Transmission dynamics of re-emerging rabies in domestic dogs of rural China

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Data Availability Statement: All relevant data are within the paper and its Supporting Information files. All sequences generated in this study are available in GenBank under accession numbers listed in S1 Table.

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Abstract

Despite ongoing efforts to control transmission, rabies prevention remains a challenge in many developing countries, especially in rural areas of China where re-emerging rabies is under-reported due to a lack of sustained animal surveillance. By taking advantage of detailed genomic and epidemiological data for the re-emerging rabies outbreak in Yunnan Province, China, collected between 1999 and 2015, we reconstruct the demographic and dispersal history of domestic dog rabies virus (RABV) as well as the dynamics of dog-to-dog and dog-to-human transmission. Phylogeographic analyses reveal a lower diffusion coefficient than previously estimated for dog RABV dissemination in northern Africa. Furthermore, epidemiological analyses reveal transmission rates between dogs, as well as between dogs and humans, lower than estimates for Africa. Finally, we show that reconstructed epidemic history of RABV among dogs and the dynamics of rabid dogs are consistent with the recorded human rabies cases. This work illustrates the benefits of combining phylogeographic and epidemic modelling approaches for uncovering the spatiotemporal dynamics of zoonotic diseases, with both approaches providing estimates of key epidemiological parameters.

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Introduction

Rabies remains a significant threat to public health in the 21st century [1], causing around 60,000 human fatalities worldwide each year [2]. Rabies control in the developing world is currently hindered by a lack of timely and accurate data about rabies cases in both humans and animals [3]. It is thought that the number of human deaths due to rabies virus (RABV) infections is underestimated, and that the dynamics of the virus in dog populations is poorly understood. These uncertainties inevitably hamper improvements in disease control strategies and the evaluation of control measures.

China is second only to India [4] in terms of the national number of human rabies cases, and in recent years the prevalence of rabies has increased in some areas of China [5]. More than 90% of human rabies cases in China occur in rural regions [6] where the proportion of vaccinated dogs is very low [7]. Additionally, China has a growing population of dogs, currently estimated at 80–200 million animals [6], and the breeding, management, and vaccination of dogs in the country is uncontrolled [8]. A better quantitative understanding of rabies epidemiology in dogs is needed to help predict future vaccine demand in China and other developing countries.

Dogs are the primary reservoir and vector of human rabies throughout most of Africa and Asia [9] and are responsible for more than 99% of human rabies cases [1]. Therefore, understanding the dispersal dynamics of rabies in dogs is essential for quantifying dog-to-human transmissions and for disease prevention. Previous studies have documented “traveling waves” of rabies among wildlife populations [10–16] and uncovered the genetic signature of spatial expansion in RABV genomes [17–21]. Recent studies [22–24] have attempted to combine epidemiological and genomic data to study the transmission dynamics of RABV. However, rabies spread in rural areas of China, like Yunnan Province, remains poorly documented despite an increasing incidence of human rabies. Yunnan Province in Southwest China first reported human rabies in 1956 and eliminated the disease by integrated dog management and control measures (including dog registration and the use of dog enclosures, dog vaccination, and culling of rabid/suspected rabid dogs and stray dogs) in the late 1990s. However, a re-emergence of rabies in this province was reported in 1999.

In this study, we combine phylogeographic approaches and mathematical modelling to examine the dispersal dynamics of RABV in Yunnan, a rural province of China. We analyse publicly available and newly-generated RABV gene sequences from domestic dogs, cattle, and humans sampled between 2006 and 2015, together with comprehensive rabies epidemiological data dating back to 1999, when the first re-emerging human rabies case in Yunnan was reported (Fig 1A). Specifically, we aim to use both types of data to reconstruct the demographic and dispersal history of RABV spread in Yunnan, as well as to estimate key
epidemiological parameters that can be compared to other previously-studied RABV outbreaks. Finally, we use phylogeographic inference to investigate which environmental factors may have impacted the spatial dispersal dynamics of RABV in Yunnan.

Results

Re-emergence of rabies in Yunnan

Analyses of case records date the first documented rabies case in Yunnan Province back to 1956. These data indicate that an epidemic wave of rabies occurred in the 1980s and that, from the mid-1990s onwards, human rabies cases were reported only sporadically in the region. However, starting in 1999, a new RABV epidemic emerged. More than 900 human rabies cases were reported during 1999–2015, with the highest incidence of 0.3 cases per 100,000 people occurring in 2010 (Fig 1A). Human rabies in Yunnan was first reported in 1999 in the Southeast prefecture of Wenshan, which borders Vietnam and the Chinese province of Guangxi and spread across the province in the next decade [25].

Two RABV lineages dominate the Yunnan epidemic

We first used the discrete diffusion model implemented in BEAST 1.8 to undertake a continental-scale phylogeographic analysis of a broad data set of Asian RABV sequences (S1 Table; see also the Materials and Methods for details). Genetic histories of the RABV N and G genes reveal two individual lineages (YN-A1 and YN-A2, Fig 2) that represent the majority of sampled RABV infections in the post-1999 Yunnan epidemic. We then used a GLM (generalised linear model) implementation, also available in BEAST 1.8, to measure the correlation between viral effective population size and RABV cases counts in Yunnan. This analysis estimated a GLM coefficient of 0.021 for the association between case counts and the trajectory of log effective population size of clade YN-A1 (which is the main RABV clade identified in Yunnan). The credible interval associated with this GLM coefficient estimate excludes zero (95% HPD: [0.005, 0.036]), and therefore indicates a significant association. The time to most recent common ancestor (tMRCA) estimates of the YN-A2 clade in the N and G gene largely overlap (N gene: 2007.8, 95% HPD: 2006.1–2009.2; G gene: 2008.3, 95% HPD: 2007.0–2009.4). The G gene estimate of the tMRCA of the YN-A1 lineage predates the N gene estimate (N gene: 1999.2, 95% HPD: 1996.6–2008.2; G gene: 1994.7, 95% HPD: 1992.5–1996.6).

