

1 **Making sense of the virome in light of evolution and ecology**

2

3 Keywords:

4 *Virome, viruses, co-phylogenetic mixed model, virus discovery, virus metagenomics, virus*
5 *ecology, virus evolution, virus communities, virus prevalence*

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27

28 Abstract

29

30 Understanding the patterns and drivers of viral prevalence is of key importance for
31 understanding pathogen emergence. Over the last decade, metagenomic sequencing has
32 exponentially expanded our knowledge of the diversity and evolution of viruses associated
33 with all domains of life. However, as most of these 'virome' studies are primarily descriptive,
34 our understanding of the predictors of virus prevalence and diversity, and their variation in
35 space and time, remains limited. For example, we do not yet understand the relative
36 importance of ecological predictors (e.g., seasonality, habitat) versus evolutionary predictors
37 (e.g., host and virus phylogenies) in driving virus prevalence and diversity. Few studies are
38 set up to determine what factors predict the composition of the virome of individual hosts,
39 populations, or species. In addition, most studies of virus ecology represent a snapshot of
40 single viromes at a single point in time and space. Fortunately, recent studies have begun to
41 use metagenomic data to directly test hypotheses about the evolutionary and ecological
42 factors which drive virus prevalence, sharing and diversity. By synthesising evidence across
43 studies, we present some over-arching ecological and evolutionary patterns in virome
44 composition, and illustrate the need for further work to quantify the drivers of virus
45 prevalence and diversity.

46 1. Introduction

47 Viruses are ubiquitous across life on earth, but we have much to learn about what
48 determines the abundance and distribution of viruses (i.e. the “virome” or “virosphere”)
49 across hosts and ecosystems. Large scale metagenomic sequencing projects have
50 expanded our knowledge of the diversity and composition of eukaryotic viromes [1], with the
51 number of published viral metagenomics papers increasing more than six-fold in the last
52 decade, and the number of classified virus species increasing more than three-fold (5542
53 released virus RefSeq genomes on NCBI in September 2014 versus 18,668 in September
54 2024). This number will increase dramatically following the implementation of discovery
55 models that utilize protein structure as well as sequence data, with a single recent study
56 using an AI-based approach identifying >160,000 novel virus species [2]. Consequently, the
57 rate of virus discovery is greatly out-pacing virus classification. Despite this revolution in
58 virus discovery, the field is only just beginning to move from being purely descriptive
59 “molecular natural history” to being hypothesis driven.

60

61 Over the last decade the metagenomic sequencing of animal, plant and soil-associated
62 bacterial communities – often referred to as microbiome research – has transitioned from a
63 descriptive state toward directed hypothesis testing (see reviews [3,4]). Continuous
64 monitoring of wild populations has allowed the analysis of long-term data sets to study the
65 determinants and fine-scale variation in microbiomes. Examples include global variation in
66 amphibian skin bacterial communities linked to climate [5], variation in the bacterial
67 microbiomes of birds linked to foraging behaviour [6], and seasonality in gut parasite
68 communities [7] and bacterial microbiotas [8] in mammals. In contrast, most virus-focused
69 metagenomic studies can only be interpreted as a single snapshot of a current individuals’,
70 populations’, species’ or environments’ virome (or microbiome) at that point in time and
71 space (e.g. [3]). Testing explicit ecological and evolutionary hypotheses on the causes and
72 consequences of variation in the virome requires that we i) integrate extensive spatial and

73 longitudinal virome sampling alongside ecological data; and ii) embed the virosphere in a
74 whole community context, by considering viruses not only as potential zoonotic diseases, but
75 as participants in their wider ecosystems. Collectively, this will allow us to determine their
76 importance in maintaining whole ecosystem functionality and stability [9]. Addressing this
77 knowledge gap is currently hampered by biases in the metagenomic literature, which could
78 lead us to overstate broad-scale patterns or drivers of virome diversity [10]. For example,
79 large databases of host-virus associations (e.g. [11,12]) are biased towards mammalian
80 viruses, and groups such as bats with high research interest. Such biases can lead to
81 dogmas in the literature, for example that more zoonotic diseases originate from particular
82 host clades because of their inherent traits or ecology, when in fact, we have simply
83 identified a larger number of the viruses infecting them [12]. As a result, compilation of
84 databases is urgently needed for less well sampled groups, as currently being attempted in
85 insects [13].

86

87 Studying the evolution and ecology of viromes presents some unique challenges. However,
88 revealing the determinants of and barriers to successful cross-species virus transmission is
89 crucial to understanding the potential of a virus to emerge in a new species. For example,
90 host ecology and behaviour affect contact rates among hosts, with more frequent contacts
91 increasing the potential for virus diversification and spread [14]. Likewise, closely related
92 hosts offer a more similar physiological and cellular environment for a virus, and may have
93 similar niches, habitats or diets, leading to more similar viromes [15]. Identifying the factors
94 that promote virus diversification and spread involve taking an whole ecosystem (i.e., One
95 Health) approach [16] and so have a broad implications for public and agricultural health. For
96 example, the evolutionary and ecological factors that structure species viromes influence
97 disease emergence in wildlife [17–20], pollinators [21,22], and livestock [23,24], and have
98 clear connections to spill-over into humans. Herein, we summarise current knowledge of the

99 ecological and evolutionary factors determining virome composition and propose how we
100 can expand this with future research.

