

A decade after SARS: strategies for controlling emerging coronaviruses

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Abstract | Two novel coronaviruses have emerged in humans in the twenty-first century: severe acute respiratory syndrome coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus (MERS-CoV), both of which cause acute respiratory distress syndrome (ARDS) and are associated with high mortality rates. There are no clinically approved vaccines or antiviral drugs available for either of these infections; thus, the development of effective therapeutic and preventive strategies that can be readily applied to new emergent strains is a research priority. In this Review, we describe the emergence and identification of novel human coronaviruses over the past 10 years, discuss their key biological features, including tropism and receptor use, and summarize approaches for developing broadly effective vaccines.

Coronaviruses are potentially lethal pathogens, and several novel strains have emerged or have been identified in animal and human populations in the past 10 years. The global severe acute respiratory syndrome coronavirus (SARS-CoV) epidemic was recognized in early 2003 and caused 10–50% mortality in infected individuals, depending on their age^{1–3}. SARS-CoV probably originated in bats, and the search for this reservoir has resulted in the vast expansion of the library of known coronaviruses. Many of these viruses infect various bats and other animal species, and several are phylogenetically similar to known pathogenic human coronaviruses^{4–13}, which suggests that additional emergence events are highly likely to occur. Indeed, recent reports have confirmed the emergence of a novel coronavirus, designated Middle East respiratory syndrome coronavirus (MERS-CoV), which causes ~50% mortality in patients who seek medical attention, is transmissible on close contact and has caused transmission clusters and cases in several countries, including Saudi Arabia, Jordan, Qatar and the United Kingdom¹⁴. Bat species have been implicated as reservoirs of MERS-CoV, but these species are distinct from those that are thought to have been involved in the emergence of SARS-CoV^{6,12,15–18}. Coronaviruses can also be major pathogens in animal populations; for example, porcine epidemic diarrhoea virus (PEDV), which is related to certain bat alphacoronaviruses (FIG. 1), is a major cause of economic loss in the swine industry in Europe and Southeast Asia and has now been detected in US herds ([National Pork Board statement on PEDV](#)).

The recent coronavirus emergence events that are summarized in this article indicate that coronaviruses have the potential to rapidly adapt and stably transmit to new species (see BOX 1 for a discussion on the structure and proteins of coronaviruses). These observations, when paired with an apparently extensive zoonotic reservoir and the propensity of coronaviruses to emerge as highly virulent human pathogens, have spurred the development of animal models to investigate coronavirus replication, pathogenesis and vaccine efficacy. In addition, because coronavirus vaccines have historically exhibited poor capacity for cross-protection¹⁹, the design of methods to generate safe, effective vaccines that can be rapidly implemented during an emerging epidemic is a high priority. In this Review, we summarize the human coronavirus emergence events that have taken place over the past decade, highlight key biological properties that are unique to coronaviruses, discuss the development of animal models for characterizing coronavirus replication, pathogenesis, transmission and vaccine efficacy, and examine the various strategies that have been implemented for the production of safe and effective coronavirus vaccines.

Emergence of novel coronaviruses

Before 2003, two coronaviruses were known to cause human disease: human coronavirus 229E (HCoV-229E) and HCoV-OC43, both of which were identified in the 1960s. HCoV-229E and HCoV-OC43 cause comparatively mild common colds, except in infants, the elderly and the immunocompromised, in whom symptoms can

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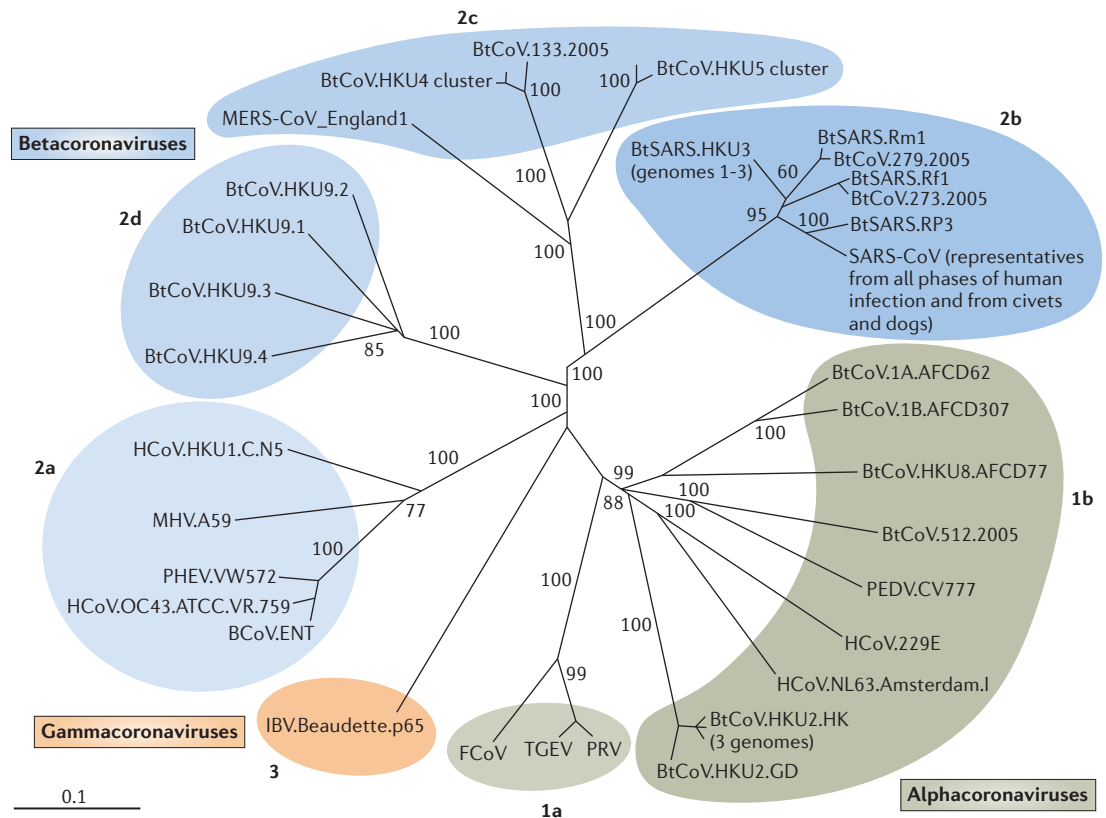


Figure 1 | Whole-genome phylogeny of representative coronaviruses. The full genomic sequences of 50 coronaviruses were aligned and phylogenetically compared. Three distinct phylogenetic groups are shown: alphacoronaviruses (green), betacoronaviruses (blue) and gammacoronaviruses (orange). This taxonomical nomenclature replaced the former group 1, 2 and 3 designations, respectively. Deltacoronaviruses are newly characterized and are not shown. Classic subgroup clusters are marked as 2a–2d for the betacoronaviruses and 1a and 1b for the alphacoronaviruses. The tree was generated using maximum likelihood with the PhyML package. The scale bar represents nucleotide substitutions. Only nodes with bootstrap support above 70% are labelled.

be more severe^{20,21}. However, in early 2003, a previously uncharacterized virus that was associated with the development of SARS, which often progressed to severe lung disease, was isolated from humans. The infected patients exhibited atypical pneumonia that was characterized by diffuse alveolar damage and that had the potential to progress to acute respiratory distress syndrome (ARDS) (BOX 2). SARS was first reported in Guangdong Province, China, and the disease quickly spread worldwide. After an unprecedented global containment effort, the final statistics included more than 8,000 infected individuals and nearly 800 deaths^{1–3,22}. The virus, a lineage B betacoronavirus, was eventually named SARS-CoV and was found to have crossed to humans from zoonotic reservoirs, such as bats¹², Himalayan palm civets (*Paguma larvata*) and raccoon dogs (*Nyctereutes procyonoides*)^{23–25}.