Expanding and more localised sub-epidemics

We next performed separate continuous phylogeographic reconstructions [26] for the YN-A1 and YN-A2 clades that were identified above in the discrete phylogeographic analysis. As shown in Fig 3, the inferred RABV diffusion histories estimated from the N and G gene alignments are consistent with each other. The YN-A1 clade underwent a larger spatial expansion and is more widely distributed within dog populations, while the YN-A2 clade appears to be geographically more limited. We compared the amino acid and nucleotide sites that vary between the YN-A1 and YN-A2 clades. There were no amino acid sites changes between these clades, suggesting that the differences in lineage distributions are not associated with pathobiological features, but rather over-representation of local area induced by stochastic founder events or by potential sampling bias (see Fig 1B).

Statistics of spatial dispersal estimated from continuous phylogeographic analyses are reported in Table 1. Estimation of diffusion coefficients [27] allows different outbreaks to be directly and quantitatively compared [19]. Specifically, we found that the diffusion coefficient
of RABV in Yunnan ($D = 1733 \text{ km}^2/\text{year}$; 95% HPD = 1082–2928) is on average substantially lower than that previously estimated for RABV among North African dogs ($D = 2874 \text{ km}^2/\text{year}$; 95% HPD = 1900–5420) [28].

**Impact of environmental factors on spatial dispersal dynamics**

Several environmental variables were tested as potential factors that could affect RABV lineage dispersal velocity. As detailed in Materials and Methods, we estimated the correlation between phylogenetic branch durations and environmental distances computed on different rasters (S1 Fig). These correlations were then compared to the correlation between branch durations and distance computed on a “null” raster (i.e. an empty raster with no spatial heterogeneity whose

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Fig 1. Current wave of rabies in Yunnan Province, China. A. Upper panel: human rabies in Yunnan Province from 1976 to 2015; lower panel: sampling of RABV samples in Yunnan Province. B. Locations of the RABV sequences sampled from dogs, 2006–2015. Circle areas are proportional to the number of gene-specific sequences sampled in each location and coloured according to a given clade (dark green: N gene clade SEA-1, green: N gene clade SEA-2, yellow: N gene clade SEA-3, red: G gene clade Asian-1, orange: G gene clade Asian-2).

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cell values are set uniformly to "1"). Finally, the difference between these two correlations was assessed using a randomisation procedure. The rationale of this approach was to investigate which environmental factors can explain dispersal velocity heterogeneity better than simple geographic distances [28–30]. Complete results are reported in S2 Table, and S2 Fig reports results for the environmental factors that are most likely to have impacted virus lineage dispersal velocity. Only the forest coverage variable, tested as a potential conductance factor, was associated with a Bayes factor $>20$. Although significant, the correlation is weak: using the “forest coverage” raster to compute environmental distances only increases the correlation with branch durations by 3%, relative to the uniform null raster. This means that forest coverage (treated as a conductance factor) would be a minor contributor to the observed heterogeneity in virus lineage dispersal velocity.

In addition to the analyses investigating the impact of environmental factors on dispersal velocity, we also investigated the impact of these factors on the dispersal tendency, i.e. we investigated if lineage translocation events tended to terminate in particular environmental conditions. Specifically, we compared environmental values at the beginning (oldest node) and end (youngest node) of each branch. Differences between the environmental values at each end of phylogeny branches tested using a randomisation procedure (see Materials and Methods). This analysis also indicated...
Fig 3. Spatiotemporal diffusion of Yunnan RABV clades estimated from continuous phylogeographic reconstructions. The plots show mapped trees and 95% HPD regions based on 100 trees subsampled from the post burn-in posterior distribution of trees. Nodes of the trees are coloured according to a colour scale ranging from red (the time to the most recent common ancestor, TMRCA) to green (most recent sampling time). 95% HPD regions were computed for successive time layers and then superimposed using the same colour scale reflecting time. National and Yunnan Province borders are shown by solid and dashed lines, respectively.

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Table 1. RABV dispersion statistics estimated from continuous phylogeographic analyses.

| Statistic      | Definition                              | Median value | 95% HPD       |
|----------------|-----------------------------------------|--------------|---------------|
| $v_{\text{branch}}$ | mean branch velocity                     | 57.5 km/year | [39.2, 85.1]  |
| $v_{\text{weighted}}$ | weighted dispersal velocity             | 23.4 km/year | [2.4, 32.6]   |
| $D_{\text{original}}$ | original diffusion coefficient (Pybus et al. 2012) | 1733 km²/year | [1082, 2928]  |
| $D_{\text{weighted}}$ | weighted diffusion coefficient (Trovao et al. 2015) | 1064 km²/year | [116, 1638]   |

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Dynamic modelling from epidemiological data and dog surveillance

We used mathematical transmission models and data from human and dog surveillance of RABV outbreaks in Yunnan Province to quantify rabies spread in a rural area of China. Dog demography and human rabies data were collected by the Yunnan Institute of Endemic Disease Control and Prevention. Our model-based simulation of dog-to-human RABV transmission captures the rapid increase in human rabies at the end of the year 2000, reaching a peak around 2010, before rapidly declining from 2011 onwards (Fig 4). Moreover, we find that the simulated density of rabid dogs is highly correlated with reconstruction of viral effective population size through time. Table 3 shows the fitted parameters and values used in the model. Trace plots as well as Gelman and Rubin diagnostic indicate convergence of the MCMC chains [31] (S3 Fig). The estimated carrying capacity (K) is 13.59 dogs/km² [95% credible interval (CI): 7.80–19.37 dogs/km²]. The dog-to-dog transmission rate (β_d) is 6.62/year, [95% CI: 5.60–7.64/year]. The dog-to-human transmission rate (β_dh) is 0.0004/year, [95% CI: 0.0002–0.0006/year]. The average basic reproductive number (R₀) of the most recent wave of infection is estimated at 1.05 during the early stages of the outbreak. High values for the transmission parameters and initial conditions are positively correlated with a high peak of human rabies infections (S4 Table), and the simulations are sensitive to these parameters. Our precise field measures are crucial for reliable model simulations, in particular the dog-to-human transmission rate and dog population size.

Discussion

Human rabies is one of the biggest public health risks facing China. While current rabies surveillance mostly reflects the RABV prevalence in humans, very little is known about its prevalence in dogs. In most of mainland China, diagnosis of dog rabies is currently effectively impossible in rural areas [8], because dogs are not leashed, can move freely and have a low vaccination coverage [6]. Because of this, rural dog populations are at increased risk of exposure
and are crucial to rabies prevention and control efforts as they are the main reservoir for zoonoses [36]. Analysis of high-resolution epidemiological and genomic data provides an opportunity to explore the dynamics of re-emerging rabies dispersal and RABV transmission. Continuous phylogeographic analyses yield a lower diffusion coefficient than that previously estimated for dog RABV in northern Africa [19,28].

