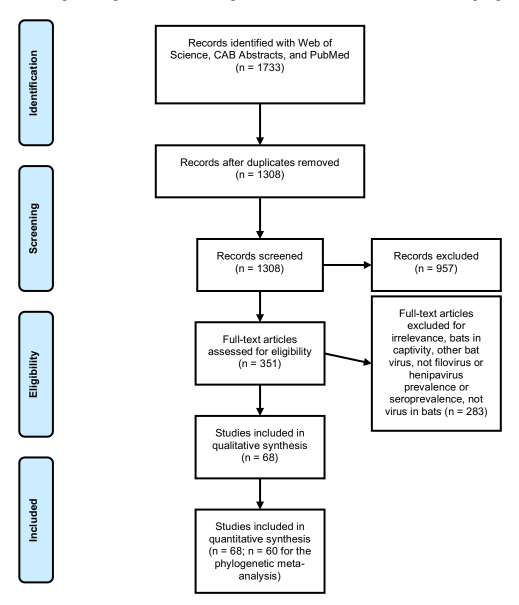
# Temporal and spatial limitations in global surveillance for bat filoviruses and henipaviruses: Online Appendix

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- S1. Systematic search
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#### S1. Systematic search

Figure S1. The data collection and inclusion process for studies of wild bat filovirus and henipavirus prevalence and seroprevalence (PRISMA diagram). Searches used the following string: (bat\* OR Chiroptera\*) AND (filovirus OR henipavirus OR "Hendra virus" OR "Nipah virus" OR "Ebola virus" OR "Marburg virus" OR ebolavirus OR marburgvirus). Searches were run during October 2017 and again in August 2019, supplemented by extracting data from references cited in identified studies. Publications were excluded if they did not assess filovirus or henipavirus prevalence or seroprevalence in wild bats or were in languages other than English.



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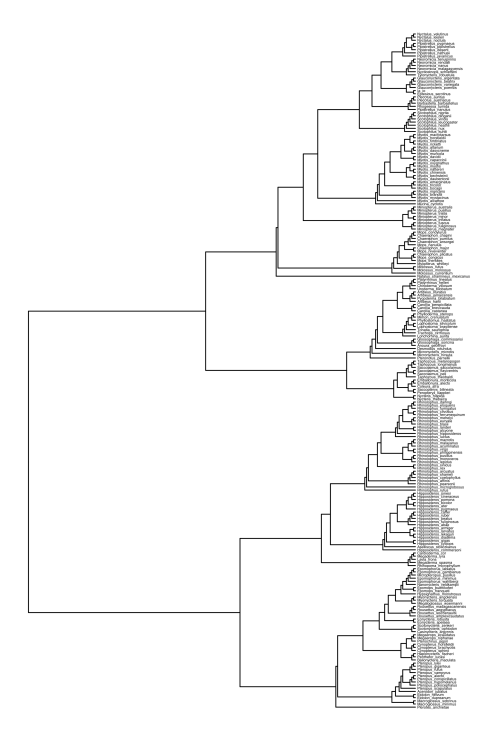
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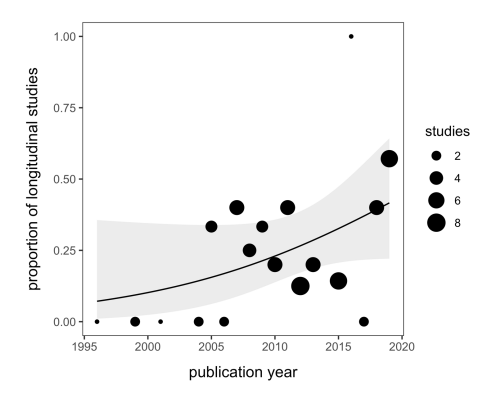
# S3. Bat phylogeny

Figure S2. Phylogeny of the 215 bat species included in the phylogenetic meta-analysis of filovirus and henipavirus prevalence and seroprevalence.

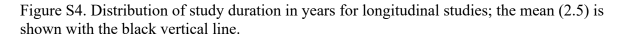


# S4. Temporal patterns in longitudinal sampling and reporting

Figure S3. Relationship between publication year and the proportion of studies reporting longitudinal data. Fitted values (line) and 95% confidence intervals (grey) from a generalized additive model (binomial errors) are shown with data scaled by the number of studies per year.