Subsequent to the SARS epidemic, two other coronaviruses that are capable of causing disease in humans, HCoV-NL63 and HCoV-HKU1, were identified from archived nasopharyngeal aspirates^{26–28}. These viruses cause mild to serious lower respiratory tract infections — including croup, bronchiolitis and pneumonia — in infants, children and adults, although the precise disease prevalence and severity, especially in the very young,

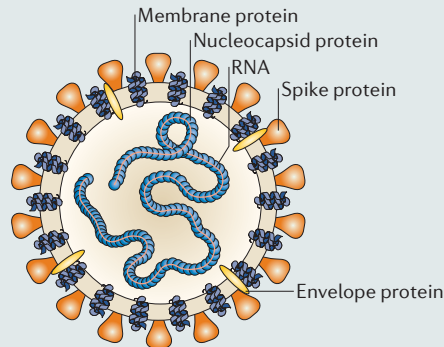
are still being studied²⁹. Although the characterization of these viruses was carried out very recently, molecular clock analyses indicate that HCoV-NL63 probably diverged from its nearest relative, HCoV-229E, around 500–800 years ago; however, as with all molecular clock analyses, these periods of time might be vastly underestimated or over-estimated because of mutation masking and rate changes owing to purifying selection^{7,30,31} (FIG. 1; FIG. 2 (TIMELINE)).

In June and September 2012, two cases of severe infections with another novel coronavirus were identified in the Eastern Mediterranean region^{6,16,17}. Both patients succumbed to severe respiratory illness and, at the time of writing, 145 cases and 62 deaths associated with the novel MERS-CoV have been confirmed by the US Centers for Disease Control and Prevention (CDC MERS-CoV incidence updates). MERS-CoV infection causes atypical pneumonia, ARDS and in some cases renal failure, which are often fatal; however the mortality rate cannot yet be accurately estimated owing to the small number of confirmed cases. Additionally, there is evidence that infection can cause less severe illness in cluster infections (such as those that occur among families and in hospitals) and can even be asymptomatic^{32–34}.

Purifying selection
(Also called negative selection).
A type of natural selection that removes deleterious alleles.

Box 1 | Coronavirus virion structure and proteins

Coronaviruses are enveloped RNA viruses that infect and cause disease in a broad array of animals and humans. Virus particles range from 70 to 120 nm in diameter and are surrounded by characteristic spike-shaped glycoproteins, as shown in the figure. Coronaviruses contain the largest single-stranded, positive-sense RNA genomes currently known, which range from 25.5 to nearly 32 kb in length. Whereas all coronaviruses encode strain-specific accessory genes in ORFs downstream of ORF1 (the replicase–transcriptase), the order of essential structural genes is remarkably well-conserved: all coronaviruses encode the replicase–transcriptase (gene 1), spike, envelope, membrane and nucleocapsid proteins^{17,130,131}. The ~180 kDa spike glycoprotein mediates host cell attachment and entry, utilizing virus- and host-specific cell receptors. The receptor-binding domain (RBD) of the spike glycoprotein is poorly conserved among viruses and, as a result, host receptor usage varies between viral genera and species^{37,38,132}. This variability contributes to the broad host range of coronaviruses, as the viruses can breach cell type, tissue and host species barriers with relative ease^{133–137}. Severe acute respiratory syndrome coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus (MERS-CoV) recognize exopeptidases as receptors for docking and entry, and other human coronaviruses use aminopeptidases or carbohydrates. Crystal structures of both SARS-CoV and MERS-CoV RBDs complexed with their corresponding receptors have been solved^{36,116}. Post-entry events are dependent on one or more cellular proteases, such as cathepsins, human airway trypsin-like protease (HAT) or transmembrane protease serine 2 (TMPRSS2)^{138,139}, which cleave the spike protein and initiate a variety of conformational changes that are important for membrane fusion and entry.



In addition to human infections, high seropositivity for MERS-CoV has been reported in camels, although the exact role of camels in virus transmission and maintenance in populations remains uncertain³⁵. As is the case for SARS-CoV pathogenesis, the elderly seem to be especially vulnerable to poor disease outcomes as a result of MERS-CoV infection, particularly in the presence of co-morbidities such as diabetes, cardiac disease, hypertension and renal disease³⁴. MERS-CoV is a lineage C betacoronavirus and is phylogenetically distinct from all known human coronaviruses, including SARS-CoV. MERS-CoV also uses a receptor that is distinct from that used by SARS-CoV and other human coronaviruses: MERS-CoV uses dipeptidyl peptidase 4 (DPP4), whereas SARS-CoV and HCoV-NL63 have been shown to use angiotensin-converting enzyme 2 (ACE2)^{36–38}. This difference in receptor usage seems to affect tropism, as SARS-CoV infects type I pneumocytes, whereas MERS-CoV infects type II pneumocytes and non-ciliated bronchial cells^{39–41} (TABLE 1). Additionally, MERS-CoV is capable of both animal-to-human and human-to-human transmission: evidence shows that multiple animal-to-human transmission events have occurred in Saudi Arabia in addition to human-to-human transmission^{42,43}, which suggests that MERS-CoV is still undergoing selection in animal and human hosts and will probably continue to do so if the virus is to firmly establish itself in humans. Furthermore, zoonotic isolates that might be

Seropositivity

The positive reaction of a serum sample tested for a pathogen: for example, detection of antibodies against a virus.

Type I pneumocytes

Epithelial cells that line the lung alveoli; type I cells are flat and thin to enable efficient gas exchange.

Type II pneumocytes

Epithelial cells that line the lung alveoli; type II cells are round and produce surfactants.

Synanthropy

The process of an organism, particularly a wild animal, becoming ecologically associated with humans.

related to MERS-CoV have been identified in bat species that are native to the Arabian peninsula, Mexico, Ghana and Europe¹⁵.