**Table 3.** RABV epidemiological parameters estimated from occurrence data.

| Parameter       | Definition                                             | Mean value | SD | Reference |
|-----------------|--------------------------------------------------------|------------|----|-----------|
| $b_d$           | dog birth rate                                         | 0.73/year  | 0.12 | Estimated |
| $d_d$           | dog death rate                                         | 0.26/year  | 0.12 | Estimated |
| $K$             | dog carrying capacity                                  | 11.41 dogs/km$^2$ | 0.83 | Estimated |
| $\sigma_d$      | $1/\sigma_d$ is the average latent period              | 13.43/year | 1.82 | Estimated |
| $\alpha_d$      | death rate of rabid dogs                               | 63.96/year | 1.96 | Estimated |
| $\beta_d$       | dog-to-dog transmission rate                           | 6.62/year  | 0.52 | Estimated |
| $\beta_{dh}$    | dog-to-human transmission rate                         | 0.0004/year | 0.0001 | Estimated |
| $S_0$           | initial susceptible dog population size                | 9.50 dogs/km$^2$ | 0.96 | Estimated |
| $E_0$           | initial exposed dog population size                    | 0.003 dogs/km$^2$ | 0.003 | Estimated |
| $I_0$           | initial infectious dog population size                 | 0.003 dogs/km$^2$ | 0.003 | Estimated |
| $\nu_d$         | dog vaccination rate                                   | 0.07/year  | -   | [32] Fixed |
| $\epsilon$      | dog vaccination success rate                            | 0.60/year  | -   | [32] Fixed |
| $\lambda_d$     | dog loss of vaccination immunity rate                  | 1/year     | -   | [33] Fixed |
| $\lambda_h$     | human loss of vaccination immunity rate                | 1/year     | -   | Fixed     |
| $\sigma_h$      | $1/\sigma_h$ is the average human incubation period    | 6/year     | -   | [34] Fixed |
| $\alpha_h$      | rabid humans mortality rate                            | 1/year     | -   | Fixed     |
| $\beta_h$       | human birth rate                                       | 0.013/year | -   | [35] Fixed |
| $d_h$           | human mortality rate                                   | 0.006/year | -   | [35] Fixed |

*standard deviation

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Africa data set also revealed the importance of human-related environmental factors (human population density, accessibility to nearest major cities) as explanatory factors for the heterogeneity in RABV lineage dispersal velocity in Africa [28]. The higher diffusion coefficient of the northern African RABV data set was thus suspected to be related to human-based connectivity and/or mobility. Consequently, the lower diffusion coefficient estimated for the Yunnan data set could, in turn, indicate that human-related movements are less important contributors to the spatial dissemination of RABV in Yunnan, although we note that this is only one possible interpretation. Here, we also investigated the potential impact of several environmental factors and found a significant but weak effect of forest coverage acting (directly or indirectly) as a factor favouring RABV lineage dispersal velocity in Yunnan. One potential interpretation of this result is that in sparsely populated forest areas a few rapid (and potentially human-related) movements of infected dogs may have occurred. Such lineage dispersal movements could have happened via the road network crossing forest areas. However, even if identified as a significant conductance factor, the forest coverage in itself is not necessarily the causal factor and could instead be correlated with the true causal factor that is not included in the present analysis.

The potential importance of forest coverage was further indicated by comparing environmental conditions at branch termination locations. This analysis mainly revealed that viral lineages did not tend to spread towards forest areas. However, this result could also reflect, to some extent, the impact of a potential sampling bias arising if there were a lower sampling probability in less densely human-populated regions, such as forest areas. Furthermore, even under the assumption that viral lineages would be less likely to spread towards forest areas, our analysis based on dispersal velocity revealed that these areas do not act as barriers decreasing lineages dispersal velocity among infected areas.

Our dynamic model provides estimates of epidemiological parameters for dog rabies in Yunnan. The simulated dog population indicated a low level of RABV transmission, different from the oscillations observed in N’Djamena, Africa [11]. We also obtained an estimate of the transmission rate among dogs ($\beta_d$), and from dogs to humans ($\beta_{dh}$); key epidemiological parameters for assessment of rabies outbreaks and incidence among dogs. The estimate ($\beta_d$) of 6.62/year was higher than the 4.20/year estimated in N’Djamena, Chad [34], and lower than 14.68/year estimated in Machakos District, Kenya [37,38]. The transmission rate from dogs to humans ($\beta_{dh}$) was estimated to be 0.0004/year, which is consistent with our field investigation of dog bites (0.0004/year) in the study area [39], but much lower than the 0.0107/year estimated in N’Djamena [34]. In addition, the estimated carrying capacity (13.59 dogs/km$^2$, 95% CI: 7.80–19.37 dogs/km$^2$) was also lower than 33.6 dogs/km$^2$ estimated in N’Djamena, Chad, but was similar to the value estimated in Tamil Nadu, India (12.69 dogs/km$^2$, estimated using livestock census data) [40]. We suspect these may result from socio-economic differences, e.g. human population density [41].

Despite a national rabies control and prevention program that was implemented in China in 1985, which resulted in a drastic reduction in human rabies cases in the 1990s, a new RABV epidemic started in China in 1999 (Fig 1). Phylogeographic reconstructions confirm the East-to-West invasion history of RABV in Yunnan, which was previously suggested by epidemiological records [25]. Furthermore, we find a time lag between the peaks of viral effective population size and the corresponding peaks of rabid dog numbers (Fig 4). This may be due to the substantial proportion of undetected dog rabies infections; epidemiological surveillance of dog rabies in the region is far from exhaustive.

This study illustrates how RABV dispersion dynamics can be analysed by two complementary approaches based on genetic and epidemiological data. Overall, our work further highlights that domestic dogs play a key role in transmission and expansion of rabies in rural areas.
of China. The data reveal a low level of RABV transmission among dogs, and from dogs to humans, with the basic reproductive number estimated at 1.05 during the early stages of the outbreak. Our results indicate that interventions in the dog population would be effective in reducing transmission to humans, in particular because they have the potential to subvert the self-sustaining capacity of epidemics in dogs. A better understanding of RABV spread in terms of spatial and temporal dynamics is necessary to help inform the prevention and control of human rabies in the vast rural areas of China.

