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Vector species richness increases haemorrhagic disease prevalence through functional diversity modulating the duration of seasonal transmission

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SUMMARY

Although many parasites are transmitted between hosts by a suite of arthropod vectors, the impact of vector biodiversity on parasite transmission is poorly understood. Positive relationships between host infection prevalence and vector species richness (SR) may operate through multiple mechanisms, including (i) increased vector abundance, (ii) a sampling effect in which species of high vectorial capacity are more likely to occur in species-rich communities, and (iii) functional diversity whereby communities comprised species with distinct phenologies may extend the duration of seasonal transmission. Teasing such mechanisms apart is impeded by a lack of appropriate data, yet could highlight a neglected role for functional diversity in parasite transmission. We used statistical modelling of extensive host, vector and microparasite data to test the hypothesis that functional diversity leading to longer seasonal transmission explained variable levels of disease in a wildlife population. We additionally developed a simple transmission model to guide our expectation of how an increased transmission season translates to infection prevalence. Our study demonstrates that vector SR is associated with increased levels of disease reporting, but not via increases in vector abundance or via a sampling effect. Rather, the relationship operates by extending the length of seasonal transmission, in line with theoretical predictions.

Key words: Parasite transmission, vector-borne disease, mathematical model, functional diversity, disease-diversity relationship.

INTRODUCTION

Vector-borne diseases represent a large fraction of emerging and re-emerging infectious diseases (Jones *et al.* 2008), yet the role of the composition of vector communities (assemblages of distinct species) in influencing parasite transmission has received relatively little attention. Many infectious diseases are transmitted by a suite of vectors that are often closely related (e.g. congeners), and realized vector communities can vary in their species composition across a landscape. While species-level competencies, including transmissibility and longevity, can directly influence parasite transmission (Sardelis *et al.* 2001), there are additionally emergent properties of vector communities, particularly community abundance and activity period, that may promote transmission. If species have distinct phenologies, then there is the potential for a longer window of transmission.

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If instead, species have similar phenologies then communities have the potential to reach greater abundances, provided interspecific competition is not too strong. Studying the effects of community composition of vectors on wildlife diseases relies on the rare combination of spatially and temporally resolved data on the vector and disease.

Haemorrhagic disease (HD) is a viral disease vectored by biting midges in the genus *Culicoides* (Gibbs and Greiner, 1989). It predominantly affects white-tailed deer (*Odocoileus virginianus*) (Nettles and Stallknecht, 1992) and can lead to asymptomatic infection, morbidity (sickness) or mortality, with morbidity being by far the most common outcome in the Southeastern USA (Park *et al.* 2013), our study region. Disease occurrence in white-tailed deer in the USA is reported annually at the county level. Additionally, *Culicoides* vector surveillance has been conducted in the Southeastern USA during the late 2007–2012 (Vigil *et al.* 2014), involving over 5500 trap nights (number of traps \times number of nights) that have recorded approximately 50 species in the genus. We characterize the vector communities by their overall activity period and abundance, and these metrics are related to measures of reported disease in

host populations in the same locations. We find that no single species is driving patterns of disease. Additionally, while species richness (SR) is broadly positively associated with abundance, the estimated number of *Culicoides* individuals in a region is not predictive of levels of disease reporting. Rather, we find that communities composed of species with different phenologies (thereby generating a longer seasonal window of transmission) are associated with higher rates of disease reporting. These findings point to an important role for functional diversity of vectors in helping parasite populations invade and persist in host populations.

To lend explanatory and predictive power to the role of vector diversity in parasite transmission, we introduce a simple vector–host model where the vector population (which could be composed of one or several species) exhibits susceptible–infected (*SI*) dynamics and the host exhibits *SIS* dynamics (i.e. hosts recover back to the susceptible state); and where vectors spend a continuous fraction of the year (ϕ) in an active phase, including biting (i.e. transmission) and the remaining fraction ($1-\phi$) inactive (i.e. overwintering). The model allows for the fraction ϕ to vary, representing vector communities composed either of species with overlapping or distinct activity periods. This framework does not assume any differences in vector species regarding their ability to transmit the parasite. The model provides predictions for how levels of infection should increase in host populations as a function of vector community phenology and abundance.

