Bat-borne virus diversity, spillover and emergence

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Abstract | Most viral pathogens in humans have animal origins and arose through cross-species transmission. Over the past 50 years, several viruses, including Ebola virus, Marburg virus, Nipah virus, Hendra virus, severe acute respiratory syndrome coronavirus (SARS-CoV), Middle East respiratory coronavirus (MERS-CoV) and SARS-CoV-2, have been linked back to various bat species. Despite decades of research into bats and the pathogens they carry, the fields of bat virus ecology and molecular biology are still nascent, with many questions largely unexplored, thus hindering our ability to anticipate and prepare for the next viral outbreak. In this Review, we discuss the latest advancements and understanding of bat-borne viruses, reflecting on current knowledge gaps and outlining the potential routes for future research as well as for outbreak response and prevention efforts.

Bats are the second most diverse mammalian order on Earth after rodents, comprising approximately 22% of all named mammal species, and are resident on every continent except Antarctica. Bats have been identified as natural reservoir hosts for several emerging viruses that can induce severe disease in humans, including RNA viruses such as Marburg virus, Hendra virus, Sosuga virus and Nipah virus. In addition to direct isolation of these human pathogens from bats, accumulating evidence suggests that other emerging viruses, such as Ebola viruses, severe acute respiratory syndrome coronavirus (SARS-CoV), SARS-CoV-2 and Middle East respiratory coronavirus (MERS-CoV), also originated in bats, even if other hosts, such as civets for SARS-CoV and camels for MERS-CoV, are proximate reservoirs for human infection

A growing list of emergent coronaviruses, including the Swine acute diarrhoea syndrome coronavirus, which emerged from horseshoe bats and killed >20,000 pigs, and the ongoing COVID-19 pandemic, further underscores the ongoing threat of bat-borne viral emergence.

Bats harbour a high viral diversity relative to other mammalian orders; indeed, recent studies have suggested that viral diversity is reflective of the number of species, with Rodentia (rodents) and Chiroptera (bats) containing the most species among mammals. This viral diversity flags bats as an important taxonomic group for global viral discovery and zoonotic disease surveillance efforts. These efforts, ultimately aimed at identifying and mitigating future emergence events of bat-borne diseases, have identified thousands of novel bat-derived viral genomic sequences over the past decade. However, as most of these sequences span polymerases and not the surface proteins that often govern cellular entry, little progress has been made towards translating sequence data from novel viruses into a risk-based assessment to quantify zoonotic potential and elicit public health action. Further hampering this effort is an incomplete understanding of the animals themselves, their distributions, behaviours and interactions with the environment, and the processes that lead to contact with humans.

In this Review, we discuss the current state and knowledge gaps of bat virus ecology (Box 1) and the molecular barriers to zoonotic disease emergence; we also review advances and challenges in pandemic preparedness and provide a framework for addressing critical deficits in our understanding of bat-borne viruses.

Viral diversity in bats

Research on bat viruses dates back to the 1930s, when Joseph Pawan first identified rabies virus in bats and experimentally infected several different bat species with the virus in Trinidad. The following decades saw a slow accumulation of newly discovered bat viruses and an exponential increase after the discovery and isolation of SARS-related coronaviruses (SARSr-CoVs) from bats in 2002 and the concomitant rise of next-generation sequencing technologies. Field research on other bat-borne emerging pathogens, including Nipah virus and Marburg virus, in combination with the decreasing cost of next-generation sequencing technologies, has spawned the current era of intensive bat viral discovery efforts. These efforts have led to the identification of whole clades of viruses and genomic sequences closely
Bats as unique virus hosts

**Viral persistence in bat cells and populations.** Shedding of zoonotic viruses from bat populations can vary considerably in space and time, with peaks in shedding sometimes coinciding with spillover to other species. Understanding the mechanisms that drive the circulation of bat viruses in populations, including causes of peaks in shedding, is necessary to predict when and where spillover may occur. Another key requirement is understanding the biology of these infections in bats. If zoonotic infections cause a simple dynamic of acute infection followed by recovery and resistance, as originally assumed for henipaviruses and sometimes assumed for filoviruses, then epidemic cycles will be driven by oscillating and connectivity, population size and other factors that drive transmission among bats. Some observations have been inconsistent with this framework; for example, the lack of an association between high virus prevalence and large host population size, whereas other observations suggest that this dynamic may be possible, for example, the short infectious periods in bats inoculated in captivity. Other studies hint at persistent and/or recurrent infection in bats. If bats are persistently infected, immune competence may control shedding, and factors such as stress would drive shedding peaks.