**Material and methods**

**Ethics statement**

It was determined by the Yunnan Institute of Endemic Diseases Control and Prevention, that the collection of data from rabies cases was part of continuing public health surveillance of a notifiable infectious disease and was exempt from institutional review board assessment. Experimental procedures were performed in compliance with guidelines established by the Chinese Center For Disease Control And Prevention and have been approved by ethics committee of Yunnan Institute of Endemic Diseases Control and Prevention.

**New samples from Yunnan**

From 2008 to 2015, we collected 1392 brain tissue samples. These samples were obtained from 252 dogs suspected of having rabies, 1129 apparently healthy domestic dogs, 2 cows, and 9 human patients within 24 hours of death in 14 prefectures and 43 counties of Yunnan Province (Fig 1B). In addition, 18 saliva samples and 1 cerebrospinal fluid sample were obtained from surviving patients.

All brain specimens were tested using direct immunofluorescence assay (DFA) and RABV nucleoprotein monoclonal antibody (Rabies DFA Reagent; Chemicon, Temecula, CA, USA). Total RNA was extracted from the original brain samples with the Trizol reagent (Invitrogen, USA) according to the manufacturer’s instructions [25]. PCR products were purified by using a QIAquick PCR Purification Kit (QIAGEN, Germany). Complete nucleoprotein gene (N gene; 1 cow, 84 dog and 6 human isolates) and glycoprotein gene (G gene; 2 cow, 110 dog and 8 human isolates) sequences were obtained by using previously described primers [21,42]. Newly generated RABV sequence data were submitted to GenBank (accession numbers KP072009–KP072030, KP202418- KP202448, KT932670-KT932698 and KX096992-KX097000 for N gene, and JF819597-JF819602, JQ040570-JQ040581, JX276383-JX276404, KP072031-KP072052, KP202402-KP202417, KT861554-KT861586 and KX096983–KX096991 for G gene). An overview of the newly sequenced isolates is provided in S1 Table.

**Incidence data from the Yunnan RABV epidemic**

Human rabies case records (Fig 1A) in Yunnan from 1999 to 2015 were obtained from the Yunnan Center for Disease Control and Prevention (CDC) and the Chinese CDC. In China, human rabies is a class B notifiable infectious disease and all human cases must be reported to the Chinese CDC. Rabies cases were confirmed according to diagnostic criteria (WS281–2008) from the Ministry of Health of the People’s Republic of China.

**Discrete phylogeographic analyses**

To identify Yunnan-specific RABV circulation, nucleotide sequences of the N and G genes of all available RABV sequences sampled from non-flying mammals in Asia were downloaded from NCBI GenBank and aligned [43] together with our newly generated sequences. The
resulting data sets included 543 N sequences (the entire N gene coding sequence, 1350 nt long) and 491 G sequences (the entire G gene coding sequence, 1575 nt long). RABV clades specific to Yunnan were identified using the discrete trait analysis model implemented in BEAST 1.8 [44,45]. For the molecular clock phylogeographic analyses, we specified a general time-reversible GTR+I nucleotide substitution model [46–48], a skygrid coalescent model [49] and a relaxed uncorrelated lognormal (UCLN) molecular clock model across branches [50]. The data sets lack a clear temporal signal [51] (S1 Fig). Therefore, we specified informative prior distributions on the gene-specific mean clock rate parameter. These prior distributions were based on previously published estimated substitution rates [52]: $1.88 \times 10^{-4}$ substitutions/site/year (95% highest posterior density, HPD: [1.37×10^{-4}, 2.41×10^{-4}]) for the N gene and $2.13 \times 10^{-4}$ substitutions/site/year (95% HPD: [1.56×10^{-4}, 2.73×10^{-4}]) for the G gene. The MCMC chain was run for 250 million states and mixing and convergence were inspected using Tracer [http://tree.bio.ed.ac.uk/software/tracer/]. TreeAnnotator 1.8 [44] was used to infer maximum clade credibility (MCC) summary trees.

Testing covariates of viral population size

We used the GLM (generalised linear model) extension of the skygrid coalescent model [53], implemented in BEAST 1.8, to simultaneously infer viral effective population sizes and measure the association between estimated effective population size and cases counts. For this analysis, we focused only on sequences associated with the largest Yunnan clade identified by the discrete phylogeographic analysis (clade YN-A1, see the Results section). Furthermore, we considered the N and G genes as two independent markers (see Appendix S1 in S1 Text for the detailed procedure).

Continuous phylogeographic analyses

The history of virus lineage dispersal in Yunnan was recovered from geo-referenced phylogenies, which were estimated using the continuous phylogeographic method [26] implemented in BEAST 1.8. A separate continuous phylogeographic analysis was performed for each gene and each Yunnan clade identified by the preliminary discrete phylogeographic analyses; however, the nucleotide substitution and molecular clock models were linked across the clades to avoid over-parameterisation. For these models, we used the same models as described above, together with a relaxed random walk model for inference of the continuous spatial locations. This relaxed random walk model assumed a log-normal probability distribution among phylogeny branches of diffusion rate scalars. The spatiotemporal information contained in inferred phylogenetic trees was extracted with the R package “seraphim” [29,30]. For the present study, we extracted spatiotemporal information from a subset of 1,000 trees sampled at regular intervals from the posterior distribution of trees (after burn-in had been removed). This was done for each gene and for each Yunnan clade. After this extraction step, each phylogeny branch is represented as a distinct movement vector [27]. “seraphim” was also used to estimate statistics of spatial dispersal based on these extracted movement vectors. We estimated the mean branch velocity, the weighted dispersal velocity, the diffusion coefficient (as originally defined in Pybus et al. [27]) and the weighted diffusion coefficient (as defined by Trovão et al. [54]). Further details regarding these statistics can be found in the “seraphim” package [30].