STUDY SYSTEM

The vector-borne, HD is caused by serotypes of epizootic HD virus and bluetongue virus, closely related double-stranded RNA orbiviruses (Roy, 1996). In the USA, the disease is mainly associated with white-tailed deer (*O. virginianus*) (Nettles and Stallknecht, 1992). Transmission dynamics in the southeast USA are primarily associated with morbidity in hosts (Park *et al.* 2013).

Virus is transmitted between animals by biting midges in the genus *Culicoides* (Mellor *et al.* 2000). Of the approximately 150 species thought to occur in USA (Borkent and Grogan, 2009) about 50 species occur in the southeast. The relative role of each species in transmitting virus is unknown, though certain species have been shown to become experimentally infected (Smith *et al.* 1996). The Southeastern Cooperative Wildlife Disease Study (SCWDS) at the University of Georgia has been conducting surveys of *Culicoides* in the Southeastern USA since the late 2007 (Vigil *et al.* 2014). We use these data (covering full years 2008–2012) to determine vector species presence and absence in each light trap together with overall

Culicoides abundance. The light trap data are aggregated by site to estimate vector community composition and mean abundance. Following infection, host animals are infectious and then recover, after which they may become re-infected, especially by different virus strains (Stallknecht *et al.* 1991; Gaydos *et al.* 2002). Infected animals maintain relatively high viraemia for several weeks, meaning that activity periods of vectors need not be continuous to support transmission throughout a season, and year-round transmission may be possible in some locations.

We also make use a disease data set collated at the University of Georgia which records occurrence of white-tailed deer morbidity and mortality associated with HD at the county level from 1980–present (detailed in Park *et al.* 2013). Here, we restrict our analysis to 2008–2012, in line with data on the *Culicoides* species identified from light trap surveys. The disease data are smoothed spatially using two-dimensional kernel density estimation (kde2d function in R package MASS), generating a non-parametric estimate of the density of disease reports across the landscape. Values from this distribution are then extracted at the geographic coordinates of the *Culicoides* sampling sites as a proxy measure of prevalence. Independent deer density data collected by the state departments of natural resources (DNR; Supplemental material) are used to establish that deer density is not responsible for patterns of HD, consistent with theory on vector-borne diseases proposing that, in many systems, biting rates saturate at sufficiently high host densities (Keeling and Rohani, 2008).

Lastly, we use the *Culicoides* data to derive an estimate of the length of seasonal *Culicoides* activity period at each site. This is achieved by first calculating the weekly probability of detection of each *Culicoides* species (using all regional data). These profiles are approximately normally distributed. All weeks within 80% of the peak probability of detection are counted towards that species activity period (this is approximately 2/3 of the standard deviation). This method accounts for the fact that species themselves have different activity period durations, in addition to timing differences. The community activity period, ϕ , is then estimated as the proportion of weeks in the year when the community has at least one species active (Fig. A1, Supplemental material).

MODEL

To illustrate the expected relationship between vector abundance, length of seasonal transmission and host prevalence, we introduce a simple vector–host transmission model (Keeling and Rohani, 2008). The vector population exhibits *SI* dynamics and the host population exhibits *SIS* dynamics (i.e. hosts recover back to the susceptible state). In

addition, vectors spend a continuous fraction of the year (ϕ) in an active phase, including biting (i.e. transmission) and the remaining fraction ($1-\phi$) inactive (i.e. overwintering):

$$\frac{dS_v^*}{dt} = \lambda_v - \mu_v S_v^* - \frac{b\tau_{hv} S_v^* I_h}{S_h + I_h} - f_{\downarrow} S_v^* + f_{\uparrow} S_v^{\circ}$$

$$\frac{dI_v^*}{dt} = \frac{b\tau_{hv} S_v^* I_h}{S_h + I_h} - \mu_v I_v^* - f_{\downarrow} I_v^* + f_{\uparrow} I_v^{\circ}$$

$$\frac{dS_v^{\circ}}{dt} = f_{\downarrow} S_v^* - f_{\uparrow} S_v^{\circ} - \mu_v S_v^{\circ}$$

$$\frac{dI_v^{\circ}}{dt} = f_{\downarrow} I_v^* - f_{\uparrow} I_v^{\circ} - \mu_v I_v^{\circ}$$