# **S5.** Distribution of sampling duration



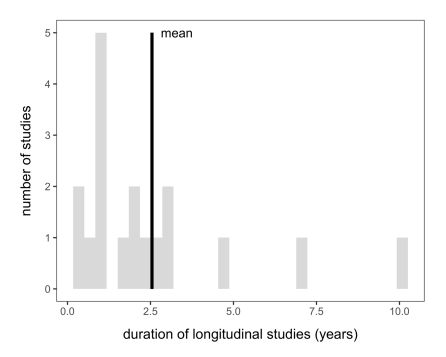
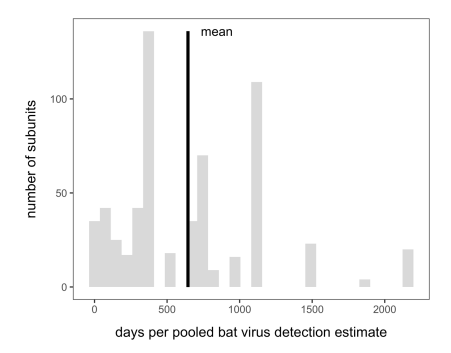


Figure S5. Distribution of days represented by each pooled viral detection estimate; the mean (644) is shown with the black vertical line.



# S6. REM results, $I^2$ , and $H^2$

Table S1. Results from the REM fit to all data and each data subset, using the random effects outlined in the main text. We show Cochran's Q, the associated p value, estimates of total  $I^2$  and the proportion of unknown variation not attributable to sampling variance for each random effect  $(I^2_{species}, I^2_{study}, \text{and } I^2_{subunit})$ , and phylogenetic heritability  $(H^2)$ .

Data	Q	р	I <sup>2</sup>	$I^{2}_{species}$	$I^{2}_{study}$	<b>l</b> <sup>2</sup> subunit	$H^2$
All data	6929	< 0.001	0.91	0.41	0.34	0.15	0.45
Filovirus prevalence	340	< 0.001	0.53	0.03	0.22	0.27	0.07
Filovirus seroprevalence	2337	< 0.001	0.85	0	0.66	0.2	0
Henipavirus prevalence	401	< 0.001	0.63	0.30	0.16	0.16	0.48
Henipavirus seroprevalence	1926	< 0.001	0.96	0.66	0.17	0.14	0.68

#### S7. Sampling design and reporting practices MEM

Table S2. ANOVA table from the MEM with a three-way interaction between sampling design and reporting practices, virus taxa, and detection method. For each term in the model, we provide Cochran's Q, the associated degrees of freedom, and the p value.

Model term	Q	df	р
Sampling and reporting practices	0.20	2	0.91
Virus taxa	0.09	1	0.77
Detection method	5.41	1	0.02
Sampling practices * virus taxa	0.46	2	0.79
Sampling practices * detection method	3.84	2	0.15
Virus taxa * detection method	0.03	1	0.87
Sampling practices * virus taxa * detection method	5.36	2	0.07

Table S3. Post-hoc analysis for the marginally significant interaction between sampling design and reporting practices, detection method, and virus taxa. MEMs with the same random effects were fit to each data subset (filovirus prevalence=193, filovirus seroprevalence=302, henipavirus prevalence=244, henipavirus seroprevalence=280). For each MEM, we provide the Q statistic and p value for sampling design and reporting practices.

Data subset	$Q_2$	р	
Filovirus prevalence	0.12	0.94	
Filovirus seroprevalence	10.30	0.006	
Henipavirus prevalence	2.03	0.36	
Henipavirus seroprevalence	1.35	0.51	

### **S8.** Spatiotemporal variation in longitudinal studies

Table S4. Signal of spatial variation within longitudinal studies. MEMs with the same random effects were fit to each data subset (filovirus prevalence=22, filovirus seroprevalence=73, henipavirus prevalence=88, henipavirus seroprevalence=90) with sampling location as a predictor. For each model, we provide the omnibus Q statistic and p value.

Data subset	Q	df	р
Filovirus prevalence	0.03	2	0.99
Filovirus seroprevalence	42.39	7	< 0.001
Henipavirus prevalence	23.96	10	0.008
Henipavirus seroprevalence	22.39	9	0.008

Table S5. Signal of temporal variation within longitudinal studies. MEMs with the same random effects were fit to each data subset (filovirus prevalence=22, filovirus seroprevalence=73, henipavirus prevalence=88, henipavirus seroprevalence=90) with month as a categorical predictor. For each model, we provide the omnibus Q statistic and p value.

Data subset	Q	df	р
Filovirus prevalence	1.44	6	0.96
Filovirus seroprevalence	23.42	11	0.02
Henipavirus prevalence	52.35	11	< 0.001
Henipavirus seroprevalence	13.12	11	0.29