101 2. Evolutionary factors driving the composition of species viromes

102

103 Vertebrate and invertebrate viromes remain largely distinct, despite their frequent
104 interactions through predation, parasitism and shared environments [25,26]. This underlines
105 that viruses, like bacteria and fungi [27], are preferentially shared between closely related
106 hosts [28]. After accounting for shared environment and ecological interactions, traits shared
107 between phylogenetically closely related species will shape the composition of the virome.

108 These traits, such as host receptors, physiology and immunity, present a similar environment
109 for a virus and are the result of the history of selection on hosts (and viruses), in part caused
110 by their exposure history [29]. The importance of host relatedness does not necessarily
111 present as a linear relationship between susceptibility and host phylogenetic distance.

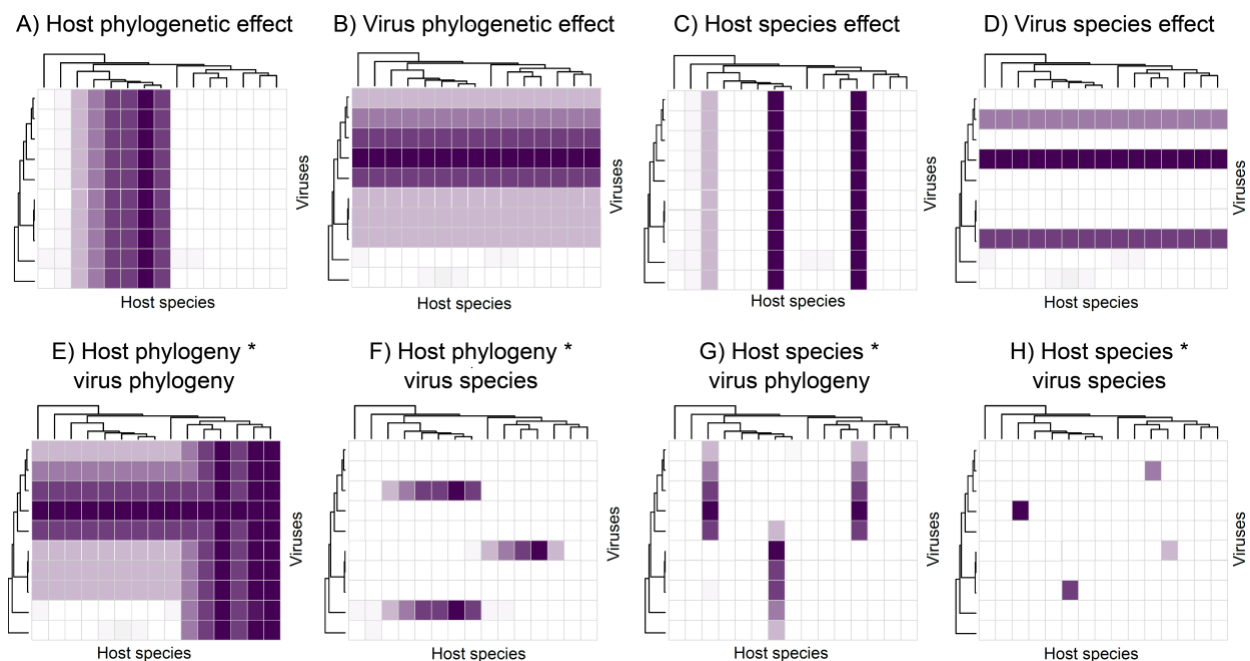
112 Closely related hosts may have similar levels of susceptibility to a given virus (or group of
113 viruses), independent of their distance from the viruses 'natural' host. This clade effect can
114 result in viruses being clustered in a patchwork of clades on the host phylogeny [15,30,31].

115

116 This concept also holds true for surveys of viral presence/absence in natural populations.

117 Host phylogenetic relatedness is a significant predictor of the likelihood of viral sharing
118 between primates [32]. Additionally, rabies virus sequence data, sampled from single viruses
119 across multiple bat host species, demonstrates that cross-species transmission events and
120 successful host shifts are more likely in closely related host species [17,33]. Importantly, in
121 this system, range overlap is less important than phylogenetic relatedness in predicting
122 sustained host shifts compared to spillover events (although current estimates of geographic
123 range are used to test this, rather than historical range). Additionally, host phylogenetic
124 effects have been demonstrated for a number of individual viruses in a range of hosts both
125 experimentally [34–37] and in nature [31].