For example, there is anecdotal evidence of captive bats seroconverting against Nipah virus after a period of seronegativity, suggesting that persistent infection and episodic shedding may be possible. A recent observation of bats synchronously shedding multiple paramyxoviruses during a pulse of Hendra virus shedding is more easily explained by a population-level stressor affecting host immune competence; however, there could be other explanations. Curiously, naïve *Roussettus aegyptiacus* bats became infected with Marburg virus months after an inoculation and transmission experiment had ceased, suggesting that Marburg virus persisted within the small group of 36 experimental bats for 7 months.

Within-host cycles of infection in bats have been extremely difficult to determine, and the data required to assess competing hypotheses have not yet been available. Results from inoculation experiments in bats have been difficult to interpret, and the limited duration of almost all bat virus experiments precludes investigations...
Type I interferon
A large group of related cytokines that bind to widely expressed interferon-α receptors and are responsible for regulating the immune response to infection.

into viral persistence within hosts. Ideally, genetic data from viruses infecting individually marked bats over time could be used to determine if viruses persist within individuals, but recapturing most bats is extremely difficult, and few studies collect data longitudinally. Recently, researchers have been able to make inferences about viral circulation in bats by fitting mathematical models of disease dynamics to longitudinal serological data. A study using such methods determined that persistence or reinfection of a circulating henipavirus was likely in Eidolon helvum bats. Research combining longitudinal sampling of bats with viral genomics, antibody surveys and mathematical models will be required to infer zoonotic pathogen circulation in bats.

Intrinsic bat resistance. Bats are seemingly refractory to viral pathogenesis, and their metabolism has been at the centre of the long-standing ‘flight as fever’ hypothesis underlying this phenomenon. Several groups have speculated that the high-energy metabolic demands of flight lead to elevated body temperatures in bats, mimicking the fever that occurs in other animals during immune activation, which may broadly impact viral pathogenesis. However, experimental studies have shown that filoviruses replicate similarly in bat cells regardless of ambient temperatures. Beyond body temperature, knowledge gaps on bat reservoir species and their flight behaviour, immunity and metabolism obscure how bat metabolism relates to immunity.

Innate bat immunity. Although viruses such as Nipah virus and Marburg virus have been experimentally shown to replicate in and shed from their bat host species, a striking feature of these infections is that the bats lack overt signs of pathology. The observation that bats may be refractory to, or tolerant of, viral infection was noted as early as 1936, yet the immunological mechanisms that underpin this phenotype have only begun to be elucidated in the past few years. Current data suggest that the classical pathology caused by strong activation of the immune system in response to viral infection that is seen in humans and laboratory animal models does not occur in bats. The lack of pathology observed in bats is likely due to a combination of differences in viral tissue tropism and host immune responses. Viral replication and shedding in bats in combination with an apparent lack of disease may allow for the efficient maintenance and dissemination of viruses.

Interferon-α (IFNa), IFNβ and IFNγ pathways vary in their level of activation between bat and human cells in response to viral infection. Some of these studies have shown dampened immune responses in bats, whereas others have shown heightened responses to infection. The consequences of these differences for overall pathology in bats are still to be determined. A notable finding common to all of these studies is that, regardless of the host species, all of the bat cell lines tested support filovirus infection, suggesting that the innate immune pathways assessed in these cell culture assays do not form barriers to infection.

Broader characterizations of bat innate immunity have provided some insights into the differences between bat and human immune responses. For example, Pteropus spp. bats have a substantially smaller type I interferon genomic locus than other mammals, yet they have constitutive basal expression of their IFNα genes, regardless of stimulation. How a smaller type I interferon locus might influence viral disease is

![Virus families with fewer than 50 different sequences](image)

**Fig. 1 | Currently described bat virus diversity.** Publicly available genetic sequence data for bat-derived viruses (database of bat-associated viruses) were pooled and categorized by viral family. Of note, large parts of the bat virus diversity remain uncharacterized, and discovery efforts have prioritized virus families with known zoonotic potential such as the Coronaviridae, ssDNA, single-stranded DNA.
Box 2 | Bats as animal models