$$\frac{dS_h}{dt} = \lambda_h - \mu_h S_h - \frac{b\tau_{vh} S_h I_v^*}{S_h + I_h} + \gamma I_h$$

$$\frac{dI_h}{dt} = \frac{b\tau_{vh} S_h I_v^*}{S_h + I_h} - \mu_h I_h - \gamma I_h$$

where closed (open) circles indicate active (inactive) vectors, and $f_{\uparrow}(f_{\downarrow})$ denotes the emergence (overwintering) rate. Dynamics of active and inactive vector states are determined by the f parameters:

$$f_{\uparrow} = \begin{cases} f_{\max} & \text{if } t \bmod 1 \leq \phi \\ 0 & \text{if } t \bmod 1 > \phi \end{cases} \quad \text{and} \\ f_{\downarrow} = \begin{cases} 0 & \text{if } t \bmod 1 \leq \phi \\ f_{\max} & \text{if } t \bmod 1 > \phi \end{cases}$$

with t measured in years (vector dynamics are illustrated in Fig. A2, Supplemental material). Parameter b is the biting rate, and τ_{vh} and τ_{hv} are transmission probabilities from host to vector and from vector to host, respectively. Demographics in the vector and host populations are captured via constant birth rates, λ_v and λ_h , and per capita mortality rates, μ_v and μ_h . Finally, parameter γ is the recovery rate of infected hosts back to the susceptible state.

For a vector population active year-round (i.e. $\phi = 1$), then the basic reproductive number (Anderson and May, 1992) for the system is easily calculated from the next generation matrix (Diekmann *et al.* 2013) (see Supplemental material):

$$R_0 = \sqrt{\frac{b^2 \tau_{vh} \tau_{hv} \widehat{S}_v}{(\mu_h + \gamma) \mu_v \widehat{S}_h}}$$

where \widehat{S}_x represent the disease-free equilibrium population sizes for vector ($x = v$) and host ($x = h$) with values determined by $\widehat{S}_x = \lambda_x / \mu_x$. For a vector population that spends a fraction of the year ($1-\phi$) inactive, the concept of a disease-free equilibrium is not defined. However, as a broad guide, we note that when vectors are active then R_0 is as defined,

and is equal to zero during inactive phases. Consequently, we see that per generation parasite fitness approaches its maximum value of R_0 as ϕ tends to 1. From this, we conclude that there are multiple facets of parasite fitness that can increase with vector SR; aside from adding competent vectors to a community (increasing biting rates, transmissibilities and longevity), R_0 can also be increased by increasing vector abundance (\widehat{S}_v) and by adding species with distinct phenologies (increasing ϕ). These latter two factors may not necessarily increase independently. In the case of overlapping active phases, competition between vector species may limit their overall abundance.

RESULTS

A series of statistical models were constructed and analysed to determine which aspects of *Culicoides* were predictive of HD reporting, i.e. the disease density estimates obtained by kernel smoothing (HDdens). *Culicoides* SR is a positive predictor of HD density estimates (HDdens, which is log-transformed to meet assumptions of normality) (linear model: $\ln(\text{HDdens}) \sim \text{SR}$, slope = 0.16, $P < 2 \times 10^{-16}$, $R^2 = 0.3$). However, SR needs to be interpreted biologically. In proposing a vector community-level explanation for disease patterns in data (i.e. length of the community-generated seasonal transmission window), it is important to consider and rule out other ways in which SR may lead to increased disease reporting. We consider three key alternative hypotheses: (i) sampling effect (Leps *et al.* 2006), where species-rich communities are more likely to contain functionally important species by chance, in this case species with high vectorial capacity; (ii) community abundance, where SR is simply leading to increased overall abundance; (iii) latitudinal effects, where species at lower latitudes exhibit relatively short overwintering periods.