126 Large databases of host-virus associations have also been used to show an increased
 127 proportion of zoonotic viruses in species that are closely related to humans [38], and that
 128 species-rich host taxonomic groups harbour more viruses [12]. This again supports that idea
 129 that viruses can preferentially jump between closely related host species. These databases
 130 have also been used to show that some virus lineages have a greater propensity to change
 131 hosts [39] and that viruses with broad host ranges have a greater propensity to jump host
 132 [40]. However, a key caveat is that these analyses are based on our current incomplete
 133 understanding of global viral diversity [10]. There is also some evidence that host species
 134 may vary in their overall susceptibility to viral infection, or cross-species transmission
 135 [18,34]. However, at least among mammalian viruses, there is no evidence that particular
 136 host taxonomic groups are inherently more likely to be virus reservoirs because of host
 137 traits. On the contrary, host taxonomic orders with greater species richness simply appear to
 138 harbour more diverse viromes, and are therefore more often the source of cross species
 139 transmission events [12,41].



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141 **Figure 1. Host and virus species level and phylogenetic effects on virus prevalence**
 142 **and viral host range.** The y axis represents a hypothetical virus phylogeny, and the x axis a
 143 hypothetical host phylogeny. Asterisk (*) indicates model interaction terms. Each panel
 144 represents different possible scenarios. A: The incidence and prevalence of viruses across

145 host species is predictable by host phylogeny (i.e. closely related host species have a similar
146 incidence of viruses). B: The incidence and prevalence of viruses across host species is
147 predictable by virus phylogeny (i.e. closely related viruses have a similar infectivity across
148 host species). C: Certain host species are inherently more or less susceptible to viruses, in a
149 way not predictable by the host phylogeny (i.e. due to ecological or physiological traits). D:
150 Certain viruses are particularly infectious, or not, irrespective of host species, in a way not
151 predictable by virus phylogeny. E: Related hosts have similar incidences of clades of related
152 viruses (i.e. virus incidence and prevalence is predictable by both host and virus
153 phylogenies). F: Related hosts have similar incidences of some viruses, but not all, and not
154 in a phylogenetically predictable manner. G: Related viruses show similar infectivity to only
155 some host species – not all – and not in a way predicted by host phylogeny. H: Host
156 susceptibility depends on specific host x virus interactions not predictable by either host or
157 virus phylogeny. Based on [42,43].

158

159 To address the roles of host and viruses relatedness requires analysis of many related hosts
160 and viruses. The evolutionary drivers of virome composition can be broken down into a
161 series of ‘species level’ and ‘phylogenetic’ effects (Figure 1) [42,43]. Host species effects
162 and phylogenetic effects capture how hosts vary in their overall prevalence of viral infection,
163 and whether related hosts tend to have similar overall viral prevalence for the host (Fig
164 1A/C). Virus species effects and phylogenetic effects capture how viruses vary in the overall
165 size of their host range, and whether related viruses have similar host ranges (Fig 1B/D). By
166 examining these effects, it is possible to ask whether some hosts are more susceptible than
167 others, whether some viruses are more generalist than others, and if these traits are similar
168 among related hosts or viruses. In addition, hosts may vary systematically in the composition
169 of their virome, and viruses may vary systematically in the composition of their host range.
170 For example, it is now well established that viruses generally transmit more easily between
171 more closely related host species [35] and that host-virus co-divergence also occurs [44],
172 although less commonly in many groups [45]. Importantly, both of these processes mean
173 that related hosts (or viruses) will have more similar viromes (or host ranges) [46,47],
174 sometimes referred to as phyllosymbiosis [48]. Moreover, we expect related hosts (or related
175 viruses) to be more similar in their virome composition (or host range). We can examine

176 these questions by looking at interactions between the terms described above (Figure 1)
177 [42].