Although a growing number of laboratories are studying infectious diseases that emerged from bats in laboratory settings, there are few examples of bat species being successfully established as animal models. This is in part owing to the unique handling requirements for volant animals, challenges with keeping many insectivorous bats fed and healthy and difficulties in supporting complex behavioural adaptations (for example, hibernation for temperate bats). These challenges are compounded by a limited availability of breeding stock, regulations on importation of live animals and the high costs of maintaining animals in biosafety level 3 or 4 facilities, which are required for virus studies. Nevertheless, several bat species have been established as animal models, including *Pteropus* spp. for Nipah and Hendra virus, *Eidolon helvum* for African Henipavirus, *Rousettus aegyptiacus* for Marburg virus, *Artibeus jamaicensis* for Zika virus, MERS-CoV and rabies virus and *Myotis lucifugus* for the fungal disease white nose syndrome. These studies have led to valuable discoveries, for example, showing that *R. aegyptiacus*, from which Marburg virus was isolated, is refractory to infection with many other viruses that are associated with other bat reservoir species, making the broad use of this bat species in disease pathology modelling questionable. Beyond the technical challenges of working with these animals, a bigger issue with bats as animal models lies in our fundamental pathology modelling questionable. Almost all currently established animal models in virology are centred on severe disease phenotypes and high levels of viral replication. This custom contrasts with our current understanding of bat virus biology, which assumes that bats exhibit minimal pathology and likely low levels or short temporal bursts of viral replication. Recent experimental infection studies with *Tacaribe virus* and *Lagos bat virus* in their respective natural reservoir bat species resulted in severe disease and mortality, showing that the paradigm that bats are resistant to highly pathogenic viruses should be addressed at the level of specific host–pathogen interactions rather than as a generalization for a complete animal order. Comparative studies between animal models of human disease and bat animal models are needed to understand the mechanisms responsible for the differences in disease severity of bat-borne viruses observed in natural reservoir and spillover host species.

Adaptive bat immunity. The adaptive humoral immune response, mediated by antibodies, is as enigmatic as the innate immune system in bats. Long-term laboratory experiments with Marburg virus challenge in the Egyptian fruit bat and with rabies virus challenge in the insectivorous big brown bat showed that, although antibodies to the respective pathogens arose, they rapidly waned below detectable levels after ~3 and 5 months, respectively. Although bats had low levels of antibodies to Marburg virus 22 months after the initial challenge, re-challenge resulted in dramatically reduced dissemination and viral spread. Interestingly, *R. aegyptiacus* challenged with Ebola virus, Marburg virus or *Sosuga* virus generated detectable antibody responses post challenge, but none of the antisera could neutralize the live virus, suggesting that antibody-dependant cell-mediated responses are crucial in the viral clearance of these viruses. By contrast, bats generate virus-neutralizing antibodies to Nipah virus and rabies virus upon experimental challenge; however, live virus may be concurrently detected in saliva or urine, suggesting incomplete virus clearance. Curiously, *Sosuga* virus and Nipah virus are both paramyxoviruses and yet elicit different types of antibody responses — the mechanisms and underlying reasons are unclear. Recent studies support the notion that the immune response in bats is adapted for pathogen tolerance and regulated to mitigate immunopathology, potentially promoting incomplete viral clearance and asymptomatic infection.

Challenges of studying viral infections in bats. A major problem facing all studies of bat-derived virus cell biology is the lack of available reagents and animal models, compounded by the enormous taxonomic diversity of these animals. Most mammalian cell lines currently available are not derived from bats and do not support replication of the majority of viruses being discovered. For example, the isolation of bat-derived coronaviruses has been challenging owing to a limited ability to infect the standard, typically primate-derived, cell lines used in most laboratories. Even cell lines derived from the same bat species that the viruses were originally sequenced from may fail to support replication owing to loss of expression of the host receptor. Thus, there is an urgent need for more cell culture reagents that can better facilitate virus isolation, either new cell lines capable of supporting replication of bat viruses or genetically modified versions of existing cells, such as *Vero* cells, to increase their susceptibility to viral infection with bat viruses. Ideally, these new cell lines should be derived from a wide range of species and tissues. In addition, organoid systems incorporating multiple cell types within a 3D architecture to reproduce tissue-specific functional properties could potentially facilitate the translation from in vitro single-cell type studies to organ-specific host–pathogen interaction studies. Lastly, live animal models will also be crucial for understanding the implications of molecular findings in bat cells for the course of infection in the natural host.