Coronavirus reservoirs: the hidden bat virome

Although investigators initially focused on Himalayan palm civets and raccoon dogs as potential reservoirs of SARS-CoV infection, multiple observations suggested that palm civets were opportunistic hosts rather than the primary reservoirs of SARS-CoV-like viruses in the wild. Marketplace civets were disproportionately positive for viral RNA in screening assays⁴⁴, and samples that had been isolated from civets showed ongoing selection, which suggested that the virus was still adapting to the civet rather than persisting in equilibrium in this host^{44,45}. In fact, bioinformatic analysis suggested that there were at least three and possibly more transmission events that occurred between civets and humans: the first occurred before the 2003 epidemic, as indicated by the presence of SARS-CoV antigens in serum samples from uninfected individuals; the second occurred during the main SARS epidemic in 2003; and the third occurred in the winter of 2003–2004 and consisted of a series of sporadic infections^{45,46}. Additionally, molecular analysis of samples that were taken from healthy individuals in Hong Kong in 2001 revealed a prevalence rate of 1.8% for antibodies against SARS-related viruses, which suggests that SARS-CoV or a close ancestor circulated in humans before the 2003 epidemic⁴⁷. In 2005, two groups independently reported the identification of SARS-CoV-related RNA sequences and anti-SARS nucleocapsid antibodies in *Rhinolophus* bats, particularly *Rhinolophus sinicus* and *Rhinolophus macrotis*^{12,13,23}. Interestingly, high antibody titres correlated with low RNA levels, which suggests that active viral replication occurred in these bat species. Genomes from these viruses differ by approximately 10–15% from the 2003 epidemic SARS-CoV, which suggests that viruses with increased homology to SARS-CoV that are capable of using human and bat ACE2 orthologues for docking and entry could exist in bats.

With more than 1,100 species spanning 17 families, it is estimated that bats (order Chiroptera) comprise over 20% of all mammalian species. Their dispersion over much of the globe probably enhances their potential to act as reservoirs for pathogens, some of which are extremely virulent and potentially lethal to other animals and humans, including SARS-CoV^{48,49}. Bats' natural roosting behaviours, in combination with ecological pressures that select for synanthropy, have brought bats closer to humans and to animals that live in proximity to humans, probably increasing the chances of bat-to-animal and bat-to-human transmission^{48,50}. In fact, such a scenario has been proposed for the transmission of SARS-CoV-related coronaviruses from bats to humans: the bats roosted in or near open markets that sold civets in China; viruses were transmitted to civets via faecal and/or oral shedding; and humans handled civets and raw or improperly cooked civet meat, thus completing the transmission chain from bat to human. Alternatively, bats might have directly transmitted the virus to humans

Box 2 | Coronavirus infection and acute respiratory distress syndrome

Acute lung injury (ALI) and acute respiratory distress syndrome (ARDS) are the main causes of respiratory failure and result in high morbidity and mortality in infections involving respiratory viruses, such as H5N1 influenza A, H1N1 2009, severe acute respiratory syndrome coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus (MERS-CoV)^{16,42,140–143}. Radiological examination of SARS-CoV- and MERS-CoV-infected patients reveals atypical pneumonia^{42,128,129,140}. At the tissue level, increased permeability of the alveolar–capillary interface results in pulmonary oedema, hypoxia, type II pneumocyte hyperplasia and cellular infiltration, particularly neutrophil influx. This ‘runaway’ immune response is characterized by the upregulation of pro-inflammatory cytokines, particularly interleukin-1 β (IL-1 β), IL-8 and IL-6, CXCL10 (CXCL10), CC-chemokine ligand 2 (CCL2) and interferon- γ (IFN γ)^{144–147}. This stage of the disease is generally followed by increased fibroproliferation and hyaline membrane formation; depending on the patient, these gross pathological changes may progress or resolve¹⁴⁸. ARDS occurred in approximately 20–30% of SARS-CoV cases and has been noted in MERS-CoV case reports. However, incidence is extremely age-dependent in the case of SARS-CoV infections, and older patients are disproportionately affected: mortality among this patient population exceeded 50% during the epidemic². Among all SARS-CoV-infected patients who developed ARDS, the need for mechanical ventilation was a strong indicator for a high risk of mortality¹⁴⁹. Although no animal models fully recapitulate the symptom profile that is observed in humans infected with SARS-CoV, a few reports have noted the presence of lung injury that resembles ARDS, including in mouse-adapted SARS-CoV-infected signal transducer and activator of transcription 1 (STAT1)-knockout mice and SARS-CoV-infected African green monkeys and, to a lesser extent, cynomolgus macaques^{70,71,86}. Systems biology studies have begun to associate disease with host genetics and have implicated the urokinase pathway, along with other wound-repair pathways, in the development of severe SARS-CoV disease outcome in the ARDS pathway, including diffuse alveolar damage and ALI¹⁵⁰.

via shedding, incidentally co-infecting the civets. There is evidence in support of this hypothesis, as a genetically reconstructed *R. sinicus* bat SARS-CoV-related coronavirus was able to infect primate cells, mouse cells that expressed human ACE2 molecules and mice used to model SARS-CoV infection, and it was able to produce cross-neutralizing antibodies in these mice⁵¹.

Since 2005, dozens of previously unknown bat coronaviruses have been identified; in fact, so many bat coronaviruses are known to be distributed across the *Coronaviridae* family that there is enough evidence to suggest that bat coronaviruses were the ancestral sources of the alphacoronavirus and betacoronavirus genera^{5,9,15,31,52,53} (FIG. 1). Additionally, a recent study reported the transmission of a bat coronavirus between bats of two different suborders, which supports the hypothesis that bats can be co-infected with different coronaviruses and thus, could enable viral molecular recombination⁶. Given that two new human coronaviruses might have emerged from bats and that HCoV-NL63 has been shown to replicate in New World bat cells³¹, priorities for basic research and global health preparedness include determining the amount of virus diversity in bats and other species and, in parallel, mapping the breadth of bat coronavirus host ranges among different bat and animal species.

In addition to SARS-CoV, there is evidence that other human coronaviruses have emerged from bats, including HCoV-229E, HCoV-NL63 and the newly emerged MERS-CoV. MERS-CoV-related viruses (which can be as little as ~1.8% divergent from MERS-CoV) were identified in a recent survey of *Pipistrellus* bats