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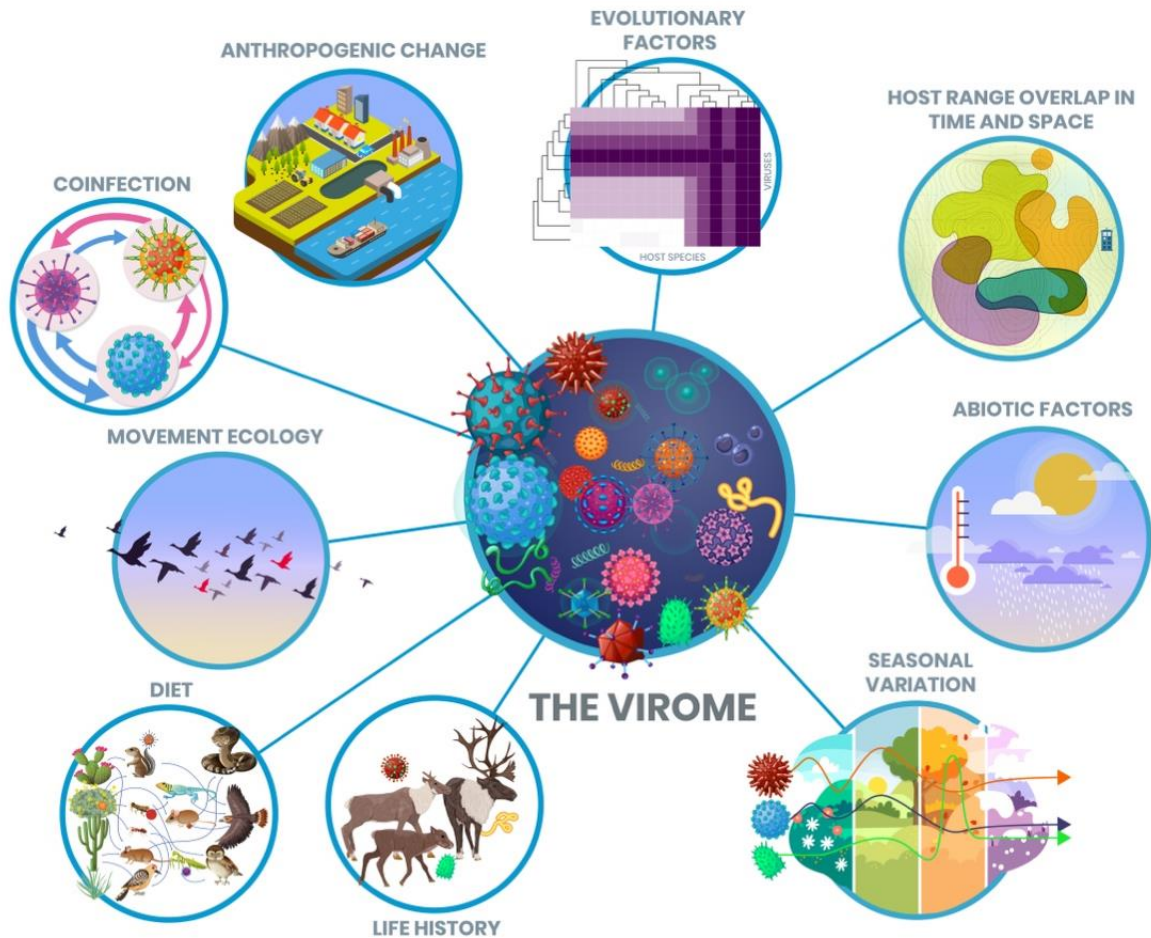
179 Interactions between host and virus species-level effects correspond to unique species-by-
180 species interactions in susceptibility or resistance that are not predictable from the relatives
181 of either the host or virus (Fig 1H). An interaction between host phylogeny and virus species
182 corresponds to an individual virus being a specialist on (or limited to) specific clades of the
183 host (Fig 1F); and an interaction between virus phylogeny and host species corresponds to
184 specific clades of viruses showing similar infectivity on a specific host (Fig 1G). The
185 phylogenetic interaction term corresponds to particular clades of the host being more prone
186 to infection by particular clades of the virus—as predicted by co-divergence or near-
187 neighbour preferential host-switching models (Fig 1E) [49].

188

189 Recent metagenomic sequencing studies are beginning to generate data that can address
190 these questions, confirming for example, that the host phylogeny predicts a significant
191 proportion of variance in the structure of virus communities. Studies of marine fish have
192 found that their viromes are predominantly shaped by the phylogenetic history of their hosts
193 [50], influencing both alpha and beta virome diversity [51]. Likewise, host taxonomy in birds
194 is important in explaining differences in virus community structure [52]. Additionally, host
195 order explains significant variation in the viral richness and abundance in wild bats, rodents
196 and shrews [18]. Likewise, viral richness in species sampled across an entire island
197 ecosystem clusters by host taxonomy, with viral order explaining the most variation in virus
198 community composition [25]. These studies also demonstrate how we can simultaneously
199 quantify the relative importance of both phylogeny and ecology in determining virome
200 composition and diversity [see also 53].

201

202 However, these studies fit taxonomic groups as categorical/random effects in models, rather
203 than the effect of the host phylogeny directly. A more sophisticated – although data intensive
204 – approach is to simultaneously fit species level and phylogenetic (relatedness) effects. For
205 example, in a study of 13 bumblebee species and 20 viruses approximately a quarter of the
206 variation in virus prevalence was explained by terms accounting for the evolutionary histories
207 of the hosts and viruses [43]. However, individually each of the host and virus effects
208 explained only a small proportion of the variance in prevalence with large amounts of
209 uncertainty around these estimates, which may reflect a lack of power to detect such effects
210 on a relatively small number of hosts and viruses. Indeed, even when sampling a larger
211 number of hosts and viruses, the best-fit models of the predictors of viral richness and
212 abundance in wild rodents, bats, and shrews explained less than 40% of deviance,
213 highlighting the challenges in accurately explaining the patterns of viral diversity and
214 abundance across species [18].



215

216 **Figure 2. Illustration of the ecological and evolutionary drivers of viromes.** Factors
 217 influencing the virome may interact, for example; seasonal changes in host range may
 218 coincide with seasonal peaks in infection burden, with coinfection interactions shifting
 219 components of the virome.