Whereas large-scale CRISPR–Cas9-mediated knockout and activation screens have been valuable for identifying human proteins involved in viral infection, they have yet to be applied to other host species, including bats. Given that an annotated transcriptome is now available for the *R. aegyptiacus* bat, similar screens could be performed in *Rousettus* cells to elucidate the factors involved in filovirus infection, for example. The ambitious Bat1K project has begun the process of generating genome sequence data for all extant bat species, and the results of this effort will undoubtedly be an invaluable resource in elucidating specific bat–pathogen interactions.

Many of the earliest studies of bat immunity focused on the humoral adaptive immune response in bats, yet we still lack an understanding of the regulation of B cell proliferation, affinity maturation and the mechanisms

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**Type III interferon**
A small group of related cytokines that bind to the interferon-κ receptors found on epithelial cells and are responsible for regulating the immune response to infection.

**Bat1K project**
A global research initiative to sequence and annotate the genomes of all bat species, starting with more than 1,000 of the most relevant species for global health.
underlying the transitory nature of bat antibody responses. Countless studies have conducted serological surveillance for antibodies to specific viruses in wild bats; however, there is relatively little experimental data with bats as a model system, particularly spanning longer periods of time. The information available from field-collected samples is limited by the unknown history of sampled bats with regard to their reproductive status, age, sex, nutritional status and infection status, with any number of pathogens potentially influencing the immune response.

**Viral spillover from bats**

The spillover of bat-associated viruses requires a combination of factors, including ecological opportunity for contact, virus–host molecular and cellular compatibility, and a permissive or circumvented immune response, as described in detail herein. Yet, despite the various potential barriers and the fact that many spillover events may go undetected by surveillance systems, there is a growing list of recent bat-borne zoonotic spillover events. This list includes examples of direct bat-to-human spillover, which are supported by both epidemiological evidence and molecular detection of monophyletic viruses between bat and human populations; the examples include near annual outbreaks of Nipah virus in Bangladesh since 2001 (REF. 66), several Marburg virus outbreaks across Africa and outbreaks of rabies virus and other novel Lyssaviruses globally. Other examples of indirect bat-to-human spillover that involve intermediate hosts are also supported by epidemiological and molecular evidence, including Hendra virus in 1994 via horses and Nipah virus in Malaysia in 1997 and 1998 via pigs. The SARS-CoV 2002–2003 outbreak in southern China and the 2019 SARS-CoV-2 emergence in central China were retrospectively connected to bat populations via molecular evidence and appeared to involve intermediate hosts. Several viruses closely related to SARS-CoV were found in bats, and actual SARS-CoV was directly isolated from animals in open-air markets. Analogously, several bat viruses have now been identified to have a high similarity to SARS-CoV-2 (REF. 72). In several other cases, bat-to-human spillover was assessed retrospectively via serological human cohort studies; for example, Henipavirus spillover among bat hunters in Cameroon, bat-borne reovirus (Melaka virus and Pulau virus) exposure in people living in close proximity to bat roosts on Tioman Island, Malaysia, as well as in a random sample screened in Singapore, Filovirus exposure of bat hunters in India and ongoing human exposure to SARSCoVs in rural communities in China occurring after the 2003 SARS outbreak.

**Molecular biology of transmission**

All viruses, regardless of classification or origins, must be able to subvert and overcome various molecular factors within their hosts in order to replicate and spill over into new species. Every stage of the viral life cycle relies on numerous protein interactions with the host cell, including viral binding and entry, recruitment of host factors essential for viral replication, suppression of antiviral host factors, assembly and egress from the cell, and evasion of the host immune system (FIG. 2). Viral–host protein interfaces have been shown, through functional and structural studies, to be remarkably specific and involve multiple points of contact. Even single amino acid variations can impact or abrogate a viral–host protein interaction between different species and form a molecular block, or species barrier, to viral replication. The complexity of viral–host protein interactions is compounded by how many interactions occur during any given infection. Recent proteomics studies have identified at least 194 protein interactions between Ebola virus and human host cells, 198 virus–host protein interactions for Zika virus, 101 for Nipah virus and over 300 for influenza A virus. Given that even small perturbations in these complex networks of virus–host interactions can make the difference between a dead-end infection or viral emergence in a new host species, it is likely that the majority of bat-borne viruses fail to infect novel species as a result of within-host barriers.