Impact of environmental factors on spatial dispersal dynamics

We next sought to investigate the impact of environmental factors on lineage dispersal velocity. These analyses followed a similar structure to those used in previous studies [28,29,55]. All
scripts for these analyses are available in the R package "seraphim" [30]. Here, we investigated the impact of the following environmental variables (S1 Fig): elevation, annual mean temperature, annual precipitation, key land cover variables (e.g. "grasslands", "savannas", "forests", "croplands", "urban areas"); land cover categorised according to the International Geosphere Biosphere Program, IGBP), human population density, human footprint, major roads, and inaccessibility (quantified as the time it takes to travel to the nearest major city of >50,000 inhabitants). The sources of the data in the original raster files are listed in S3 Table. All factors were tested as potential conductance factors (i.e. factors facilitating movement) and as potential resistance factors (i.e. factors impeding movement). Correlations between phylogenetic branch durations and environmentally-scaled distances were quantified as a statistic representing the difference between two coefficients of determination (R²): (i) the R² obtained when branch durations are regressed against environmentally-scaled distances, and (ii) the R² obtained when branch durations are regressed against distances computed on a “null” raster, i.e. a raster with a value of “1” assigned to every cell. An environmental factor was only considered as potentially explanatory if both its distribution of regression coefficients and its associated distribution of Q values were positive [56]. In a positive distribution of estimated Q values (i.e. with at least 90% of positive values), statistical support was then evaluated against a null distribution generated by a randomisation procedure and formalised using a Bayes factor (BF) value [28] (see Appendix S2 in S1 Text for the full procedure). Due to computational limits, this analysis was based on 100 trees subsampled from each post-burn-in posterior distribution obtained by continuous phylogeographic inference.

Addition to the analyses based on lineage dispersal velocity, we also used a new analytical procedure to investigate the impact of several environmental factors on dispersal tendency. In order words, this procedure aims at testing if virus lineages tend to disperse towards particular environmental conditions. In this framework, environmental conditions are compared between the locations of the two nodes connected by a phylogeny branch. For each branch and environmental factor, we computed the difference in raster cell values between the start (oldest node) location and at the end (youngest node) location. These differences were then averaged within each sampled tree and evaluated against a null distribution generated by the same randomisation procedure used for the analyses of the impact on dispersal velocity (see Appendix S3 in S1 Text for the detailed procedure).

**Dynamic modelling based on epidemiological data and dog surveillance**

In the dynamic modelling part of this study, we extended the discrete susceptible-exposed-infected (SEI) model for dog rabies [10,11,34]. Our discrete-time model framework was developed by dividing a closed dog population into four rabies classes, susceptible (Sₗ), exposed (Eₗ), and infectious (Iₗ), and vaccinated (Vₗ). We then extended the model for dog rabies to include dog-to-human RABV transmission. A similar modelling framework has been applied to quantitatively assess the dynamics of RABV transmission [34]. Sₗ is the number of people susceptible to disease, Pₗ is the annual vaccinated individuals in Yunnan (based on medical records from Chinese Center For Disease Control and Prevention), Eₗ and Vₗ represents the number of exposed humans and immunized humans during the time period, respectively.

\[
S_{d,t+1} = S_{d,t} + b\dot{N}_{d,t} + \dot{\lambda}_d V_{d,t} - d_d S_{d,t} - \gamma N_d S_{d,t} - \beta_d S_{d,t} I_{d,t} - ve S_{d,t}
\]

\[
E_{d,t+1} = E_{d,t} + \beta_d S_{d,t} I_{d,t} - d_d E_{d,t} - \gamma N_d E_{d,t} - \sigma E_{d,t} - ve E_{d,t}
\]

\[
I_{d,t+1} = I_{d,t} + \sigma E_{d,t} - d_d I_{d,t} - \gamma N_d I_{d,t} - \alpha_d I_{d,t}
\]
of 1% of the total dog population based on empirical estimation, due to the temporal irregular
mates of the initial susceptible dog population size,
and changing intensity of potential culling. The susceptible reconstruction provided initial esti-
ation rate ($\lambda$) be density-dependent, with birth ($b_d$) and death ($d_d$) rates. The density-dependent mortality
parameter ($\gamma$) is a function of the dog population growth rate per year ($b_d - d_d$) and the carrying
capacity density of the dog population in km$^{-2}$ ($K$) is defined by $\gamma = (b_d - d_d) / K$. Carrying capacity
is defined as the maximum number or density of individuals that can be sustainably supported
by a given environment. In rural areas of China, many domestic dogs are free-roaming and there
has been no population control for most of the time. We can then assume that dog populations
(including owned and stray dogs) are regulated through represented by carrying capacity deter-
mined by some unknown combination of environmental and human geographic factors. $\beta_d$ and
$\beta_{dh}$ are the dog-to-dog and dog-to-human transmission rate, respectively. $\sigma_d$ is the latent-to-infect-
ious rate per year and $\alpha$ is the rabies-induced mortality. At the beginning ($t = 0$), the basic repro-
ductive number ($R_0$) is $\alpha \beta_d S_0 / (\sigma_d + \alpha + \beta_d)$). Informative prior distributions are assumed for death
rate for rabid dog/human, latent period, incubation period, as well as dog-to-human transmission
rate ($\beta_{dh}$), which is assessed from our field investigation of dog bites [39]. We assumed dog vacci-
nation rate ($\nu$) of 0.07 and dog vaccination success rate ($c$) of 0.60 in Yunnan by previous surveil-
ance [32]. $\lambda$ is loss of vaccination immunity rate. Culling was simulated assuming the elimination
of 1% of the total dog population based on empirical estimation, due to the temporal irregular
and changing intensity of potential culling. The susceptible reconstruction provided initial esti-
mates of the initial susceptible dog population size, $S_{d0}$, for which we chose a uniform prior of
5–15 dogs/km$^2$, according to our surveillance [32]. Initial numbers for $E_{d0}$ and $I_{d0}$ were unknown
and fitted in the model with a wide prior 0–1 dogs/km$^2$.

We fitted the dynamic model using the Bayesian state space framework using Metropolis-
Hastings Markov chain Monte Carlo algorithms implemented in the MATLAB code [57,58]
toolbox DRAM (Delayed Rejection Adaptive Metropolis) [57,58]. The chain was initiated
using published estimates of dog demography and RABV transmission rates [34,38,59–61]
with an uninformative prior that varied from 0 to infinity, and then was run for 5 million iter-
ations and sampled every 5000th step, after a burn-in of 500,000 iterations.