220 *3. Ecological drivers of virome composition*

221

222 Host ecological traits have a major impact on the composition and diversity of viromes,
 223 operating primarily through influencing the likelihood of exposure at multiple scales and
 224 interacting with evolutionary factors (Figure 2). First, differences in a host's distribution in
 225 time and space (phenology and geographical range) affect the likelihood of exposure and
 226 virus sharing. A number of studies have revealed a positive relationship between host
 227 geographic range overlap and the likelihood of viral sharing, cross species transmission, and

228 viral richness [11,17,38,54]. Second, within communities of sympatric organisms, biotic
229 factors limit exposure between host individuals through food webs or trophic networks,
230 dietary preferences, age structures, and predator-prey networks. At both scales,
231 anthropogenic driven climate and land-use change will alter host dynamics, with knock-on
232 effects on virome composition and diversity. Within-host ecological interactions can also
233 modulate the likelihood of virus acquisition [55], for example through co-infection with other
234 viruses or non-viral pathogens, or through interactions with the resident microbiota. These
235 interactions can alter infection outcomes and onward spread, and therefore larger population
236 level virus diversity or abundance [56].

237

238 *3.1. Abiotic associations with virome diversity and abundance*

239

240 Key abiotic factors such as temperature, humidity, and rainfall, all drive virus prevalence in
241 single virus studies by modulating host population behaviour or viral
242 transmission/environmental persistence [57]. We might therefore expect that virus
243 prevalence and diversity will follow similar trends to those seen in other microbes, exhibiting
244 broad-scale elevational/depth and latitudinal gradients, with these abiotic factors driving
245 changes in virome diversity and composition.

246

247 Ocean temperature modulates the abundance and composition of both marine
248 bacteriophage communities [58] and the viral communities of fish [51]. In terrestrial
249 organisms, increases in elevation are associated with a decline in viral richness in vampire
250 bats, with colonies at lower elevations in the Amazon rainforest having higher viral richness
251 and distinct community composition [59]. Given that host species richness generally
252 increases towards the equator via the latitudinal diversity gradient, latitude (as well as
253 longitude) has been identified as a modulator of virus communities, acting as a proxy for
254 both the biotic and abiotic variables described above. For example, marine virus diversity

255 showed higher diversity at lower latitudes, with decreasing virus diversity moving poleward,
256 mirroring that of most aquatic and terrestrial host diversity patterns [60]. In addition,
257 longitude was a significant factor in explaining virus diversity in bats [59], while both latitude
258 and longitude had a very strong impact on the human gut virome even when accounting for
259 ethnicity and other demographic factors [61]. However, in contrast to these clear latitudinal
260 and longitudinal trends, the viruses infecting fish species and individuals in Antarctica are
261 just as diverse and abundant as those from warmer marine environments [62], despite the
262 host diversity gradient. It may be the case that our relative lack of knowledge on virus
263 diversity in many groups is obscuring caveats to the assumption that virome diversity
264 increases with host diversity. For example, phylogenetic rarity (the phylogenetic distance
265 between species in a community) may be more important in determining virome diversity,
266 and temperate areas may facilitate larger aggregations of species, increasing contact rates,
267 and the transmission of viruses [63].

268

269 4.1.2 Seasonal variation in viromes

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272 If temperature, humidity and rainfall can drive species viral diversity and composition, then
273 viromes will vary seasonally, particularly in temperate regions. Indeed, evidence from single-
274 virus studies suggests that we should see seasonal trends in virus prevalence [64],
275 particularly with respiratory viruses, with the highest prevalence in autumn months [65].
276 From the few virome studies which have addressed seasonality, it appears that viral
277 abundance, evenness and richness also display seasonal trends [66]. Additionally, in
278 surveys of wastewater, viral alpha and beta diversity varies significantly by season [67].
279 However, in one study of seasonality in the *Picornaviridae* component of rodent viromes,
280 evenness appears to peak in spring/summer, pre-dating peaks in virus prevalence seen in
281 autumn [68]. Indeed, the intrinsic link between seasonal trends and abiotic factors such as
282 temperature, humidity and rainfall, and biotic factors such as host immune response or

283 behaviours, make the precise drivers of these trends both difficult to disentangle, and worthy
284 of further detailed study. Even for the best-studied viruses of humans, such as influenza, we
285 are only just beginning to unravel the complexity of seasonal trends [69].

Box: What data do we need?

How can we design experiments and sampling schemes that allow the quantification of the ecological and evolutionary factors that structure species' viromes?

Sampling Design

How can we create balanced sampling from both a virus and host perspective?