Across all the sites where *Culicoides* were sampled ($n = 207$) there is considerable variation in disease reporting (measured by the kernel density estimate at each site). No single vector species strongly influences the level of disease reporting across sites. This was established by comparing the fit of linear models ($\ln(\text{HDdens}) \sim \text{SR}$) in which each species was left out in turn. The difference in model R^2 does not point to an obvious keystone species (difference in R^2 between adjacent models ordered by R^2 : min = 0.0, mean = 0.0027, max = 0.027). Additionally, average abundance of *Culicoides* at sites did not explain the disease kernel density estimate (linear model: $\ln(\text{HDdens}) \sim \text{mean}(\text{abundance})$, $P = 0.55$, $R^2 = 0.0018$), although *Culicoides* abundance is related to SR in a saturating, non-linear fashion (Fig. A3, Supplemental material; Spearman rank correlation: $\rho = 0.56$, $P < 2.2 \times 10^{-16}$).

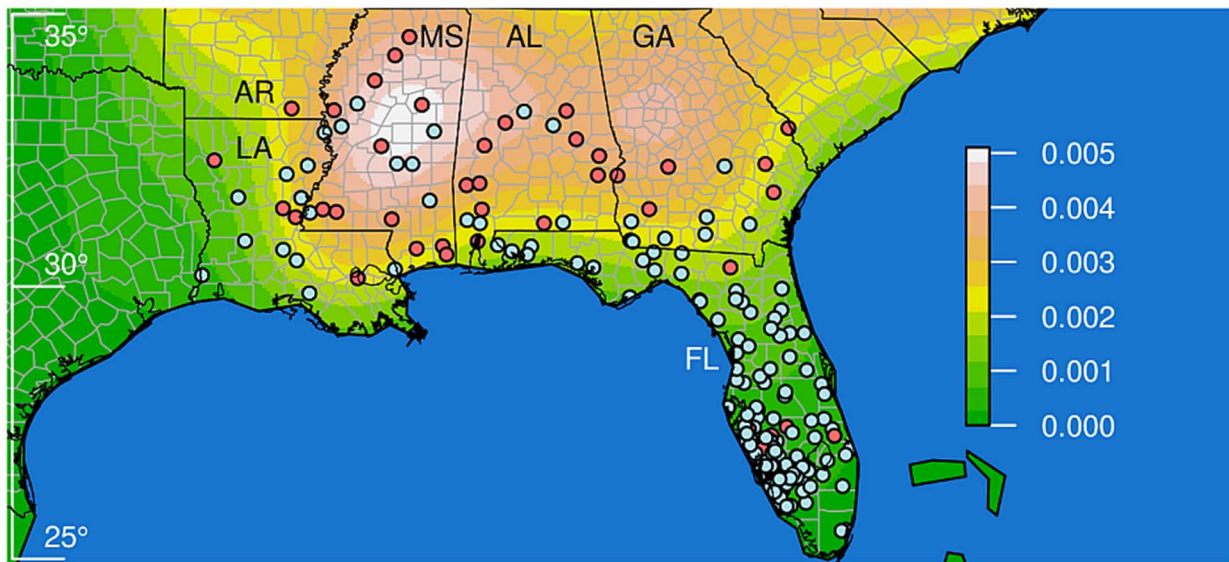


Fig. 1. Kernel density estimate of HD based on occurrence data in the Southeast USA (colourbar: highest density of records in white, lowest in green). *Culicoides* sampling sites are overlaid in their respective positions in the states of Arkansas (AR), Louisiana (LA), Mississippi (MS), Alabama (AL), Georgia (GA) and Florida (FL). Sampling sites are coloured according to their ranked position of estimated length of annual *Culicoides* community activity period, ϕ (red = highest 25%, cyan = remaining 75%). Longitude is indicated on the left-hand vertical axis.

Simple mapping reveals that disease density is heterogeneous in space with localized high density in Mississippi, continuing into Alabama and Georgia (Fig. 1). *Culicoides* sampling sites are overlaid on this map and coloured according to the estimated length of the seasonal transmission window, ϕ (red = longest 25% of transmission windows, which corresponds to $\phi > 0.5$). We acknowledge that space itself is naturally a strong predictor of such a disease density pattern (e.g. tensor product smoothed generalized additive model: $\ln(\text{HDdens}) \sim (\text{latitude}, \text{longitude})$, family = Gaussian, $P < 2 \times 10^{-16}$, $R^2 = 0.98$). However, evidence suggests that the interpretation of the importance of space is in terms of the community-generated length of transmission window (ϕ) and not latitude.

