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Abstract

The recent emergence of bat-borne zoonotic viruses warrants vigilant surveillance in their natural hosts. Of particular concern is the family of coronaviruses, which includes the causative agents of severe acute respiratory syndrome (SARS), Middle East respiratory syndrome (MERS), and most recently, Coronavirus Disease 2019 (COVID-19), an epidemic of acute respiratory illness originating from Wuhan, China in December 2019. Viral detection, discovery, and surveillance activities were undertaken in Myanmar to identify viruses in animals at high risk contact interfaces with people. Free-ranging bats were captured, and rectal and oral swabs and guano samples collected for coronaviral screening using broadly reactive consensus conventional polymerase chain reaction. Sequences from positives were compared to known coronaviruses. Three novel alphacoronaviruses, three novel betacoronaviruses, and one known alphacoronavirus previously identified in other southeast Asian countries were detected for the first time in bats in Myanmar. Ongoing land use change remains a prominent driver of zoonotic disease emergence in Myanmar, bringing humans into ever closer contact with wildlife, and justifying continued surveillance and vigilance at broad scales.

Introduction

Infectious diseases are considered to be “emerging” if they appear in a new population or geographic region or are occurring with greater frequency than the expected background rate [1–3]. Emerging infectious diseases (EIDs) are capable of causing debilitating health effects and financial instability, especially in less developed countries with insufficient capacity to mount health interventions, and thus pose a significant global public health challenge in the 21st
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An estimated 60–75% of EIDs are comprised of zoonotic diseases; of these, more than 70% have purportedly originated in wildlife species [3–5]. Spillover has been largely attributed to changes in anthropogenic activity subsequent to exponential human population growth since the latter half of the 20th century. Large-scale land use change, such as deforestation and land conversion for agriculture, can alter host-pathogen relationships and increase human encounter rates with wildlife and their pathogens, making cross-species transmission events more likely [6,7]. For established pathogens, human-mediated biodiversity loss often leads to reduced populations of suboptimal host species and increased numbers of competent or amplifying hosts, potentially precipitating higher infection rates in people [8]. In addition, intensification of livestock and poultry production systems results in artificially dense populations of domestic animals, which can lead to pathogen amplification and spillover to humans [7]. Approximately two-thirds of human pathogens occupy complex, multi-host systems, and pathogens with multiple animal hosts, including some wildlife species, are more likely to become emergent [9].

Bats are increasingly recognized as the natural reservoirs of viruses of public health concern [10–13]. The capacity of bats to carry and transmit zoonotic pathogens has been hypothesized to be due to their unique life history traits, including their ability for sustained flight, potential for long-distance dispersal, aggregation into densely populous colonies, and adaptation to peri-urban habitats [11,12]. Historically, bats have been linked to highly pathogenic viruses that pose a serious threat to human health, including the coronaviruses responsible for severe acute respiratory syndrome (SARS) and Middle East respiratory syndrome (MERS), the hemorrhagic ebola and Marburg filoviruses, and paramyxoviruses such as Nipah virus [10,11,13–18]. More recently, a pandemic of an acute respiratory syndrome originating in Wuhan, China in December 2019 was linked to a coronavirus (designated “SARS-CoV-2”) that shared 96% identity with a bat-borne coronavirus at the whole-genome level [19]. In some cases, these viruses can subsequently spread through person-to-person contact following spillover from animals, increasing their epidemic potential [10,11,19].

The 2002–2003 SARS epidemic, the emergence of MERS in people in 2012, and the ongoing COVID-19 pandemic have prompted substantial interest in detecting coronaviruses of bat origin due to public health concern and their pandemic potential [10,13–18]. Coronaviruses (CoV) are a family of enveloped, single-stranded RNA viruses that commonly infect the respiratory and gastrointestinal tracts of their mammalian and avian hosts [10]. The Alphacoronaviruses and Betacoronaviruses are of particular importance to human health, with SARS-CoV, SARS-CoV-2, and MERS-CoV—which have caused the most severe disease in humans to date—belonging to the latter group [10,20,21]. Mounting evidence indicates that bats are the evolutionary hosts and origin for these CoV lineages [10,19–22]. In addition to human-associated CoVs, bats are also hosts of coronaviruses that infect production animals, and have been implicated in the emergence and origin of swine acute diarrhea syndrome (SADS), transmissible gastroenteritis virus (TGEV) in pigs, and porcine epidemic diarrhea (PED), which can cause considerable losses [23–26]. Thus, bat-borne CoVs can pose a significant threat to human health and food production.