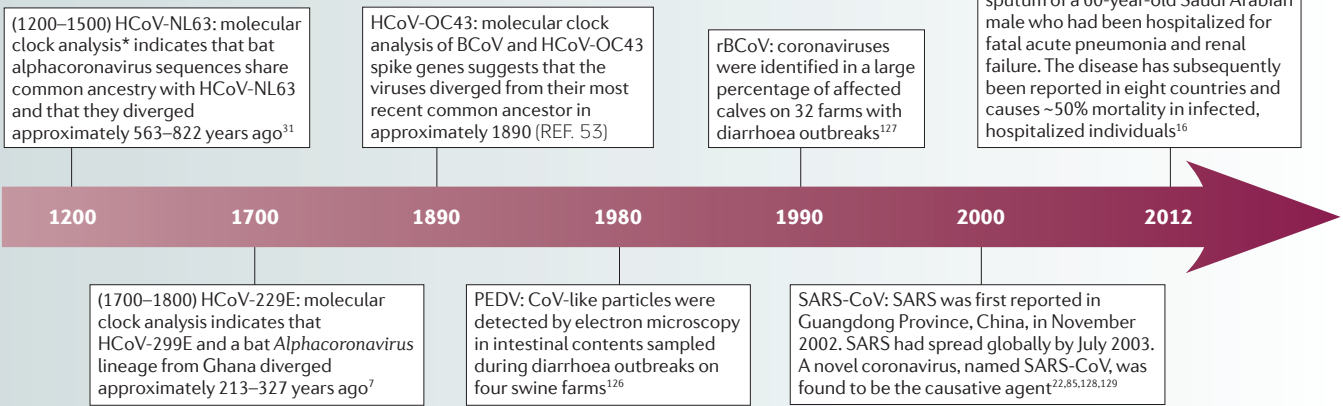
that were captured in Ghana and Europe¹⁵. Bats that were sampled in Ghana also carried viral sequences that are phylogenetically similar to HCoV-229E¹⁵. Viral sequences that were sampled from North American *Perimyotis* bats have phylogenetic similarity to HCoV-NL63 (REF. 31). Additionally, immortalized cells from *Perimyotis* bats are capable of hosting HCoV-NL63 infection, which supports the hypothesis that bats serve not only as reservoirs but also as reverse transmission conduits³¹. Clearly, there is substantial evidence that a broad array of coronaviruses persists in bats and that these viruses have the capacity to recombine and emerge as novel animal and human pathogens. Indeed, MERS-CoV is closely related to both HKU4 (which is a *Tylonycteris* bat coronavirus) and HKU5 (which is a *Pipistrellus* bat coronavirus); all three viruses probably emerged from a common ancestor several centuries ago. Additionally, many polymorphisms have been detected in HKU5 spike protein sequences, which suggests that this virus and, by extension, other bat coronaviruses are capable of generating mutants to occupy new ecological niches when they encounter novel hosts⁵⁴.

Animal models for human coronaviruses

The continuing emergence of virulent human coronaviruses emphasizes the need for animal models for studying viral replication, pathogenesis and transmission. Thus far, this work has largely focused on SARS-CoV infection. SARS-CoV replication has been reported in mice, hamsters, cats, civets and primates, and the most severe disease symptoms have been observed in aged animals. MERS-CoV has a broad host range *in vitro*⁵⁵, which provides some promise for the development of a small animal model for human disease in the near future. However, a functional small animal model for MERS-CoV replication or pathogenesis has not yet been characterized or reported; the possibility of ongoing selection in the receptor-binding sequence in the spike protein or other sequences that are important for host specificity might contribute to this limitation. Inoculation of rhesus macaques with 7×10^6 half-maximal tissue culture infectious doses (TCID₅₀) of a MERS-CoV isolate results in transient mild-to-moderate clinical disease, which includes localized or widespread pneumonia⁵⁶; however, the lack of a small animal model is clearly a major obstacle to furthering our understanding of viral pathogenesis and to testing vaccines and therapeutics.

SARS-CoV mouse models. Epidemic SARS-CoV strains that have been isolated from humans encode receptor-binding domains (RBDs) that interact well with human ACE2. These strains, particularly SARS-CoV Urbani, are capable of interacting with the mouse ACE2 orthologue and of replicating in mouse lungs and small intestine, but disease is limited to mild respiratory symptoms and minimal (<5%) weight loss. However, passage of SARS-CoV Urbani in BALB/c mice produced mouse-adapted variants (SARS-CoV MA15, SARS-CoV MA20 and SARS-CoV v2163) that cause severe and lethal respiratory disease that closely resembles the clinical illness observed in patients and that leads to pneumonitis and

Timeline | **Timeline of coronavirus emergence events**



BCoV, bat coronavirus; HCoV, human coronavirus; MERS-CoV, Middle East respiratory syndrome coronavirus; PEDV, porcine epidemic diarrhea virus; rBCoV, respiratory bat coronavirus; SARS, severe acute respiratory syndrome; SARS-CoV, severe acute respiratory syndrome coronavirus. *Molecular clock analysis estimates the time at which two gene sequences diverged from each other based on known factors, such as mutation rate.

diffuse alveolar damage^{57–59}. In young mice, adaptation requires 6–9 mutations, is focused on specific mutation sites (in the spike and membrane proteins) and takes approximately 15–25 passages; however, in aged mice, mouse-adapted strains emerge within five passages and are defined by fewer mutations, which occur at various locations across the genome. SARS-CoV mouse-adapted strains are significantly more pathogenic in 1-year-old, aged animals, as shown by ~100- to 1,000-fold reductions in median lethal dose values, increased clinical disease severity, extensive ARDS-associated pathological lesions and increased mortality^{19,57,60–63}. Viraemia is rare and transient in mice but is common and long-lasting in patients⁶⁴. Sublethally infected mice develop neutralizing antibody responses, and passive transfer of sera from these mice protects recipients from subsequent lethal challenge. These observations reflect what probably occurred in infected humans during the epidemic: in a cohort study of 128 convalescent human samples, 50% were positive for T cell responses (CD8⁺ T cell responses were more frequent than CD4⁺ T cell responses), and 90% possessed strongly neutralizing antibodies. Of the T cell epitopes that were identified, most were in the spike protein⁶⁵.

Other mouse strains are also susceptible to infection with SARS-CoV and SARS-CoV mouse-adapted strains, most notably 129SvEv-lineage mice and C57BL6 mice. This breadth of susceptibility facilitates the use of genetic knockout and transgenic animals, as most relevant transgenic and knockout lines are available in these backgrounds^{66–69}; for example, myeloid differentiation primary response 88 (MYD88)-knockout and signal transducer and activator of transcription 1 (STAT1)-knockout animals, which both have defects in innate immunity, develop severe disease with weight loss, pneumonitis and bronchiolitis, and die within 9 days of infection. In STAT1-knockout animals, death follows a brief convalescent period and a resurgence of the animal's inflammatory response^{63,70}. By contrast, recombination

activating gene (RAG)-knockout and severe combined immunodeficient (SCID) mice, which have defects in adaptive immunity, become persistently infected after SARS-CoV inoculation and maintain viral titres for at least 30–60 days after infection. There is no evidence of clinical disease during the course of the entire infection, probably owing to a lack of the inflammatory responses that would normally occur as a result of lymphocyte recruitment⁷¹.

Other SARS-CoV animal models. Syrian golden and Chinese hamsters have also been evaluated as models of SARS-CoV disease^{72–75}. Infected animals develop pneumonitis, inflammation and immune cell infiltration, although signs of clinical illness do not necessarily accompany the lung pathology. In contrast to mice, hamsters experience transient (1–2-day) viraemia and become less active. In addition, the virus can be detected in the spleen and liver, although it is not associated with inflammation in these organs. Similarly to mice, hamsters also develop a protective neutralizing antibody response to subsequent SARS-CoV challenge. Reagents for measuring immune responses in the hamster are improving, but there is still limited availability of genetically defined animals and refined immunological and cellular markers for hamster models. TALEN-mediated mutagenesis and CRISPR–Cas-mediated mutagenesis of zygotes might enable the development of genetically defined hamsters and other atypical model species for pathogenesis research.