**Cell entry of zoonotic viruses.** One of the first major virus–host protein interactions that occurs during the course of infection is at the level of viral cell entry, when the virus interacts with the host receptor to facilitate the release of viral components into the cytoplasm. Depending on the virus, this process can involve one or more viral proteins, one or more host components and encompass several steps occurring at the cell surface or at an internalized membrane.

It is not surprising that many bat-borne zoonotic viruses have evolved to use highly conserved host
molecules for cell entry that have little genetic variation between different species. Henipaviruses bind to the ephrin family of signalling proteins\textsuperscript{2,3,94}. Filoviruses bind to the cholesterol transporter Niemann–pick C1 (NPC1)\textsuperscript{4,96} and Betacoronaviruses have been shown to bind different common cell-surface proteases, including angiotensin-converting enzyme 2 (ACE2) in the case of SARS-CoV and SARS-CoV-2 (REF\textsuperscript{7}) and dipeptidyl peptidase IV (DPP4) in the case of MERS-CoV\textsuperscript{98}. Indeed, these receptors are nearly identical, at least in the regions that interact with the virus, between various bat species, intermediate host species, such as camels and palm civets, and humans\textsuperscript{94,95,99}.

Restriction of cell entry is not easily overcome by viruses. For example, wild-type mice are completely resistant to infection with MERS-CoV because of differences in murine DPP4 glycosylation from human DPP4 (REF\textsuperscript{100}). Despite great effort from the animal disease-modelling community, to date, there is no MERS-CoV isolate that can use wild-type murine DPP4, likely because too many viral adaptations are necessary. However, partial blocks to cell entry are more easily overcome. Recently, it has been shown that MERS-CoV can rapidly acquire single-point mutations to increase its compatibility with DPP4 from different bat species\textsuperscript{45}. Some MERS-related CoVs discovered in bats, which appear nearly identical to MERS-CoV over most of the genome, have mutations at key binding sites across the receptor–binding spike glycoprotein and are thus unable to bind to human DPP4 (REF\textsuperscript{95}). Similar types of viral adaptation have been observed for other zoonotic viruses such as SARS-CoV\textsuperscript{114}, parvoviruses\textsuperscript{115} and avian influenza A virus\textsuperscript{116}. Taken together, the ability to use conserved host receptors and readily adapt to receptor variation between species are two hallmarks of viruses that have spilled over into the human population.

The genetic diversity of many RNA viruses can be attributed to high mutation rates, short generation times and the strong selective pressure of the host environment (during natural infections or after vaccinations); however, positive-stranded RNA viruses, including SARS-CoV and MERS-CoV, have relatively low mutation rates associated with 3′–5′ exoribonuclease proofreading activity\textsuperscript{104–108}. The rapid evolution of coronaviruses to the host environment is largely driven by the high rates of genetic recombination, which facilitate the acquisition of multiple mutations in a single event. Inheriting multiple genetic changes at once can have dramatic effects on viral replication and subsequent adaption to new host environments. Recent phylogenetic analyses have revealed that the variation in MERS-CoV circulating in camel populations is largely driven by recombination\textsuperscript{109–112}. Although SARSCoV-CoVs have been identified and isolated from bats, no single bat isolate perfectly matches the human strains. For example, some SARSr-CoVs can use the human receptor but vary drastically from SARS-CoV in the 3′ end of their genome, whereas other SARS-CoVs are nearly identical to SARS-CoV in this region but fail to interact with the human receptor\textsuperscript{113}. In further support of these findings, the entire SARS-CoV genome has now been sequenced across multiple separate but related viruses circulating in bats, strongly suggesting that the human virus is a recombinant form of these ancestral variants\textsuperscript{114}.

Outside coronaviruses, recombination in the rabies virus glycoprotein was shown to facilitate cross-species transmission from bats to skunks and raccoons\textsuperscript{115}. Thus, in addition to the rapid mutation rate characteristic of many RNA viruses, recombination provides an additional mechanism to rapidly overcome barriers in novel host species.