In spite of these infectious disease threats, bats are an indisputably essential component of ecosystems. They provide critical services such as seed dispersal, pollination, control of insect populations (including crop pests and disease vectors), and fertilization via guano, making them invaluable assets to agricultural industries and small-holder farming [27]. The importance of bats to ecosystems and human communities while being the natural reservoirs of many zoonotic pathogens presents a challenge for disease control. The potential threats posed
by bat-borne coronaviruses to human and livestock health necessitate the identification and characterization of these viruses at high-risk interfaces among humans, domestic animals, and wildlife.

Particular attention is needed in developing regions of high biodiversity, where EIDs are most likely to arise, and where substantial losses in agricultural production may be a source of financial insecurity [28–32]. Myanmar is a particularly vulnerable country due to the interplay of ecological and human factors, which increase opportunities for viral spillover. The nation is situated in the heart of the Southeast Asia region, a hotspot for EIDs, including some neglected tropical diseases and some of pandemic potential like SARS and H5N1 influenza [31,32]. A combination of biological, ecological, socioeconomic, and anthropogenic factors renders the region particularly susceptible to emerging zoonoses that could impart a considerable public health and economic burden [31,32]. Our study aimed to detect coronaviruses in free-ranging bats living in close proximity to human communities.

**Materials and methods**

### Sampling sites

Between May 2016 and August 2018, sample and data collection occurred at three selected sites in Myanmar: 1) Northern District in Yangon Region, near Hlawga National Park (spanning lat: 17.04˚N, 17.50˚N; long: 95.86˚E, 96.12˚E); 2) Hpa-An in Kayin state (spanning lat: 16.66˚N, 16.88˚N; long: 97.58˚E, 97.68˚E); and 3) Shwebo of Sagaing region (lat: 22.37˚N, long: 95.78˚E) (Fig 1). These sites were targeted as potential high-risk human-animal interfaces due to land use change increasing human proximity to wildlife and potential human exposures through livelihood, recreational, commercial, and religious or cultural activities. Two of these sites also featured popular cave systems where people were routinely exposed to bats through guano harvesting, religious practices, and ecotourism. Sites 1 and 2 consisted of several smaller sub-sites where bat capture and sampling events occurred. All surveillance activities were conducted in collaboration with three of Myanmar’s government ministries: (1) the Ministry of Livestock, Agriculture, and Irrigation; (2) the Ministry of Health and Sports; and (3) the Ministry of Natural Resources and Environmental Conservation. All work conducted was approved through a Letter of Agreement, Ethical Review Committee, and Memorandum of Understanding, respectively.

### Animal capture and sampling

Bat sampling was performed by trained field personnel in collaboration with Myanmar’s Ministry of Agriculture, Livestock and Irrigation (MOALI) and Ministry of Natural Resources and Environmental Conservation (MONREC). All bats were captured using mist nets, with each individual manually restrained for species identification, morphometric evaluation, and sample collection. No anesthetic or immobilization agents were used during capture or handling. Oral and rectal swabs were collected when possible using sterile polyester-tipped applicators (animal size often precluded rectal swab collection). Naturally voided guano samples consisting of combined urine and feces were also collected from the environment using plastic tarps. At Site 2, the tarps were placed on the floor of the caves and left overnight, with sample collection occurring the following morning. At Sites 1 and 3, the tarps were placed at cave entrances and under roosting areas in the evening as the bats emerged to forage, and samples were collected immediately. Guano pellets were collected randomly from the tarps and pooled. Tarps were disinfected between each use and gloves were changed in between each sampling event. Pooled guano samples were attributed to a presumptive host species based on field identification of species in caves when possible, otherwise were designated as "Unidentified..."
Chiropterans” when multiple species were present. All sample types were collected into 500 μL viral transport medium (ThermoScientific MicroTest tubes, Fisher Scientific, Pittsburgh, PA, USA) or 500 μL TRIzol reagent (Invitrogen TRIzol reagent, Fisher Scientific, Pittsburgh, PA, USA), transported from the field in liquid nitrogen, and transferred to a -80˚C freezer within five days and stored until time of testing. Bats were humanely trapped, handled, and sampled from according to protocols approved by the Institutional Animal Care and Use Committee of the University of California at Davis (Protocol 19300) and Smithsonian Institution (Protocol 16–05) and with approvals from MOALI and MONREC. Bats were released within 1 km of the

Fig 1. Myanmar study sites. Map of bat capture sites in Myanmar, 2016–2018. Data Sources: Natural Earth. Map created in QGIS 2.18.4. 2020.

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capture site as soon as possible upon completion of each sampling event, with net capture and pillowcase restraint between 5 to 30 minutes and handling times less than 5 minutes for each individual.