Ferrets support SARS-CoV replication in both the upper and lower respiratory tracts; however, studies have yielded variable results in terms of other signs of infection. Multifocal lesions develop over 5–10% of the lung surface area and are detected during gross pathological analysis, but clinical symptoms differ between studies^{76,77}. Fever has been reported in SARS-CoV-infected ferrets, in addition to nasal discharge, sneezing and virus shedding; ferrets are the only animals that regularly develop

TALEN-mediated mutagenesis

Mutagenesis mediated by transcription activator-like effector nucleases (TALENs); involves the use of a series of TAL effector (DNA-binding) repeats fused to a *FokI*-specific cleavage domain, which enables site-specific DNA double-strand breaks, non-homologous end joining and repair, which result in hybrid gene construction.

CRISPR–Cas-mediated mutagenesis

Clustered regularly interspaced short palindromic repeats (CRISPRs), which consist of multiple short, conserved nucleotide repeats and function as a bacterial immune system that resists the incorporation of exogenous genetic elements. For mutagenesis, the CRISPR–Cas (CRISPR-associated) system targets foreign DNA with short, complementary single-stranded RNA that directs the Cas9 nuclease to the target DNA, causing double-strand breaks and resulting in silencing of that DNA sequence.

Table 1 | **Coronavirus receptor and co-receptor usage***

Virus	Group	Receptor	Might also bind
Human coronavirus 229E (HCoV-229E)	Alpha	APN	-
Feline coronavirus (FCoV)	Alpha	APN	-
Transmissible gastroenteritis virus (TGEV)	Alpha	APN	Sialic acid
Canine coronavirus (CCoV)	Alpha	APN	-
Bat coronaviruses (BCoVs — multiple species)	Alpha	Unknown	-
Human coronavirus NL63 (HCoV-NL63)	Alpha	ACE2 (REF. 132)	-
Murine hepatitis virus (MHV)	Beta	CEACAM1a	Sialic acid
Severe acute respiratory syndrome coronavirus (SARS-CoV)	Beta	ACE2 (REF. 38)	DC-SIGN, DC-SIGNR, LSECtin
Bat SARS-related coronavirus (Bat-SrCoV)	Beta	ACE2?	-
Middle East respiratory syndrome coronavirus (MERS-CoV)	Beta	DPP4 (REF. 37)	-
Human coronavirus OC43 (HCoV-OC43)	Beta	Unknown	Sialic acid
Avian infectious bronchitis virus (IBV)	Gamma	Unknown	Sialic acid
Bird coronaviruses (multiple species)	Delta	Unknown	-

ACE2, angiotensin converting enzyme 2; APN, aminopeptidase N; CEACAM1a, carcinoembryonic cell adhesion molecule 1a; DC-SIGN, dendritic cell-specific ICAM3-grabbing non-integrin; DC-SIGNR, DC-SIGN-related protein; DPP4, dipeptidyl peptidase 4; LSECtin, liver and lymph node sinusoidal C-type lectin. *Coronavirus receptor usage has been reviewed in REFS 133,151.

fever when infected with SARS-CoV, a symptom that is a hallmark of human infections^{78,79}. Ferrets have not been evaluated as a model for studies of SARS-CoV transmission, but their use might provide interesting information about the genetic basis of the super-spreader events that were observed during the 2002–2003 epidemic. As ferret and human ACE2 molecules probably require different mutations in the SARS-CoV RBD for optimal interactions to occur, it is probable that such ‘transmission-evolved’ ferret-adapted strains would efficiently replicate in ferrets, but less so in humans; however, such studies might highlight residues or regions that have roles in transmission and altered host specificity.

SARS-CoV has also been serially passaged in young F344 rats, which yielded a virus that was capable of enhanced replication and limited pathogenesis in young rats and more severe clinical disease in adult rats. Sequence analysis of the adapted mutant showed that there was a Y442S substitution in the spike protein⁸⁰. *In vitro* passage of SARS-CoV on Chinese hamster ovary (CHO) cells that expressed rat ACE2 yielded a variant that encoded A811S and S950F substitutions in the spike protein⁸¹. Further sequence analysis of rat-adapted variants might help to identify interesting genetic correlates between serially passaged mouse and rat viruses.

Multiple non-human primates (NHPs) are susceptible to SARS-CoV infection, including rhesus macaques, cynomolgus macaques, marmosets and African green monkeys^{82–85}. Clinical symptoms and pathology vary by species but all NHPs that support infection show evidence of SARS-CoV replication in the lungs (with 10^5 – 10^6 TCID₅₀ titres in the lungs) and shedding in respiratory secretions. Most primate models develop histopathological changes and show evidence of pneumocyte infection and some degree of diffuse alveolar damage.

Disease outcomes seem to be more severe in aged macaques⁸⁶, which reflects human disease outcomes. As in humans, increased virus replication and increased lung pathology are noted in primates that are infected with wild-type SARS-CoV compared to most other animal models, except for mice that have been infected with mouse-adapted SARS-CoV. Few vaccine candidates have been evaluated in NHPs, probably owing to the cost and the challenge of achieving statistically significant sample sizes in NHP studies.

Prevention of human coronavirus disease

There are currently no approved antiviral treatments or vaccines for human coronavirus infections, including HCoV-NL63, HCoV-HKU1, SARS-CoV and MERS-CoV. Studies have described small-molecule inhibitors that have the potential to control SARS-CoV infection, and *in vitro* studies have reported that the virus has partial ribavirin and interferon sensitivity at high doses, but clinical management of severe infections (which mostly occur in people infected with SARS-CoV and MERS-CoV) is limited to supportive and palliative care^{14,87,88}. Thus, the development of safe, stable vaccines is necessary, and, because these viruses rise rapidly out of heterogeneous, zoonotic pools, vaccines would ideally be broad-spectrum and rapidly adaptable to new coronaviruses. Of particular importance is the protection of the elderly, as they might be disproportionately susceptible to severe disease from both SARS-CoV and MERS-CoV infections. In support of this hypothesis, the median age of MERS-affected patients who experience disease that is severe enough to require medical attention is currently 50 years (with an age range from 14 months to 94 years) ([WHO MERS-CoV incidence updates](#)). Additionally, in the case of SARS-CoV,

Super-spreader events

Pathogen transmission events that are characterized by the identification of a host (human or animal) that transmits and spreads infection to a significantly greater number of susceptible organisms than the statistical average for that pathogen–host combination.