**Post-entry virus–host interactions of zoonotic viruses.** Whereas our understanding of viral entry as a species barrier is becoming clearer for many emerging zoonotic viruses, cellular blocks beyond entry are more elusive and remain largely unknown. However, research over the past 20 years with well-studied zoonotic pathogens, such as lentiviruses, including HIV and its evolutionary predecessor simian immunodeficiency virus, and influenza A virus, which includes avian influenza A virus, has led to the identification of numerous human and primate intracellular species barriers in the form of dependency factors, which these viruses rely on to replicate, and restriction factors, which are antiviral proteins that specifically interfere with viral replication\textsuperscript{116}. For example, the ability of the simian immunodeficiency virus accessory protein Vif to antagonize the host restriction factor APOBEC3 is vital in determining the potential host breadth in non-human primates\textsuperscript{117,118}. The avian influenza A virus accessory protein PB1-F2 disrupts mitochondrial antiviral signalling more efficiently in the avian host than the truncated PB1-F2 common in mammalian influenza A viruses\textsuperscript{119}. Although the post-entry species barriers that limit host breadth for lentiviruses and influenza viruses are likely different from those that limit host breadth for the emerging bat-borne infectious diseases, the research framework for these well-studied host–pathogen systems can serve as a roadmap for research into bat-borne host–pathogen systems.

Recent experimental infection studies in bats provided evidence of species-level post-entry barriers, suggesting that some bat-borne viruses are likely host specific and may have a limited ability to transmit between certain bat species. For example, the *Rhinolophus* spp. bat coronavirus WIV1-CoV and Ebola virus can use the cellular receptors from *R. aegyptiacus* bats to enter cell lines, but these bats fail to support any WIV1-CoV infection and only poorly support Ebola virus replication\textsuperscript{114,120,121}. Transcriptomics studies have identified several immune signalling pathways that are activated differently in human versus *Roussetus* spp. cells\textsuperscript{115,116}. Other studies have taken more direct approaches to identify post-entry barriers to replication. Mass spectrometry has pinpointed the host E3-ubiquitin ligase, RBBP6, as a negative regulator of Ebola virus transcription, which functions by binding VP30, a viral protein that is key in replication\textsuperscript{118}. A similar study found that the key Ebola virus protein involved in antagonizing the host interferon pathway, VP35, forms an essential interaction with host TRIM6 protein and that the disruption of this interface reduces viral replication\textsuperscript{122}. Additionally, tetherin\textsuperscript{123–125} and IFITM host proteins\textsuperscript{126}...
Moving beyond virus discovery

Worldwide consortia such as the USAID PREDICT programme as well as many independent academic laboratories around the world have used a combination of consensus PCR screening and deep sequencing to characterize thousands of novel viral sequences in samples taken from healthy bats\(^\text{[127]}\). Many of these novel viruses, or viral fragments, are phylogenetically related to pathogens of interest to public health; however, the capacity for these novel viruses to cause future outbreaks typically remains unresolved. With viruses isolated or sequenced directly from bat samples and molecular approaches, including viral pseudotype studies and reverse genetics, researchers have been able to demonstrate the potential of novel viruses to replicate in human cells or use human receptors for entry — the first step to incorporate virus discovery into a more comprehensive risk-reduction framework\(^\text{[14,63,136]}\) (FIG. 3). Characterizing the breadth of naturally occurring bat-borne viruses, including close relatives to human viruses, is critical and leads to important insights. For example, the discovery and subsequent investigations of a novel non-pathogenic henipavirus\(^\text{[127]}\), Cedar virus, related to Nipah virus and Hendra virus, have proved invaluable in revealing the genetic determinants of pathogenicity in henipaviruses. Unlike Nipah virus, Cedar virus does not produce V or W proteins, which are responsible for antagonizing the host interferon pathway, and it also relies on different host cell receptors\(^\text{[16]}\). The identification of viruses related to Ebola virus in various bat species, including the novel Bombali virus, has provided additional support for bats as reservoirs for Ebola viruses and, even though there are no reports of filovirus haemorrhagic fever in China, filoviruses have also been identified in Roussettus spp. bats in China\(^\text{[14]}\). Thus, further research is needed to confirm host species range and the potential for human infection\(^\text{[17]}\). Ancestral variants of zoonotic coronaviruses similar to SARS-CoV, MERS-CoV and SARS-CoV-2 have been identified from bats, with some viruses related to SARS-CoV even being capable of using the human receptor ACE2 \(^\text{[18]}\). Although both SARS-CoV and MERS-CoV have also been isolated from intermediate hosts (palm civets\(^\text{[14]}\) and camels\(^\text{[12]}\), respectively), it is possible that these two viruses can be directly transmitted to humans from bats.