Latitudinal stratification confirms that several species ($n = 11$) have a different temporal activity pattern in the north relative to the south (with 30° latitude demarcating north from south; Fig. A4, Kolmogorov–Smirnov tests on species-level detection week in north and south groups). The difference is generally characterized by both later emergence and earlier overwintering in the north. For the other species there was either no significant difference in the temporal activity ($n = 13$) or the species occurred exclusively in one of the two regions ($n = 28$). While latitude is predictive of disease density (linear model: $\ln(\text{HDdens}) \sim \text{latitude}$, slope = 0.42, $P < 2.2 \times 10^{-16}$, $R^2 = 0.95$), it is in the opposite direction than would be expected based on species-level extended activity (since disease density is shown to be higher at high latitudes where species-level activity is relatively short, Fig. 1).

However, the community-level activity period (ϕ) is positively predictive of disease density (Fig. 1; linear model: $\ln(\text{HDdens}) \sim \phi$, slope = 3.7, $P = 2.5 \times 10^{-15}$, $R^2 = 0.27$). Each *Culicoides* species' presence and absence at sites results in spatially distinct *Culicoides* communities. Some of these communities have a relatively short window of transmission (small ϕ), typically due to a high degree of overlap in phenology of member species. Conversely other communities have long windows of transmission (large ϕ) for the complementary reasons. The range of ϕ (min = 0.19, mean = 0.42, max = 0.69) means some vector communities are active for ~ 2 months while others are active for > 8 months. The community-level window of transmission accounts for nearly 30% of variation in the disease reporting metric. In addition, it overcomes any species level reductions in activity period in relatively northern latitudes, which contain the disease hotspots. By calculating ϕ based on time periods when each species is within 80% of its peak probability of detection (Fig. A1, Supplemental material), we aim to conservatively estimate region-wide activity, rather than capture early emergence and late overwintering in southern latitudes (as could occur with a threshold $\ll 80\%$).

The vector–host transmission model predicts that equilibrium prevalence in the host population will increase with the length of the active vector season, ϕ (Fig. 2). In addition, high abundance of the vector population is also predicted to positively influence host equilibrium prevalence. As prevalence is bounded, it tends to saturate as both factors increase, and in much of the ϕ -parameter space, increases in community abundance exhibit

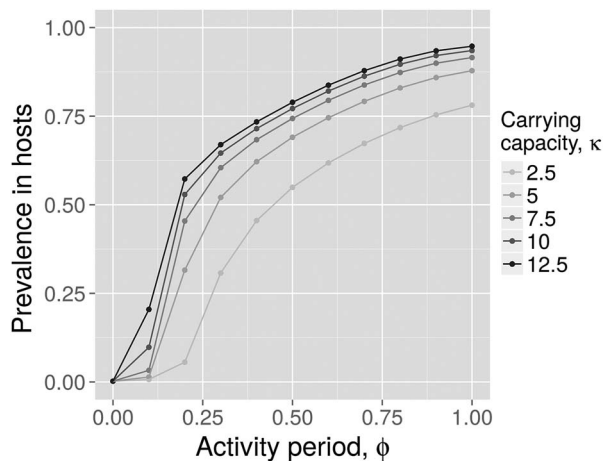


Fig. 2. Equilibrium prevalence in host populations (y -axis, proportion of hosts infected) as a function of the fraction of the year that vectors are active (x -axis, ϕ) and vector abundance (vector-carrying capacity, κ , grey-scale coded). Model parameters are $\lambda_v = k\mu_v$, $\mu_v = 2.0$, $f_{\max} = 100.0$, $\lambda_h = 1.0$, $\mu_h = 0.2$, $b = 10.0$, $\tau_{vh} = \tau_{hv} = 1.0$, $\gamma = 1.0$, $\phi = 0.0$ – 1.0 , $\kappa = 2.5$ – 12.5 .

strong diminishing returns on infection prevalence in host populations.