Table 2 | **Coronavirus vaccine strategies: advantages and disadvantages**

Vaccine strategy	How is it generated?	Advantages	Disadvantages	Constructs and/or loci tested in CoVs
Inactivated virus vaccines	Virions are inactivated with chemicals or radiation	Virion structure maintained (antigenically similar to live virus); easy to prepare; cannot infect, propagate or cause disease; can produce high-titre neutralizing antibodies ^{78,90–92} ; protective when used with adjuvant ^{78,93,94}	Can induce a T _H 2 cell-skewed immune response that leads to inflammatory immune pathology, including eosinophilia and neutrophilia ^{19,95,96}	Whole virus, with or without adjuvant
Live-attenuated virus vaccines	Genomes are mutated using chemical-driven or site-directed mutagenesis	Inexpensive; can be generated rapidly for novel emerging pathogens; cause little or no disease in vaccinees; can elicit T cell- and B cell-dependent immune responses ⁹⁸ ; can induce life-long immunity; site-directed mutations can be tailored to target unique or conserved loci	Phenotypic or genotypic reversion possible; can still cause some disease; generation of a protective immune response with high-titre neutralizing antibodies might depend on sufficient viral replication	Envelope protein deletion ^{99–101} ; non-structural protein 14 (nsp14) and exonuclease (ExoN) inactivation ⁷¹
Viral vector vaccines	An unrelated viral genome lacking packaging elements is engineered to encode the gene of interest	Infect but cannot propagate without helper genes; can induce B cell- and T-cell-dependent immune responses; can generate high-titre neutralizing antibodies (although fewer than those produced by vaccination with inactivated virus)	Intranasal and intramuscular routes of inoculation can produce different immune responses (intranasal inoculation could be necessary to induce mucosal immunity) ⁹⁴ ; protection may not be complete (although the resulting disease is less severe); protection may fail in aged vaccinees ^{66,107,108} ; nucleocapsid protein vaccination produces a T _H 2 cell-skewed immune response ¹⁰⁷	Spike and nucleocapsid proteins ^{66,107,109,111,113,114}
Subunit vaccines	Antigenic components are generated <i>in vitro</i> and harvested for vaccine use	Cannot cause disease from viral infection; can induce B cell- and T cell-dependent immune responses; can generate high-titre neutralizing antibodies ¹¹⁷	<i>In vivo</i> studies have not been completed ^{118,119}	Spike and nucleocapsid proteins ^{36,115,116}
DNA vaccines	Genes that encode antigenic compounds are directly inoculated into vaccinees	Cannot cause disease from viral infection; T _H 1 cell immune profile results from vaccination in combination with inactivated virus ¹²² ; can generate high-titre neutralizing antibodies ^{120,121}	T _H 2 cell-skewed immune response results when used alone; vaccines using nucleocapsid protein might not generate a protective response; vaccines using nucleocapsid protein could result in delayed-type hypersensitivity ^{119,123}	Spike and nucleocapsid proteins ^{120,121}

CoV, coronavirus; T_H, T helper.

the elderly respond poorly to most vaccine formulations^{14,42,89}. Multiple strategies have been used to generate coronavirus vaccines, which include inactivated virus vaccines, live-attenuated virus vaccines, viral vector vaccines, subunit vaccines and DNA or protein vaccines. Most studies have focused only on SARS-CoV vaccine development and have used animal models that do not recapitulate the severe clinical disease that occurs in humans (TABLE 2). To critically evaluate a vaccine, lethal-challenge models using viral strains that are homologous and heterologous to the vaccine strain in both young and aged animals are essential.

Inactivated virus vaccines. Inactivated virus vaccines use chemicals (such as formalin, β-propiolactone and diethylpyrocarbonate) or radiation to render the viral genome non-infectious while maintaining the virion structure, thus preserving antigenicity but eliminating the potential to cause productive infection. Thus, in concept, inactivated virus vaccines are easily prepared and antigenically similar to the live virus. In SARS-CoV research, various studies have shown that inactivated vaccines elicit the

production of neutralizing antibodies^{78,90–92}, and administration of inactivated vaccines with or without adjuvants has been shown to protect against viral replication^{78,93,94}. Vaccination and SARS-CoV challenge in primates and ferrets was reported to favour a T helper 2 (T_H2) cell response that resulted in the production of interleukin-4 (IL-4) and subsequent IL-4-driven inflammatory pathology, rather than the macrophage-driven clearance that is observed in the typical antiviral T_H1 response⁹⁵. One particular inactivated SARS-CoV vaccine, double-inactivated vaccine (DIV), which uses alum as the adjuvant, was protective against challenge with the homologous virus; however, it induced eosinophilia and T_H2 cell immune pathology with poor protection after challenge with a heterologous virus¹⁹. This type of response has been noted with other inactivated, vectored and DNA vaccines (see below). For respiratory syncytial virus, eosinophilic infiltrates are associated with increased mortality and exacerbated lung disease⁹⁶. The eosinophilia that was observed in SARS-CoV-vaccinated animals was not age-dependent but might have promoted hypersensitivity in the presence of other inflammatory

Adjuvants

Pharmacological or immunological substances that modify and enhance the effect of an agent, such as a vaccine.

responses that are enhanced in aged animals, such as neutrophilia¹⁹. Inactivated SARS-CoV vaccines have been administered to humans; the vaccines were well tolerated and induced the production of neutralizing antibodies, although the participants were all relatively young (with an age range between 21 and 40 years) and lung pathology was not assessed. Additionally, in the absence of a natural challenge, no data on vaccine efficacy are available^{78,97}.

Live-attenuated virus vaccines. Live-attenuated vaccines are produced by reducing or eliminating the virulence of a live virus, typically using chemical-driven or site-directed mutagenesis; thus, the virus is capable of productive infection but the resulting disease is either diminished or eliminated. Live-attenuated vaccines can elicit both innate and adaptive immune responses, and protection can be life-long. Additionally, their production is inexpensive⁹⁸. The development of reverse genetics for coronaviruses, particularly SARS-CoV, has greatly simplified the investigation of attenuation-associated alleles and their resistance to reversion in cell culture and in mice and the evaluation of the efficacy of attenuated viruses as vaccine candidates. The envelope protein, which is involved in viral morphogenesis, intracellular trafficking and budding and which might possess ion channel activity, is dispensable for SARS-CoV production; however, replication is attenuated in SARS-CoV lacking this protein⁹⁹. SARS-CoV lacking the envelope protein that was initially constructed in a wild-type rather than mouse-adapted backbone was protective against challenge in hamsters⁹⁹ and partially protective in mice that transgenically express the human ACE2 molecule¹⁰⁰. When the deletion of the envelope protein was engineered in the MA15 backbone, it elicited higher titres of neutralizing antibodies and was protective against challenge in aged mice¹⁰¹. This improvement in vaccine efficacy is probably facilitated by the ~2 log-increased replication of mouse-adapted strains compared to wild-type virus in mice⁵⁹.

All coronaviruses encode an exonuclease (ExoN) in the non-structural protein 14 (nsp14)-coding region of ORF1b. ExoN has been shown to mediate proofreading activity for the viral RNA-dependent RNA polymerase¹⁰². Inactivation of ExoN (Δ ExoN) in SARS-CoV yields viable, replication-attenuated virus that accumulates mutations when passaged¹⁰³. When generated in the MA15 background, the SARS-CoV MA- Δ ExoN virus is attenuated for both growth and pathogenesis in young, aged and immunocompromised mice; it is fully protective against lethal challenge and elicits high levels of neutralizing antibodies, even at low vaccination doses⁷¹. Moreover, SARS-CoV MA- Δ ExoN is stable in terms of both replication and resistance to reversion over short- and long-term passage. Its high resistance to reversion to virulence *in vitro* and *in vivo* might be due to the increased mutation frequency, which is likely to continuously introduce attenuating and/or neutral mutations that discourage and/or prohibit primary (at the site of the original mutation) and secondary (at sites other than the original mutation) reversion to virulence.