Sensitivity analysis was performed using latin hypercube sampling (LHS) and partial rank
 correlation coefficient (PRCC) techniques [62]. Two criteria were retained as outputs for the
analysis: the intensity of the peak of the human rabies infections and the date of the peak. Uni-
form distributions were used for all parameters and the same ranges as for the prior distributions
were used. A total of 1,000 parameter sets were sampled with LHS and PRCC was
estimated using the MATLAB code provided by Marino et al. [63].
Supporting information

**S1 Text.** Appendix S1. Covariate analysis. Appendix S2. Analysis of the impact of environmental factor on RABV dispersal velocity in Yunnan. Appendix S3. Analysis of the impact of environmental factor on RABV dispersal tendency in Yunnan.

**S1 Fig.** Environmental variables that were tested in the analysis of the dog RABV dataset from Yunnan (China). National and Yunnan province borders are respectively displayed by solid and dashed lines, respectively.

**S2 Fig.** Potential impact of environmental factors on RABV dispersal velocity in Yunnan. The results are based on 100 trees sampled from the posterior distribution. “C” and “R” indicate if the considered environmental variable was considered as a conductance (“C”) or resistance factor (“R”), and $k$ is the rescaling parameter used to transform the initial raster. Estimated Q distributions and related Bayes factor (BF) supports are reported here only for those model combinations with >90% of positive Q values and a BF support > 3 (see S3 Table for complete results). BF supports were estimated with the randomisation procedure detailed in Appendix S1 in S1 Text. Following Kass & Raftery (1995) we consider a BF > 3 and BF ≥20 respectively as a “positive” and “strong” evidences of the statistical significance of Q, i.e. the correlation between environmental distances and dispersal durations (see text for further details).

**S3 Fig.** Trace plots of four chains for each of the parameters. Four chains were initialized at different values, and they converge to the target distribution quickly.

**S4 Fig.** Plots of the root-to-tip genetic distance against sampling time for N gene and G gene data sets of Yunnan. TempEst was used to obtain exploratory regressions based on the maximum likelihood trees.

**S1 Table.** List of RABV sequences analysed in this study.

**S2 Table.** Impact of several environmental factors on RABV dispersal velocity. The results are based on 100 trees sampled in each posterior distribution. “C” and “R” indicate if the considered environmental raster was considered as a conductance (“C”) or resistance factor (“R”), and $k$ is the rescaling parameter used to transform the initial raster (see the Appendix S2 in S1 Text for further details). For regression coefficients and Q values we report both the median estimate and the 95% HPD interval. The Bayes factor (BF) supports based on the randomisation procedure is only reported when $p(Q > 0)$ is at least 90%. Following Kass & Raftery (1995) we consider a Bayes factor (BF) >3 as positive support for a significant correlation between the environmental distances and dispersal durations.

**S3 Table.** Source of data for each environmental raster.

**S4 Table.** Sensitivity analysis. For model parameters, we used the same range as for the prior distribution. 1000 parameter sets were sampled with latin hypercube sampling, and partial
rank correlation coefficients were estimated.

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References

1. Fooks AR, Banyard AC, Horton DL, Johnson N, McElhinney LM, et al. (2014) Current status of rabies and prospects for elimination. Lancet 384: 1389–1399. https://doi.org/10.1016/S0140-6736(13)62707-5 PMID: 24828901

2. Hampson K, Cleaveland S, Briggs D (2011) Evaluation of cost-effective strategies for rabies post-exposure vaccination in low-income countries. PLoS Negl Trop Dis 5: e982. https://doi.org/10.1371/journal.pntd.0000982 PMID: 21408121

3. Knobel DL, Cleaveland S, Coleman PG, Fèvre EM, Meltzer MI, et al. (2005) Re-evaluating the burden of rabies in Africa and Asia. Bull World Health Organ 83: 360–368. PMID: 15976877

4. Zhou H, Vong S, Liu K, Li Y, Mu D, et al. (2016) Human rabies in China, 1960–2014: A descriptive epidemiological study. PLoS Negl Trop Dis 10: e0004874. https://doi.org/10.1371/journal.pntd.0004874 PMID: 27500957

5. Ma C, Hao X, Deng H, Wu R, Liu J, et al. (2017) Re-emerging of rabies in Shaanxi Province, China, 2009 to 2015. J Med Virol: https://doi.org/10.1002/jmv.24769 PMID: 28112421