- Viruses:
 - **Identification of the host that a virus is actually infecting**, because of the existence of multiple hosts in metagenomic samples (e.g. the bacterial microbiome, host dietary components, and eukaryotic parasites or symbionts). Can be done by comparing novel virus genomes to existing viral phylogenies. Non-host associated viruses can then be used as an internal control, as they should not be affected by trends in host-associated viruses.
 - **Increased attention to DNA viruses** to ascertain whether there is a dearth of DNA viruses in some ecosystems or groups
 - **Aim to characterise the within-host diversity of viral communities**, and therefore its drivers, possibly by combining short and long read sequencing [70] to distinguish between co-circulating haplotypes and structural variants.
- Hosts:
 - **Utilising carefully designed, systematic sampling of species/ecosystems** – using power analyses to determine the number of individuals, and species sampled, rather than haphazard approach
 - **Sampling multiple individuals of a host species, and in a variety of habitat types/seasons** to estimate virus prevalence, climatic, seasonal or habitat effects, and scale dependencies [71].
 - **Sampling complete food webs/trophic networks/ecosystems** – by considering which systems allow more complete sampling of potential host taxa (e.g. islands [25,56], tree fogging, ponds)
 - **Gathering data from traditionally under-sampled ecosystems** will enable us to examine the effect of different ecologies and life history traits on the structure of the virome. Current sampling biases have likely skewed our view of the ecology of even well sampled host virospheres. Predictions of viral sharing [11] or potential host-shifting will not be able to be expanded out of

well sampled (e.g. mammalian) groups without more detail on the host range and ecological context in non-mammalian viral metagenomics.

Utilising species distribution & demographic history data

How can we expand the possible virome predictors we can test using public data?

- Testing drivers of prevalence and diversity by **making use of historic climatic data, and data on anthropogenic environmental changes** such as land use change (introduction of agriculture, urbanization) and habitat disturbance levels
- By **making use of data from public citizen science projects** [e.g. 72,73], and the expertise of local forums or naturalist communities [e.g. 74,75] we can examine how more factors, e.g., Migration/range shifts, impact virus prevalence and diversity.
- By incorporating public data on the **presence/absence of symbionts, or co-infecting macro-parasites** (e.g., Varroa mites with DWV), we can assess their impact on viral prevalence and virome composition - ultimately aiming for data from whole macro, symbiont, microbiome and virome datasets.

Analysis

How can we quantify the effect size of both ecological and evolutionary drivers of species viromes?

- **By utilising a mixed-model approach** [e.g. 76], **including co-phylogenetic mixed models**, it's possible to draw out both evolutionary and ecological predictors of virus prevalence and host range [42,77]
- By **estimating beta diversity directly from linear models**, we may be able to quantify the effect of factors on virus diversity, as well as prevalence. [78]
- By **accounting for spatial and temporal autocorrelation** in analyses of the drivers of virus prevalence, we can not only make our identified drivers more robust but quantify the influence of spatial and temporal effects.
- **Developing the equivalent tools for the analysis of virome data that are already available for the analysis of bacterial microbiomes**

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287 *3.2. Host biotic factors that shape virome composition*

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290 To date, a number of potential biotic moderators of virus community diversity and structure

291 have been identified. These include life history traits, species migration histories and

292 demography. This is an area of huge potential expansion into topics such as how social
293 networks [e.g. 79,80] and behaviour [81] impact virus transmission. Here we will address
294 four descriptive factors; the composition of the population by life history traits (here host age
295 and sex), host dietary preferences, and a host species history of range movement/migration.

296

297 *3.2.1. Life history traits*

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299

300 Host age is a key feature of virus susceptibility and host immune response, as, at least in
301 vertebrates, young animals are more susceptible to viral infections, and show a consistently
302 higher prevalence compared to adults [89,90]. Taking this out to the level of whole viromes,
303 host age is arguably the most studied demographic factor affecting virome diversity and
304 composition, across a wide range of species, from humans [84] to echinoderms [85]. In the
305 future, we need an increased understanding of the impact of host age structure on virome
306 diversity and composition in invertebrates, where antibodies do not mediate susceptibility,
307 and therefore could show vastly different trends.

308

309 In contrast to host age, host sex has not yet clearly been associated with whole virome
310 composition [59,66,86]. From studies in some individual pathogens, we might predict that
311 males would show higher prevalences of viruses due to behaviour and physiology, with
312 knock-on effects on whole virome composition, perhaps decreasing alpha diversity.
313 However, studies have not all shown a clear trend in this direction, perhaps reflecting the
314 varied impact of host sex on individual viruses [87], lack of behavioural or immune
315 differences between sexes in some systems [88], the taxonomic groups considered, or a
316 result of study design.

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318 *3.2.2. Diet*

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The impact of ecosystem food web structure on species virome diversity and composition is as yet unknown, despite viruses playing an integral part in food webs, the recycling of organic matter, and transfer of energy across trophic levels [89]. At an individual level, studies of human gut viromes have provided some limited evidence that dietary variation can impact gut virome community structure [90], as it does for bacterial microbiomes [91]. However, at a broader species level, we do not know if certain types of diet, or a more phylogenetically diverse diet, drives higher virome diversity. Dietary preferences and increased dietary phylogenetic breadth could increase the opportunity for viral host shifts, and a more diverse virome. However, when predator-prey, or herbivore-plant pairings are phylogenetically distant, our current knowledge of animal viromes suggests that viruses are not often shared during these interactions [26], and that host phylogeny plays a larger role in virus sharing [25]. In the future, the study of whole food webs, and multi-species, phylogenetically controlled comparisons, will enable the effect of species diet on virome diversity and composition to be better quantified. However, in studies of wild populations, it is extremely important to distinguish between the transient gut virome, which are likely to be actually infecting dietary or prey species, and ‘resident’ viruses that cause sustained infections and go on to persist in their new host.