The particular advantage of approaches that involve conserved alleles, such as envelope and ExoN, aside from their effects on replication and efficacy in all mouse models tested, is that the conservation of the alleles makes them prime candidates for the rapid generation of new vaccines after the genome sequence of a novel emerging coronavirus (such as MERS-CoV) is known. Further work will focus on stabilizing the attenuated Δ ExoN background and ensuring that it remains resistant to both primary and secondary reversion to virulence and recombination with virulent coronaviruses. For example, all coronaviruses possess a network of fairly well-conserved transcriptional regulatory sequences (TRSs) that are essential for the production of subgenomic mRNAs. Changing the TRS consensus yields viable viruses that have a reduced capacity to recombine with other coronaviruses — a method that shows great promise for rendering vaccine candidates refractory to recombination with naturally occurring (virulent) coronaviruses and for enhancing the safety of using genetically engineered coronaviruses as live vaccines¹⁰⁴. With the recent development of MERS-CoV reverse genetic systems, these approaches can also be tested in the newest emergent coronavirus^{105,106}.

Other vaccine approaches. Viral-vector vaccines, which function as viral gene delivery systems, rely on a host viral genome (for example, adenovirus) that typically lacks the genetic components necessary to produce new virions and that encodes antigenic components of the virus of interest to elicit an immune response. Because viral-vector vaccines persist in the host as genetic material, directly infect antigen-presenting cells and have strong inherent adjuvant activity, they can efficiently induce both innate and B cell- and T cell-mediated immune responses. Adenovirus vectors that express SARS-CoV spike and nucleocapsid proteins, which are the immunodominant coronavirus proteins, yield varying results depending on preparation, the route of administration and the animal model used; however, challenge experiments have not always been performed^{78,93}. Although intramuscular vaccination induces high serum titres of neutralizing antibodies, intranasal inoculation more effectively prevents replication of the challenge virus, which suggests that the intranasal route is more efficient at inducing mucosal immunity⁹⁴. In a side-by-side comparison with inactivated virus, an adenovirus-vectored vaccine produced significantly lower titres of neutralizing antibodies but was protective against subsequent challenge (although not as protective as inactivated virus)⁹⁴. Venezuelan equine encephalitis virus (VEE) viral replicon particles (VRPs) have also been used to express SARS-CoV spike and nucleocapsid proteins. Spike-expressing VRP (VRP-S) vaccines protect against lethal homologous challenge in both young and aged BALB/c mice. When challenged with lethal virus that expresses the heterologous spike protein, only young mice are protected and only for a short time^{66,107,108}. Other viral vectors that use the spike protein, including poxvirus^{109,110}, parainfluenza virus^{111,112}, rabies virus¹¹³ and vesicular stomatitis virus¹¹⁴, confirm

Viral replicon particles (VRPs). Virus-like particles that encode the components required to mediate replication of the genome but that do not encode genes necessary for the production of new virions. Viral replicon particles are typically generated by co-transfecting the replicon genome with helper cassettes that encode structural components, which enables a single round of replication and particle production to occur.

the production of neutralizing antibodies, and some demonstrate protection. Conversely, vaccination with VLP constructs that express SARS-CoV nucleocapsid protein results in a T_H2 cell-skewed immune response and immune pathology in the lungs, with no protection against homologous or heterologous challenge¹⁰⁷.

Similarly to vectored vaccines, subunit vaccines only utilize antigenic components from the virus of interest; in contrast to vectored vaccines, the propagating organism is grown in the laboratory rather than in the vaccinee, and the antigenic component is harvested for use in the vaccine. SARS-CoV subunit vaccines contain either a spike protein fragment (which consists of amino acids 14–762) or the nucleocapsid protein. Because the crystal structures of both the SARS-CoV and MERS-CoV spike protein RBDs have been solved in complex with their corresponding receptors and their interacting residues have been mapped, these domains are the most obvious targets for subunit vaccine design^{36,115,116}. Additionally, subunit spike vaccines can produce higher neutralizing antibody titres than live-attenuated SARS-CoV, poxvirus spike protein or DNA spike protein (DNA-S) vaccination¹¹⁷. Subunit nucleocapsid vaccines induce high levels of B cell- and T cell-mediated immune responses; however, *in vivo* challenge experiments have not yet been performed^{118,119}.

DNA vaccines consist of DNA that encodes the viral antigenic components, and they are directly injected or otherwise inoculated into the vaccinee. As in vectored and subunit vaccines, DNA vaccines also use spike peptides to elicit high titres of neutralizing antibodies^{120,121}. Interestingly, when DNA-S vaccines are used in combination with inactivated vaccines, the cellular immune response is T_H1 cell-directed (whereas vaccination with inactivated virus alone induces a T_H2 cell response)¹²². Similarly to viral nucleocapsid protein vectored vaccines, DNA vaccines that encode the nucleocapsid protein induce strong cell-mediated immunity but are not protective after high-titre challenge; additionally, and unlike spike protein vaccines, DNA-nucleocapsid (DNA-N) vaccines can induce delayed-type hypersensitivity, even in the absence of an antibody response^{119,123}. Importantly, subunit and DNA vaccines have not been rigorously tested in either young or aged animals in lethal-challenge models, so their true *in vivo* efficacy is currently unknown.

Future directions

The emergence of SARS-CoV in 2002 taught us many lessons about zoonotic reservoirs, the importance of identifying animal models that recapitulate the various aspects of human disease and the determinants of vaccine efficacy and safety. This knowledge is being applied to the recent emergence of MERS-CoV in the human population. With its strikingly high morbidity and mortality rates in hospitalized individuals, it is clear that MERS-CoV has the potential to have a profound impact on the human population. However, its low penetrance thus far suggests that the virus might ultimately fail to develop a niche in humans or it might still be adapting to human hosts and that the worst of its effects are yet to

come. Studies with SARS-CoV suggest that MERS-CoV research should focus on establishing animal models that recapitulate replication, pathogenesis and transmission in humans. A priority is to develop treatments to prevent viral-induced immune pathology, particularly in the elderly. Furthermore, vaccines that target conserved alleles and provide broad protection against strains that are both closely and distantly related to the vaccine strain should be developed. One particular concern is the possibility that MERS-CoV vaccines can elicit T_H2 cell immune pathology, as seen with inactivated, vectored and DNA SARS-CoV vaccines. Live-attenuated vaccines and vaccine combinations (for example, inactivated virus combined with DNA vaccines) do not seem to induce this immune phenomenon; thus, these approaches might hold more promise for the development of successful vaccination strategies, particularly in older populations. Adjuvants that promote more robust T_H1 cell responses should also be evaluated in more rigorous models. A second concern is that for SARS-CoV escape from neutralization is driven by spike protein variability; thus, spike-dependent vaccine strategies might require multivalent approaches¹²⁴. For MERS-CoV, the natural variation in the MERS-CoV spike glycoprotein is mostly unknown; thus, it is essential that more natural isolates are recovered and sequenced from humans and reservoir species. Focusing on these priorities will not only help to combat the emergence of MERS-CoV but will also increase our preparedness for any future coronavirus emergence events that originate from the vast, mutable zoonotic reservoir. As such, heterologous SARS-CoV- and MERS-CoV-related isolates that encode even minute variations in the spike protein will be important reagents for evaluating vaccine efficacy against future emerging strains.