6. Tang X, Luo M, Zhang S, Fooks AR, Hu R, et al. (2005) Pivotal role of dogs in rabies transmission, China. Emerg Infect Dis 11: 1970–1972. https://doi.org/10.3201/eid1112.050271 PMID: 16485494
7. Deng Z, Qiu H, Hong J (2011) Factors related to rabies prevalence in Kunming City from 1981 to 2010. China Tropical Medicine 11: 1251–1252.
8. Hu R, Tang Q, Tang J, Fooks AR (2009) Rabies in China: an update. Vector Borne Zoonotic Dis 9: 1–12. https://doi.org/10.1089/vbz.2008.0046 PMID: 18803503
9. Cleaveland S, Kaare M, Knobel D, Laurenson MK (2006) Canine vaccination—providing broader benefits for disease control. Vet Microbiol 117: 43–50. https://doi.org/10.1016/j.vetmic.2006.04.009 PMID: 16701966
10. Anderson R, Jackson H, May R, Smith A (1981) Population dynamics of fox rabies in Europe. Nature 289: 765–771. PMID: 7464941
11. Hampson K, Dushoff J, Bingham J, Brückner G, Ali Y, et al. (2007) Synchronous cycles of domestic dog rabies in sub-Saharan Africa and the impact of control efforts. Proc Natl Acad Sci USA 104: 7717–7722. https://doi.org/10.1073/pnas.0609122104 PMID: 17452645
12. Murray J, Stanley E, Brown D (1986) On the spatial spread of rabies among foxes. Proc R Soc Lond B Biol Sci 229: 111–150. PMID: 2880348
13. Childs JE, Curns AT, Dey ME, Feinstein L, et al. (2000) Predicting the local dynamics of epizootic rabies among raccoons in the United States. Proc Natl Acad Sci USA 97: 13666–13671. https://doi.org/10.1073/pnas.240326697 PMID: 11069300
14. Russell CA, Smith DL, Waller LA, Childs JE, Real LA (2004) A priori prediction of disease invasion dynamics in a novel environment. Proc Biol Sci 271: 21–25. https://doi.org/10.1098/rspb.2003.2559 PMID: 15002767
15. Smith DL, Lucey B, Waller LA, Childs JE, Real LA (2002) Predicting the spatial dynamics of rabies epizootics on heterogeneous landscapes. Proc Natl Acad Sci USA 99: 3668–3672. https://doi.org/10.1073/pnas.042400799 PMID: 11904426
16. Hampson K, Dushoff J, Cleaveland S, Haydon DT, Kaare M, et al. (2009) Transmission dynamics and prospects for the elimination of canine rabies. PLoS Biol 7: e1000053.
17. Biek R, Henderson JC, Waller LA, Rupprecht CE, Real LA (2007) A high-resolution genetic signature of demographic and spatial expansion in epizootic rabies virus. Proc Natl Acad Sci USA 104: 7993–7998. https://doi.org/10.1073/pnas.0700741104 PMID: 17470818
18. Real LA, Henderson JC, Biek R, Snaman J, Jack TL, et al. (2005) Unifying the spatial population dynamics and molecular evolution of epidemic rabies virus. Proc Natl Acad Sci USA 102: 12107–12111. https://doi.org/10.1073/pnas.0500057102 PMID: 16103538
19. Talbi C, Lemey P, Suchard MA, Abdelatif E, Elharra M, et al. (2010) Phylogeography and human-mediated dispersal of a zoonotic virus. PLoS Pathog 6: e1001166. https://doi.org/10.1371/journal.ppat.1001166 PMID: 21060816
20. Yao H-W, Yang Y, Liu K, Li X-L, Zuo S-Q, et al. (2015) The spatiotemporal expansion of human rabies and its probable explanation in mainland China, 2004–2013. PLoS Negl Trop Dis 9: e0003502. https://doi.org/10.1371/journal.pntd.0003502 PMID: 25692883
21. Tao X, Tang Q, Li H, Mo Z, Zhang H, et al. (2009) Molecular epidemiology of rabies in Southern People’s Republic of China. Emerg Infect Dis 15: 1192–1198. https://doi.org/10.3201/eid1508.081551 PMID: 19751579
22. Bourhy H, Nakouné E, Hall M, Nouvellet P, Lepelletier A, et al. (2016) Revealing the micro-scale signature of endemic zoonotic disease transmission in an African urban setting. PLoS Pathog 12: e1005525. https://doi.org/10.1371/journal.ppat.1005525 PMID: 27058957
23. Duke-Sylvester SM, Biek R, Real LA (2013) Molecular evolutionary signatures reveal the role of host ecological dynamics in viral disease emergence and spread. Philos Trans R Soc Lond B Biol Sci 368: 20120194. https://doi.org/10.1098/rstb.2012.0194 PMID: 23382419
24. Brunker K, Marston DA, Horton DL, Cleaveland S, Fooks AR, et al. (2015) Elucidating the phylogenetics of endemic rabies virus in eastern Africa using whole-genome sequencing. Virus Evol 1: vex011. https://doi.org/10.1093/ve/vex011 PMID: 27774283
25. Zhang H, Zhang Y, Yang W, Tao X, Li H, et al. (2014) Molecular epidemiology of reemergent rabies in Yunnan Province, southwestern China. Emerg Infect Dis 20: 1433. https://doi.org/10.3201/eid2009.130440 PMID: 25144604
26. Lemey P, Rambaut A, Welch JJ, Suchard MA (2010) Phylogeography takes a relaxed random walk in continuous space and time. Mol Biol Evol 27: 1877–1885. https://doi.org/10.1093/molbev/msq067 PMID: 20203288
27. Pybus OG, Suchard MA, Lemey P, Bernardin FJ, Rambaut A, et al. (2012) Unifying the spatial epidemiology and molecular evolution of emerging epidemics. Proc Natl Acad Sci USA 109: 15066–15071. https://doi.org/10.1073/pnas.1206598109 PMID: 22927414
28. Dellicour S, Rose R, Faria NR, Vieira LFP, Bourhy H, et al. (2017) Using viral gene sequences to compare and explain the heterogeneous spatial dynamics of virus epidemics. Mol Biol Evol: https://doi.org/10.1093/molbev/msx1176

29. Dellicour S, Rose R, Pybus OG (2016) Explaining the geographic spread of emerging epidemics: a framework for comparing viral phylogenies and environmental landscape data. BMC bioinformatics 17: 82. https://doi.org/10.1186/s12859-016-0924-x PMID: 26864798

30. Dellicour S, Rose R, Faria NR, Lemey P, Pybus OG (2016) SERAPHIM: studying environmental rasters and phylogenetically-informed movements. Bioinformatics 32: 3204–3206. https://doi.org/10.1093/bioinformatics/btw384 PMID: 27334476

31. Laneri K, Bhadra A, Ionides EL, Bouma M, Dhiman RC, et al. (2010) Forcing versus feedback: epidemic malaria and monsoon rains in northwest India. PLoS Comput Biol 6: e1000898. https://doi.org/10.1371/journal.pcbi.1000898 PMID: 20824122

32. Dong G, Ma K, Zhang W, Yang P (2011) Current situation and control of rabies in Yunnan Province. Modern Journal of Animal Husbandry and Veterinary Medicine: 32–33.

33. Lv Y, Lv S (2012) Some thoughts on rabies disease prevention in Yunnan Province. Yunnan Journal of Animal Science and Veterinary Medicine: 44–45.

34. Zinsstag J, Durr S, Penny M, Mindekem R, Roth F, et al. (2009) Transmission dynamics and economics of rabies control in dogs and humans in an African city. Proc Natl Acad Sci USA 106: 14996–15001. https://doi.org/10.1073/pnas.0904740106 PMID: 19706492

35. Statistical Bureau of Yunnan Province (2015) Yunnan statistical yearbook. Beijing: China Statistical Press.

36. Zinsstag J, Lechene R, Laager M, Mindekem R, Oussiguéré A, et al. (2017) Vaccination of dogs in an African city interrupts rabies transmission and reduces human exposure. Sci Transl Med 9: eaaf6984. https://doi.org/10.1126/scitranslmed.aaf6984

37. Kitala P, McDermott J, Kyule M, Gathuma J, Perry B, et al. (2001) Dog ecology and demography information to support the planning of rabies control in Machakos District, Kenya. Acta Trop 78: 217–230. PMID: 11311185

38. Kitala P, McDermott J, Kyule M, Gathuma J (2000) Community-based active surveillance for rabies in Machakos District, Kenya. Prev Vet Med 44: 73–85. PMID: 10727745

39. Hu Q, Hu H, Zhang Y, Wu X, Yang W, et al. (2016) Analysis of a surveillance system based on aggressive dog attack incidents in chuxiong prefecture. J Med Pest Control 32: 1005–1007.

