002). Comparison of vaccination strategies for the control of dog rabies in Machakos District, Kenya. *Epidemiology & Infection* 129(1), 215–222.
- Kurosawa, A., K. Tojinbara, H. Kadowaki, K. Hampson, A. Yamada, and K. Makita (2017). The rise and fall of rabies in Japan: A quantitative history of rabies epidemics in Osaka prefecture, 1914–1933. *PLoS Neglected Tropical Diseases* 11(3), e0005435.
- Kwoba, E. N., P. Kitala, L. Ochieng, E. Otiang, R. Ndung’u, G. Wambura, K. Hampson, and S. Thumbi (2019). Dog health and demographic surveillance survey in Western Kenya: Demography and management practices relevant for rabies transmission and control. *AAS Open Research* 2.
- Li, M., J. Dushoff, and B. M. Bolker (2018). Fitting mechanistic epidemic models to data: A comparison of simple Markov chain Monte Carlo approaches. *Statistical methods in medical research* 27(7), 1956–1967.
- Ma, J., J. Dushoff, B. M. Bolker, and D. J. Earn (2014). Estimating initial epidemic growth rates. *Bulletin of mathematical biology* 76(1), 245–260.
- Macdonald, G. (1952). The analysis of equilibrium in malaria. *Tropical Diseases Bulletin* 49(9), 813.

- Mariner, J. C., J. McDermott, J. A. P. Heesterbeek, A. Catley, and P. Roeder (2005, July). A model of lineage-1 and lineage-2 rinderpest virus transmission in pastoral areas of East Africa. *Preventive Veterinary Medicine* 69(3), 245–263.
- Mazeri, S., A. D. Gibson, N. Meunier, M. Barend, I. G. Handel, R. J. Mellanby, and L. Gamble (2018). Barriers of attendance to dog rabies static point vaccination clinics in Blantyre, Malawi. *PLoS Neglected Tropical Diseases* 12(1), e0006159.
- Minghui, R., M. Stone, M. H. Semedo, and L. Nel (2018). New global strategic plan to eliminate dog-mediated rabies by 2030. *The Lancet Global Health* 6(8), e828–e829.
- Mtema, Z., J. Chagalucha, S. Cleaveland, M. Elias, H. M. Ferguson, J. E. Halliday, D. T. Haydon, G. Jaswant, R. Kazwala, G. F. Killeen, et al. (2016). Mobile phones as surveillance tools: implementing and evaluating a large-scale intersectoral surveillance system for rabies in Tanzania. *PLoS Medicine* 13(4), e1002002.
- Rupprecht, C., J. Barrett, D. Briggs, F. Cliquet, A. Fooks, B. Lumlertdacha, F. Meslin, T. Müller, L. Nel, C. Schneider, et al. (2008). Can rabies be eradicated? *Developments in biologicals* 131, 95–121.
- Taylor, L. H., K. Hampson, A. Fahrion, B. Abela-Ridder, and L. H. Nel (2017). Difficulties in estimating the human burden of canine rabies. *Acta Tropica* 165, 133–140.
- Tepsumethanon, V., H. Wilde, and F. X. Meslin (2005). Six criteria for rabies diagnosis in living dogs. *J Med Assoc Thai* 88(3), 419–22.
- Wallace, R. M., H. Reses, R. Franka, P. Dilius, N. Fenelon, L. Orciari, M. Etheart, A. Destine, K. Crowdis, J. D. Blanton, et al. (2015). Establishment of a canine rabies burden in Haiti through the implementation of a novel surveillance program. *PLoS Neglected Tropical Diseases* 9(11), e0004245.
- Wallinga, J. and M. Lipsitch (2006). How generation intervals shape the relationship between growth rates and reproductive numbers. *Proceedings of the Royal Society B: Biological Sciences* 274(1609), 599–604.