Linking surveillance and control

Although whole genome data are needed for downstream and comparative studies, even knowing the taxonomic family of a novel pathogen causing an outbreak can help narrow down control and treatment strategies. Novel diagnostic platforms such as the GeneXpert\textsuperscript{TM} (Cepheid Inc.), which is a semi-automated PCR-based test, enable rapid, multiplexed detection of a wide panel of human pathogens and are constantly being improved to increase sensitivity and pathogen coverage. Virus discovery efforts are producing a wealth of genome sequencing data that are then made publicly available through online data repositories. Pan-serological assays are also facilitating virus discovery and providing insights into antibody cross-reactivity\(^\text{[19]}\). The resulting datasets provide insight into the existing variation in

Viral pseudotype

A genetically modified virus that has incorporated the glycoprotein of another virus and can produce a measurable reporter, such as GFP or luciferase, upon infection of a cell.

**Fig. 3** | Using functional viromics to move beyond zoonotic virus discovery. Current viromics research, mostly based on the targeted amplification of viral sequences or sometimes metagenomics, stops after the identification of novel viral sequences in animals. Large-scale functional screens of viruses in vitro will facilitate transmission and pathogenesis studies in vivo and ultimately lead to the development of ‘One Health’ intervention strategies, such as vaccination of humans or reservoirs or intermediate animal hosts, as well as other measures to reduce the risk of contact and viral transmission at the animal–human interface.

which have been shown to inhibit lentiviruses and influenza A virus, respectively, have broad antiviral effects to Ebola virus and SARS-CoV. Tetherin from fruit bats inhibits Nipah virus but not Ebola virus\(^\text{[127]}\). Compared to more well-studied viruses with a zoonotic origin, such as influenza A virus and HIV, there is still much to uncover for post-entry species barriers of emergent, bat-derived viral pathogens. Large-scale CRISPR–Cas9-mediated knockout and activation screens in human cells have recently identified specific host factors that are essential to flaviviruses\(^\text{[128–130]}\), HIV\(^\text{[131]}\), Epstein–Barr virus\(^\text{[132]}\) and influenza A virus\(^\text{[133,134]}\), but such data have not yet been generated for bat-derived viruses.
viral families, allowing for the development of diagnostic and surveillance assays broadly targeting virus clades. For example, Bombali ebolavirus was initially discovered in bat samples using a consensus PCR assay developed to target a region of the viral genome for which the nucleic acid sequence is conserved between related viruses. The identification of key reservoir species and the development of better models to predict virus spill-over events could enable targeted prophylactic vaccination campaigns of humans and potential reservoir hosts or intervention strategies to minimize contact between bats and humans. For example, high-risk populations that geographically overlap and have high levels of contact with bats and other wildlife could be vaccinated against Ebola virus to prevent outbreaks rather than responding to them after spillover, and local communities could benefit from education campaigns on how to live safely around bats and reduce direct contact

Alongside these measures, efforts should be taken to reduce bat habitat destruction, which results in increased contact between bats and humans and is considered a cause of viral spillover

Next-generation vaccine technologies are platforms that are broadly and rapidly adaptable for different types of viral pathogens. Importantly, several of these platforms use genetically modified viruses, such as the vesicular stomatitis virus (VSV) and ChadOx1 platforms, which can induce protective immunity in humans, mice, guinea pigs, non-human primates and livestock to a number of pathogens, including bat-borne Ebola virus and Nipah virus. Vaccine efficacy in animals, for example, horse vaccination for Hendra virus, including live-stock and other peri-domestic animals, may even enable proactive measures to reduce cross-species transmission of bat viruses to humans. Although still nascent and only in early clinical trials, novel platforms such as DNA-based and mRNA vaccines offer the potential for an incredibly rapid response time from pathogen discovery to therapeutic intervention. Using these technologies, researchers were able to test the first Zika vaccine in mice and non-human primates within 3.5 months of the initial outbreak in 2015 and, more recently, a similar RNA-based vaccine was designed for SARS-CoV-2 and entered human clinical trials only 2 months after the virus sequence was published. Other platforms based on VSV or adenovirus are already FDA approved and have been shown to be effective in several species and for most of the major emerging viruses identified to date.