40. Gwatkin DR, Rutstein S, Johnson K, Suliman E, Wagstaff A, et al. (2007) Socio-economic differences in health, nutrition, and population within developing countries. Washington, DC: World Bank.

41. Yu J, Li H, Tang Q, Rayner S, Han N, et al. (2012) The spatial and temporal dynamics of rabies and canine rabies in China. PLoS Negl Trop Dis 6: e1649. https://doi.org/10.1371/journal.pntd.0001640 PMID: 22565318

42. Katoh K, Standley DM (2013) MAFFT multiple sequence alignment software version 7: improvements in performance and usability. Mol Biol Evol 30: 772–780. https://doi.org/10.1093/molbev/mss265 PMID: 23529690

43. Drummond AJ, Suchard MA, Xie D, Rambaut A (2012) Bayesian phylogenetics with BEAUti and the BEAST 1.7. Mol Biol Evol 29: 1969–1973. https://doi.org/10.1093/molbev/mss075 PMID: 22367748

44. Lemey P, Rambaut A, Drummond AJ, Suchard MA (2009) Bayesian phylogeography finds its roots. PLoS Comput Biol 5: e1000520. https://doi.org/10.1371/journal.pcbi.1000520 PMID: 19779555

45. Yang Z (1994) Maximum likelihood phylogenetic estimation from DNA sequences with variable rates over sites: approximate methods. J Mol Biol 39: 306–314. PMID: 7932792

46. Tavaré S (1986) Some probabilistic and statistical problems in the analysis of DNA sequences. In: Miura R. M., editor. Some Mathematical Questions in Biology—DNA Sequence Analysis. Providence, RI: Amer Math Soc. pp. 57–86.

47. Shoemaker J, Fitch W (1989) Evidence from nuclear sequences that invariable sites should be considered when sequence divergence is calculated. Mol Biol Evol 6: 270–289. https://doi.org/10.1093/oxfordjournals.molbev.a010550 PMID: 2622335

48. Gill MS, Lemey P, Faria NR, Rambaut A, Shapiro B, et al. (2013) Improving Bayesian population dynamics inference: a coalescent-based model for multiple loci. Mol Biol Evol 30: 713–724. https://doi.org/10.1093/molbev/ms3265 PMID: 23180580

49. Drummond AJ, Ho SY, Phillips MJ, Rambaut A (2006) Relaxed phylogenetics and dating with confidence. PLoS Biol 4: e88. https://doi.org/10.1371/journal.pbio.0040088 PMID: 16683862
51. Rambaut A, Lam TT, Carvalho LM, Pybus OG (2016) Exploring the temporal structure of heterochronous sequences using TempEst (formerly Path-O-Gen). Virus Evol 2: vew007. https://doi.org/10.1093/ve/vew007 PMID: 27774300

52. Zhang Y, Vrancken B, Feng Y, Dellicour S, Yang Q, et al. (2017) Cross-border spread, lineage displacement and evolutionary rate estimation of rabies virus in Yunnan Province, China. Virol J 14: 102. https://doi.org/10.1186/s12985-017-0769-6 PMID: 28578663

53. Gill MS, Lemey P, Bennett SN, Biek R, Suchard MA (2016) Understanding past population dynamics: Bayesian coalescent-based modeling with covariates. Syst Biol 65: 1041–1056. https://doi.org/10.1093/sysbio/syw050 PMID: 27368344

54. Trovão NS, Suchard MA, Baelde G, Gilbert M, Lemey P (2015) Bayesian inference reveals host-specific contributions to the epidemic expansion of Influenza A H5N1. Mol Biol Evol 32: 3264–3275. https://doi.org/10.1093/molbev/msv185 PMID: 26341298

55. Laenen L, Dellicour S, Vergote V, Nauwelaers I, De Coster S, et al. (2016) Spatio-temporal analysis of Nova virus, a divergent hantavirus circulating in the European mole in Belgium. Mol Ecol 25: 5994–6008. https://doi.org/10.1111/mec.13887 PMID: 27862516

56. Jacquot M, Nomikou K, Palmarini M, Mertens P, Biek R (2017) Bluetongue virus spread in Europe is a consequence of climatic, landscape and vertebrate host factors as revealed by phylogeographic inference. Proc Biol Sci 284: 20170919. https://doi.org/10.1098/rspb.2017.0919 PMID: 29021180

57. Chib S, Greenberg E (1995) Understanding the metropolis-hastings algorithm. Am Stat 49: 327–335.

58. Haario H, Laine M, Mira A, Saksman E (2006) DRAM: efficient adaptive MCMC. Stat Comput 16: 339–354.

59. Cleaveland S, Dye C (1995) Maintenance of a microparasite infecting several host species: rabies in the Serengeti. Parasitology 111: S33–S47. PMID: 8632923

60. Kitala P, McDermott J, Coleman P, Dye C (2002) Comparison of vaccination strategies for the control of dog rabies in Machakos District, Kenya. Epidemiol Infect 129: 215–222. PMID: 12211590

61. Hua M, Zhang Y, Zhang Z, Liu R, Yang W, et al. (2013) Rabies virus carriage in domestic dogs and distribution of positive dogs in Lufenng, Yunnan. Disease Surveillance 28: 853–856.

62. Blower SM, Dowlatabadi H (1994) Sensitivity and uncertainty analysis of complex models of disease transmission: an HIV model, as an example. International Statistical Review/Revue Internationale de Statistique 62: 229–243.

63. Marino S, Hogue IB, Ray CJ, Kirschner DE (2008) A methodology for performing global uncertainty and sensitivity analysis in systems biology. J Theor Biol 254: 178–196. https://doi.org/10.1016/j.jtbi.2008.04.011 PMID: 18572196