DISCUSSION

We have introduced a study system in which many species of vectors from the same genus can transmit virus to hosts, and where the competency of the vector species is largely unknown. Communities comprising species with distinct activity periods (over richness *per se* and over abundance) are spatially coincident with relatively high levels of disease reporting. This is a community-level effect since empirical and modelling results do not require any differences in the vector species in terms of their competency; just that they have distinct activity periods thereby generating longer transmission windows. Similar mechanisms have been put forward to explain the epidemiology of West Nile virus using a two vector species approach (Lord, 2010). However, the role of vectors is largely absent from discussions on the disease-diversity relationship, in spite of the large impacts that vector diversity can have on parasite transmission.

While our approach is neutral with respect to assumptions about the ability of a given vector species to transmit virus, more broadly, community-driven seasonality in transmission may interact with species-level variation in vector competence creating temporal hotspots for parasite transmission. Such mechanisms have the potential to operate broadly in human and animal infectious disease systems since ecological sensitivity of vectors means that community composition is likely to turn over along environmental gradients influencing parasite transmission (To e *et al.* 1997; Lambin *et al.*

2010). Community abundance and the duration of seasonal transmission, in addition to vectorial capacity (Dye, 1986), can lead to both positive and negative effects of vector SR on parasite fitness.

Awareness of the importance of functional diversity above SR is increasing (D az and Cabido, 2001; Petchey and Gaston, 2002). By far the most active areas of research concern plant communities, ecosystem services and potential responses to climate change (Tilman, 1997; Sala *et al.* 2000; Cadotte *et al.* 2011; Craine *et al.* 2012). However, we have demonstrated that parasite transmission may also be promoted or inhibited by functional diversity, especially through the temporal activity of vector species in a community.

As in many studies of functional diversity, it is important to consider the interplay between SR and abundance. Temporal niche partitioning may facilitate the presence of multiple species since it alleviates interspecific competition (Chesson, 2000). Consequently, communities with longer transmission windows than those attained by any one vector species may be common. However, an increase in seasonal transmission due to SR may trade off against vector abundance, a factor that can also increase parasite transmission; communities comprising species with similar activity periods may only attain higher abundance provided interspecific competition is not too strong, as we demonstrated with our mathematical model.

The quality of the data associated with this wildlife disease have shed light on vector-borne parasites, and suggest that similar inquiries could be made in parasites of human health concern, where data collection is typically more extensive. We contend that vector species diversity should form part of the disease-diversity discussion, and recognize novel ways in which vector SR may manifest functionally, in addition to classic components of competency.

SUPPLEMENTARY MATERIAL

To view supplementary material for this article, please visit <https://doi.org/S0031182015000578>.