Depending on the route of transmission, pre-emptive control strategies may also include low-cost, low-technology (that is, ecological) countermeasures. For example, Nipah virus is believed to be transmitted to humans through date palm sap collection containers that have been contaminated with virus-containing urine from visiting fruit bats. One proposed intervention is covering the containers to prevent bat feeding and contamination with bat urine. Proactive wildlife and livestock mortality surveillance, such as great ape Ebola virus carcass surveillance in the Republic of the Congo, could function as an early warning system preceding spillover to the human population.

Challenges to outbreak control. ‘One Health’ approaches involve addressing zoonosis at the human, animal and environmental levels. One of the biggest hurdles to preventing zoonosis at the animal level is the limited feasibility of wildlife vaccination. Given that filoviruses and coronaviruses are likely hosted in a variety of different animal populations, including multiple bat species and other mammals, covering large geographic regions, current vaccination delivery methods are impracticable and likely insufficient to induce effective herd immunity. Although effective vaccines are now in development for Ebola virus, Hendra virus and rabies virus, there are no effective vaccines currently available for both human and animal use. Some progress has been made on this front, for example, in the form of oral vaccines against rabies in dogs and bats as well as against plague in black-tailed prairie dogs. Applying similar efforts to other viruses in lesser-studied and more remote bat populations and other mammals will require a greater understanding of host ecology and behaviour.

After it was discovered that horses are susceptible to Hendra virus and can amplify the virus and infect humans, the Australian government invested in developing a highly effective vaccine that could be given to horses. It was hoped that reducing transmission of the virus to horses would reduce transmission to humans. However, vaccination efforts in Australia have been hindered by antivaccination sentiment, the public perception of the vaccine being too costly for such a rare pathogen and by anecdotal evidence of unwanted side effects. This lack of adoption of Hendra virus vaccination has allowed for sporadic Hendra virus outbreaks in horses to continue, threatening human and animal health.

The geopolitical climate represents an even bigger challenge to outbreak prevention. Despite the existence of multiple, experimental therapeutic options, the latest
Ebola virus outbreak in the Democratic Republic of Congo (ongoing since 2018) has been stymied by civil war breaking down the health-care system and militant groups targeting health-care workers and outbreak response teams146. The recombinant VSV Ebola virus vaccine has been successfully used in response to Ebola virus outbreaks and is now FDA approved. The full licensure of the VSV vaccine will allow for a broader pre-emptive rather than reactive vaccination approach and marks the first licensure of a human vaccine for a bat-borne infectious disease since rabies. Another example of geopolitical disruption was seen during the emergence of SARS-CoV-1 in China in 2002, when the Chinese government delayed reporting the health crisis to the international community well after the outbreak had begun to spread147. The timely release of public health data remains a critical issue during the current COVID-19 pandemic and appears to have improved with the release of case data and full genome viral sequences as the outbreak develops (novel 2019 coronavirus on virological.org).

Conclusions

Disease X, or the as yet unknown pathogen poised to cause the next pandemic, poses a grand challenge in outbreak prevention and response. Bats represent an important but largely uncharacterized source of known human pathogens. Despite a limited understanding of bats, the viruses they carry, and the molecular and ecological forces driving viral spillover, the tools to develop next-generation vaccines and antiviral technologies are maturing such that researchers will be able to respond to the next outbreak with unprecedented speed. In order to transition bat virus research from reactive to predictive with the ability to determine which pathogens represent the greatest threat to global health, vast advancements are needed across a multitude of disciplines (BOX 3). Here, we have identified the important progress made in the past decade and the gaps remaining in our understanding of bat virus ecology, genetic diversity and the molecular mechanisms underlying zoonotic infection and immunity. The emergence and re-emergence of zoonotic bat pathogens demonstrates the inextricable link between the health of humans, animals and the environment. Therefore, efforts to mitigate the public health impacts of bat-borne viruses must integrate research across these disciplines, applying a ‘One Health’ approach, from field to lab, to address the problem. The future of bat virus research lies in a combined and concerted effort to evaluate the molecular and macro-ecological risk factors of transmission, shine light on which viruses carry the potential to spill over and conduct large-scale, longitudinal surveillance studies that will support the deployment and evaluation of next-generation interventions. Nevertheless, the emergence of SARS-CoV-2 continues to present new and sobering challenges. The ongoing COVID-19 pandemic has clearly demonstrated that a dramatic increase in knowledge on pathogen emergence combined with a rapid all-out aggressive response aimed at tracing zoonotic spillover events is needed to prevent the repetition of current events.

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Competing interests

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