Coronavirus phylogeny demonstrates an incredible diversity in antigenic variants, which leads to limited cross-protection against infection with different strains, even within a phylogenetic subcluster. Consequently, the risk of introducing novel coronaviruses into naive human and animal populations remains high. Despite this antigenic breadth, it is revealing that alphacoronaviruses and betacoronaviruses often use receptor orthologues and/or recognize carbohydrates for cross-species transmission. Given the large number of bat species and an ecology that enables potential pathogens to spread between bat populations, a number of bat coronaviruses might be naturally able to recognize human orthologue receptors for docking and entry. A recent study that analysed viral biodiversity in the flying fox *Pteropus giganteus* supports this hypothesis, as coronavirus species were among the several potentially novel viral species that were identified in PCR assays¹²⁵. Moreover, the phylogeny of coronaviruses that have appeared in the human population indicates an accelerating pattern of emergence and disease outbreaks from zoonotic sources. Molecular clock analyses have estimated the dates of the emergence of HCoV-NL63 (~500–800 years ago)³¹, HCoV-229E (~200–300 years ago)⁷, HCoV-OC43 (~120 years ago)⁵³, PEDV (~30 years ago)¹²⁶, respiratory bovine CoV (rBCoV) (~20 years ago)¹²⁷, SARS-CoV

Delayed-type hypersensitivity (Also called type IV hypersensitivity). A CD4⁺ T cell-mediated immunological response rather than an antibody-mediated response. The immune reaction develops 24–72 hours after exposure to an immunogen.

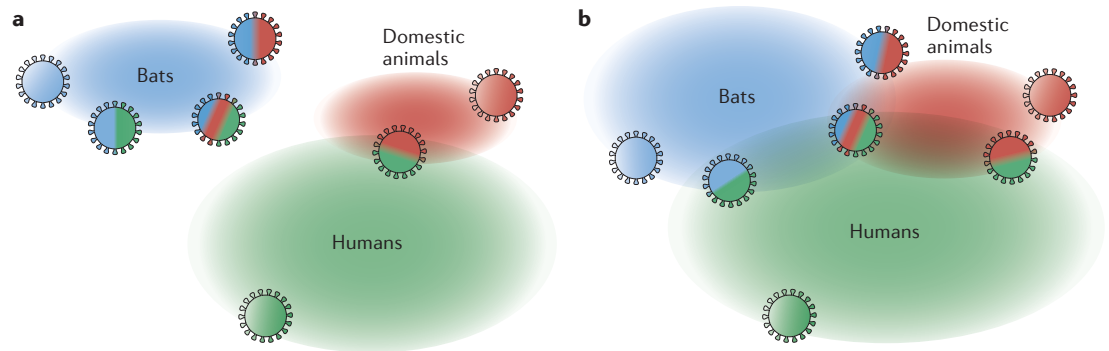


Figure 3 | Changing viral ecology in expanding human and animal populations. Alphacoronaviruses and betacoronaviruses have historically been shown to have the capacity to utilize orthologous receptors and carbohydrates for entry in different species. Thus, in host pools (bats, blue; domestic animals, red; and humans, green), viruses that are capable of cross-species transmission might exist or be generated by the intrinsic error rate of the RNA-dependent RNA polymerase. Bat populations that are largely separated from human and domestic animal populations result in limited viral emergence into new populations (part a). However, the expansion of human populations into previously unsettled areas, intensive domestic animal farm management practices that result in increased herd and flock sizes in small areas and in closer proximity to their human caretakers, and the encroachment of bats into human-populated regions contribute to the increased incidence of pathogenic human viruses from zoonotic pools (part b).

(~10 years ago)^{22,85,128,129} and MERS-CoV (<1 year ago)¹⁶ (FIG. 2), which demonstrates a potential paradigm shift in the relationships that coronaviruses have with domesticated animals and human hosts. Bat coronaviruses have probably evolved to transmit in close proximity and with broader host tropism as a result of the large multispecies bat populations that exist across much of the globe. This pattern of transmission facilitates the spread of coronaviruses to other hosts; for example, intensive farm-management practices result in thousands of animals being housed together in a closed environment. Human population density has increased over the past 100 years, encroaching on wild animal habitats. In addition, the average age of the human population is increasing, and with it the proportion of immunocompromised individuals who are susceptible to severe lung infections. These conditions provide more vulnerable hosts for severe disease and, if animal models are representative of the process in humans, can lead to rapid *in vivo* evolution of virulence and transmissibility (FIG. 3). In fact, most early cases of SARS-CoV and MERS-CoV have been described in the elderly and arose after close contact in hospital and family settings^{42,89}. Emergence of virulence in the SARS-CoV background in older hosts could require fewer mutations and less passage than is required in young hosts⁵⁷. Moreover, if close proximity is an important determinant for efficient coronavirus cross-species transmission and dissemination,

it is possible that the current and projected ecological and demographic conditions are approaching a critical point. The *Coronaviridae* family is poised to excel in these conditions and colonize this growing niche. If these hypotheses are correct, accelerated transmission of bat and animal coronaviruses to humans can be expected to continue and possibly escalate. Public health authorities should prepare by developing broadly applicable platform strategies for the rapid diagnosis, containment and treatment of, and vaccination against, emerging coronavirus infections. We have been warned twice in recent years, and the belief that coronaviruses are highly vulnerable to public health intervention strategies is not supported by the increasing incidence of coronavirus emergence in livestock animal populations and the identification of novel coronaviruses in reservoir species, despite the application of widespread disease management practices and intervention strategies.

Note added in proof

The recent identification of bat SARS-like coronaviruses in Chinese horseshoe bats that are capable of using both the bat ACE2 and the human ACE2 receptor for entry¹⁵² strongly supports the argument that SARS-CoV emerged as a human infection directly from a bat reservoir. These data emphasize the importance of surveying synanthropic wildlife populations for potential zoonotic candidates.

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Competing interests statement

The authors declare no competing interests.

FURTHER INFORMATION

National Pork Board statement on PEDV infections in the US:

<http://www.pork.org/News/3904/NationalPorkBoardStatementOnPEDV.aspx#.UdsYeRZnS-I>

US Centers for Disease Control and Prevention — MERS-CoV incidence updates:

<http://www.cdc.gov/coronavirus/mers/>

World Health Organization — MERS-CoV incidence

updates: http://www.who.int/csr/disease/coronavirus_infections/update_20130813/en/index.html