3.2.3. *Movement ecology and virome composition*

Species movement ecology and history, including migration, dispersal, and a species history of invasion or introduction are likely to have significant impacts on current virome composition. The idea that an individual’s movement ecology and demographic history influence the prevalence and diversity of parasites is not a novel one [92]. However, our understanding of the way in which species viromes are influenced by this is still in its infancy.

348 There is a pressing need to understand the role of species' histories of introduction and
349 dispersal in shaping the current virome, given how rapidly distributions are shifting in tandem
350 with climate change, and the frequency of introductions via global trade and travel [93]. For
351 example, increasing ocean temperatures are likely to drastically shift marine species'
352 distributions [94]. As these ranges shift poleward in response to changing climates, species
353 will be pushed into contact with novel viruses [95]. They will also expose native and naïve
354 host species to novel viruses, perhaps with devastating consequences. Species invasions
355 may also change species-virus relationships, and the diversity of the whole host ecosystem
356 – with potential knock-on effects such as the 'dilution effect' [96]. Additionally, the outcome of
357 these species-virus interactions will be influenced by the phylogenetic relationships between
358 hosts, including their evolutionary rarity (i.e., how phylogenetically distant a host is from the
359 rest of the host community)? For example, introduced hosts that are phylogenetically
360 isolated from other members of the host community have lower disease pressure [97].
361 However, we still lack a comprehensive understanding of whether dispersing individuals act
362 as sources or sinks for viral infection, and how these patterns vary with host taxonomy. For
363 instance, do recent arrivals in an ecosystem exhibit reduced virus diversity or abundance, or
364 do they tend to act as sources of novel viruses?

365

366 Species movement also affects virome composition through the life history strategy of
367 migration. For example, migration can lead to escape from pathogens, may lead to infected
368 individuals being removed from populations, may allow recovery from infection or spatially
369 isolated infected and uninfected individuals [98]. Studies of single viruses, such as avian
370 influenza and West Nile virus, have shown that migration might have increased the spread of
371 disease in general by increasing contacts and thus, virus exposures [82]. Future work should
372 investigate how migratory strategies shape variation in both virome composition and risk of
373 transmission at the individual level.

374

375 *3.3. Anthropogenic factors that influence virome community structure*

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377 Human driven changes to the natural environment, such as climate change, urbanisation,
378 habitat disturbance, and altered nutrient cycling, have a profound impact on host
379 biodiversity, and alter the ecology of systems governed by the abiotic and biotic factors
380 described above. However, we poorly understand the broader consequences of such
381 changes to virome community diversity.

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383 Studies in humans suggest that urbanisation can have a profound impact on the diversity
384 and composition of viromes [99], with studies illustrating important differences in the viromes
385 of urban and rural-living humans [102]. An important caveat is that studies involving humans
386 have largely focussed on the gut virome (i.e. bacteriophages) rather than viruses that infect
387 human cells, so what we are observing could be driven by differences in the bacterial
388 microbiome.

389

390 In wild populations, anthropogenic factors have a profound impact on the distribution and
391 home ranges of many host species, with the potential to facilitate the cross-species
392 transmission of viral pathogens, affecting wildlife conservation, agriculture and human health
393 [103]. While few studies have assessed the impact of host biodiversity changes on the
394 virome, our limited evidence suggests some pristine/undisturbed habitats have increased
395 viral diversity, likely related to an increase in host species diversity in some systems [104].

396

397 *3.4. Coinfection as a modulator of species viromes*

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400 Coinfections, the simultaneous infection of a host with multiple viruses, parasites or
401 symbiotic microbiota are common in nature and may alter the outcomes of individual
402 infections [105]. Within coinfecting hosts, viruses can interact directly – such as in the
403 transactivation of one virus's gene expression by the proteins of another [106]. Similarly
404 viruses can interact indirectly (with other viruses, microbes or parasites) through modulation
405 of host components such as immune activity or resource availability [107]. These
406 interactions can create synergistic, exploitative, or competitive relationships between
407 pathogens.

408 There is not yet a clear link between coinfection interactions, coinfection prevalence, and
409 virome composition. However, the consequences of interactions between pathogens in
410 individual hosts can affect the prevalence of viruses across host populations. For example,
411 negative interactions between influenza A virus and Rhinovirus in humans can lead to
412 fluctuating and asynchronous seasonal prevalences of each virus [108], and has been
413 suggested to have delayed the introduction of the 2009 H1N1 influenza virus pandemic to
414 Europe [109,110] (after which H1N1 is thought to have disrupted the epidemic transmission
415 of another respiratory virus [111]).