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REFERENCES

- Anderson, R. M. and May, R. M.** (1992). *Infectious Diseases of Humans: Dynamics and Control*. Oxford University Press, Oxford.
- Borkent, A. and Grogan, W. L.** (2009). Catalog of the New World biting midges north of Mexico (Diptera: Ceratopogonidae). *Zootaxa* **2273**, 1–48.
- Cadotte, M. W., Carscadden, K. and Mirotchnick, N.** (2011). Beyond species: functional diversity and the maintenance of ecological processes and services. *Journal of Applied Ecology* **48**, 1079–1087.
- Chesson, P.** (2000). Mechanisms of maintenance of species diversity. *Annual Review of Ecology and Systematics* **31**, 343–366.
- Craine, J. M., Ocheltree, T. W., Nippert, J. B., Towne, E. G., Skibbe, A. M., Kembel, S. W. and Fargione, J. E.** (2012). Global diversity of drought tolerance and grassland climate-change resilience. *Nature Climate Change* **3**, 63–67.
- Diekmann, O., Heesterbeek, H. and Britton, T.** (2013). *Mathematical Tools for Understanding Infectious Disease Dynamics*. Princeton University Press, Princeton, NJ.
- Diaz, S. and Cabido, M.** (2001). Vive la différence: plant functional diversity matters to ecosystem processes. *Trends in Ecology & Evolution* **16**, 646–655.
- Dye, C.** (1986). Vectorial capacity: must we measure all its components? *Parasitology Today* **2**, 203–209.
- Gaydos, J. K., Davidson, W. R., Elvinger, F., Howerth, E. W., Murphy, M. and Stallknecht, D. E.** (2002). Cross-protection between epizootic hemorrhagic disease virus serotypes 1 and 2 in white-tailed deer. *Journal of Wildlife Diseases* **38**, 720–728.
- Gibbs, E. P. J. and Greiner, E. C.** (1989). Bluetongue and epizootic hemorrhagic disease. *The Arboviruses: Epidemiology and Ecology* **2**, 30–70.
- Jones, K. E., Patel, N. G., Levy, M. A., Storeygard, A., Balk, D., Gittleman, J. L. and Daszak, P.** (2008). Global trends in emerging infectious diseases. *Nature* **451**, 990–993.
- Keeling, M. J. and Rohani, P.** (2008). *Modeling Infectious Diseases in Humans and Animals*. Princeton University Press, Princeton, NJ.
- Lambin, E. F., Tran, A., Vanwambeke, S. O., Linard, C. and Soti, V.** (2010). Pathogenic landscapes: interactions between land, people, disease vectors, and their animal hosts. *International Journal of Health Geographics* **9**, 54.
- Leps, J., De Bello, F., Lavorel, S. and Berman, S.** (2006). Quantifying and interpreting functional diversity of natural communities: practical considerations matter. *Preslia* **78**, 481–501.
- Lord, C. C.** (2010). The effect of multiple vectors on arbovirus transmission. *Israel Journal of Ecology & Evolution* **56**, 371–392.
- Mellor, P. S., Boorman, J. and Baylis, E.** (2000). Culicoides biting midges: their role as arbovirus vectors. *Annual Review of Entomology* **45**, 307–340.
- Nettles, V. F. and Stallknecht, D. E.** (1992). History and progress in the study of hemorrhagic disease of deer. Transactions of the North American Wildlife and Natural Resources Conference **57**, 499–516.
- Park, A. W., Magori, K., White, B. A. and Stallknecht, D. E.** (2013). When more transmission equals less disease: reconciling the disconnect between disease hotspots and parasite transmission. *PLoS ONE* **8**, e61501.
- Petchey, O. L. and Gaston, K. J.** (2002). Functional diversity (FD), species richness and community composition. *Ecology Letters* **5**, 402–411.
- Roy, P.** (1996). Orbiviruses and their Replication. In *Fields Virology* (ed. Fields, B.N.), pp. 1709–1731. Lippincott-Raven, Philadelphia, PA.
- Sala, O., Chapin, F., Armesto, J., Berlow, E., Bloomfield, J., Dirzo, R., Huber-Sanwald, E., Huenneke, L., Jackson, R., Kinzig, A., Leemans, R., Lodge, D., Mooney, H., Oesterheld, M., Poff, N., Sykes, M., Walker, B., Walker, M. and Wall, D.** (2000). Global biodiversity scenarios for the year 2100. *Science* **287**, 1770–1774.
- Sardelis, M. R., Turell, M. J., Dohm, D. J. and O'Guinn, M. L.** (2001). Vector competence of selected North American Culex and Coquillettia mosquitoes for West Nile virus. *Emerging Infectious Diseases* **7**, 1018–1022.
- Smith, K. E., Stallknecht, D. E. and Nettles, V. F.** (1996). Experimental infection of Culicoides lahillei (Diptera: Ceratopogonidae) with epizootic hemorrhagic disease virus Serotype 2 (Orhivirus: Reoviridae). *Journal of Medical Entomology* **33**, 117–122.
- Stallknecht, D., Kellogg, M., Blue, J. and Pearson, J.** (1991). Antibodies to bluetongue and epizootic hemorrhagic disease viruses in a barrier island white-tailed deer population. *Journal of Wildlife Diseases* **27**, 668–674.
- Tilman, D.** (1997). The influence of functional diversity and composition on ecosystem processes. *Science* **277**, 1300–1302.
- Toé, L., Tang, J., Back, C., Katholi, C. R. and Unnasch, T. R.** (1997). Vector–parasite transmission complexes for onchocerciasis in West Africa. *Lancet* **349**, 163–166.
- Vigil, S. L., Grogan, W. L., Wlodowski, J. C., Parris, J., Edwards de Vargas, S., Shaw, D., Cleveland, C. and Corn, J. L.** (2014). New records of biting midges of the genus *Culicoides latreille* from the Southeastern United States (Diptera: Ceratopogonidae). *Insecta Mundi* **394**, 1–14.