416

417 Compared to single infections, coinfections can alter the relative fitness of different viruses
418 and virus genotypes [112,113], and may play a role in generating and maintaining virus
419 diversity. For example, in nucleopolyhedrovirus coinfections of pine beauty moths (*Panolis*
420 *flammea*), the relative fitness of virus genotypes during single infection does not correspond
421 to fitness during coinfection, and are further influenced by the ecological context of the
422 infected host [114]. At a population level, coinfection-induced changes in the rank order of
423 virus fitness are expected to fluctuate with coinfection prevalence [115]. Additionally, the
424 outcome of coinfections is likely to be heavily influenced by the sequential timing of
425 infections [116], with within-host viral diversity sometimes dependant on the order in which
426 viral infections occur. In this way, coinfection may be an important – yet relatively

427 understudied – mechanism for the maintenance of virus diversity and the shaping of host
428 viromes [117].

429

430 In the future, more studies of the role of co-infections are likely to shift away from PCR
431 detection of known viruses, and will use the detection of unknown viral infections in the wild
432 using metagenomic sequencing, across a food web, and the sampling of multiple individuals
433 from each host species (e.g. [77,129]). Such studies will enable us to simultaneously
434 quantify the relative impact of both within-host (co-infections) and between host
435 (diet/predator-prey) interactions on virome composition.

436

437 4. Perspectives

438

439

440 Whilst we have attempted to synthesise the current evidence on what drives the diversity
441 and composition of species viromes, the majority of data still come from single-host, single-
442 virus studies that might not generalise to whole virus communities, could be focused on viral
443 ‘oddities’, such as extremely virulent viruses, and hence are unlikely to represent the
444 majority of the virome. With ever decreasing costs of RNA sequencing, hypothesis-driven
445 and structured sampling of viromes from multiple individuals in host populations, and
446 multiple host species in a community, is becoming more affordable.

447

448 Despite the unique challenges that virome studies bring, there are many exciting areas for
449 expansion in this field, and many outstanding questions about the basic relationships driving
450 the distribution of viruses across host species. For example, do areas with a greater diversity
451 of host species generate higher virome diversity, or is this dependant on the phylogenetic
452 composition of the host community? Are species with more diverse viromes more likely to

453 acquire more viruses, and are generalist viruses more likely to infect new species than
454 specialist ones [120]? At a population level, what is the relationship between population size,
455 and virome diversity [e.g. 59]? This is particularly interesting to consider in the tropics, as
456 numerous studies have shown the role temperature or UV play on virus transmission by
457 reducing environmental persistence. Another unexplored aspect of the drivers of virome
458 composition are social networks, and how associations within social networks drive virus
459 transmission [e.g. 80]. In the future, can we determine the mechanistic basis of the host-
460 virus associations, in particular, the phylogeny-related variation? Can we use trait, gene, or
461 motif-based models/phylogenies of viruses to test the predictive power of these features in
462 driving the distribution of viruses? Perhaps we can also move towards a more holistic,
463 whole-microbial community approach to these studies, with exciting opportunities to study
464 covariation among viruses, bacteria and fungi across a broad host phylogeny [27].

465

466 These questions are particularly timely due to ongoing global and climate change. Will
467 increasing urbanisation and global movement drive an increase in the virome diversity of the
468 urban populations of wildlife, or a decrease in virome diversity due to lower host diversity?
469 With global changes in non-urban areas, such as conversion to monoculture, what are their
470 impacts on virome diversity downstream? Or, as in the case of habitat fragmentation, will the
471 break-up of diverse ecosystems result in increased prevalences for the most abundant viruses
472 and a corresponding reduction in virome evenness?

473

474 By understanding the evolutionary and ecological drivers of the virosphere, and in particular,
475 the proliferation of zoonotic pathogens through communities and landscapes, we can also
476 seek to mitigate these risk factors. For example, methods of reducing the prevalence of
477 harmful viruses, such as reducing the prevalence of vector-borne viruses through dilution
478 effects (selectively increasing livestock densities), have been proposed [121]. However, their

479 effectiveness will depend on the degree to which virus prevalence is driven by specific host
480 densities, and how this changes with local spatial and temporal variation in abiotic factors.
481 With a greater understanding of these drivers we can not only aim to control viruses with
482 impacts on human, agricultural and wildlife health, but also understand and appreciate the
483 role viruses play as components of whole ecosystems.

484

485 **Acknowledgements**

486 Thanks to Darren Obbard for discussion.

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488 any Author Accepted Manuscript version arising from this submission.

489

490 **Funding**

491 BL, MAW and RMI are funded by a Sir Henry Dale Fellowship jointly funded by the
492 Wellcome Trust and the Royal Society (grant no. 109356/Z/15/Z)

493 <https://wellcome.ac.uk/funding/sir-henry-dale-fellowships>.

494 JLG is funded by a New Zealand Royal Society Rutherford Discovery Fellowship (RDF-20-
495 UOO-007) and the Webster Family Chair in Viral Pathogenesis.

496 ECH is funded by a National Health and Medical Research Council (NHMRC) Investigator
497 award (GNT2017197) and by AIR@InnoHK administered by the Innovation and Technology
498 Commission, Hong Kong Special Administrative Region, China.

499 XAH is funded by the Leverhulme Trust (RPG-2020-320).

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