**RNA extraction, viral detection, and sequencing**

Sample testing was performed at the UC Davis One Health Institute Laboratory and the Veterinary Diagnostic Laboratory, Livestock, Breeding, and Veterinary Department (LBVD) in Myanmar. 250 μl was used from each sample for RNA extraction per kit instructions, and to ensure availability of an additional aliquot should a second extraction or other downstream analyses be needed. RNA was extracted using Direct-Zol RNA columns (Zymo Research Corp), and 8 μl RNA was used for cDNA transcription using Superscript III (Invitrogen). Sam- ples were screened for coronaviruses using two broadly reactive consensus conventional polymerase chain reaction (PCR) assays targeting two non-overlapping fragments (434 bp and 332 bp) of the RNA-dependent RNA polymerase (RdRp) of orf1ab of CoVs [33,34]. Bands of the expected size were cloned (pCR4-TOPO vector; Invitrogen Corp.) and Sanger sequenced (ABI 3730 Capillary Electrophoresis Genetic Analyzer; Applied Biosystems, Inc., Foster City, CA).

Sequences were analyzed and edited using Geneious Prime (Version 2019.1.3), uploaded to Genbank (S1 Table), and compared with known sequences in the database. Coronavirus sequences were classified as belonging to viral taxa according to established cut-offs and methods [28]. Virus sequences that shared less than 90% identity to a known sequence were labelled sequentially as PREDICT_CoV-1, -2, -3 etc; while groups sharing ≥90% identity to a sequence already in GenBank were given the same name as the matching sequence. Based on these criteria, the CoV sequences detected were assigned to discrete viral taxa. Viral culture and isolation were not attempted for any positive samples.

**Host DNA barcoding**

Bat samples positive for a CoV—including positive pooled guano samples—were barcoded to confirm the host species using PCR assays targeting fragments of the cytochrome B gene (cytB) and the cytochrome oxidase subunit 1 genes (CO1) [35]. One PCR amplicon was selected for sequencing and compared to reference sequences in GenBank using BLAST tools. A threshold of 97% sequence identity was used to confirm the species. Sequences with <95% sequence identity were classified to the genus. DNA barcoding was also performed on a subset of the CoV-negative pooled guano samples. Pooled guano samples were assigned a presumptive origin species based on host barcoding.

**Results**

A total of 464 bats representing at least 11 species across eight genera from six families were captured and sampled (Table 1). Both insectivorous microbats and fruit bats were represented in our study population. A total of 759 samples were collected and tested (464 oral swabs, 140 rectal swabs, 155 guano samples). A total of 461 samples were collected in the dry-season sampling (244 oral swabs, 117 rectal swabs, and 100 guano samples) and 298 samples (220 oral swabs, 23 rectal swabs, and 55 guano samples) in the wet season.

CoVs were detected in 48 samples: one oral swab and seven rectal swabs from seven individual bats and 40 pooled guano samples (Table 1). Viral fragments were detected from one unidentified tomb bat (Taphozous sp.), three Horsfield’s leaf-nosed bats (Hipposideros larvatus), and three greater Asiatic yellow house bats (Scotophilus heathii). Thirty-six of the 40 positives detected in guano were attributed to H. larvatus, while the host species for the remaining four positive pooled guano samples was identified as wrinkle-lipped free-tailed bats...
Overall viral prevalence across all bat taxa and all coronaviral genotypes was approximately 1.5%. The vast majority of positive detections (83.3%) were made from pooled guano samples, while oral swabs had the lowest yields. Positive detections were made from 40 samples collected during the dry season (83.3%), while wet-season sampling resulted in positive detections from eight samples (16.7%). Both Sites 1 and 2 accounted for positive detections, while no coronaviral sequences were detected at Site 3.

Fifty-four total sequences were recovered, clustering within seven distinct coronaviral genotypes. Using established cut-offs and methods [28], we detected four alpha coronaviruses (PREDICT_CoV-35, 47, 82, and 90) and three betacoronaviruses (PREDICT CoV-92, 93, and 96). Of these, the alphacoronavirus PREDICT_CoV-35 was previously known, having been found in Scotophilus kuhlii, unidentified Myotis, and other unspeciated host bats in the neighboring countries of Cambodia and Vietnam from 2013 to 2017 [36]. The remaining six coronaviruses were novel (three alphacoronaviruses and three betacoronaviruses).

PREDICT_CoV-92 was the most commonly detected coronavirus, found in 36 pooled guano samples attributed to H. larvatus (Table 1). Interestingly, three coronaviruses were only found as co-infections: PREDICT_CoV-90 was detected with PREDICT_CoV-35, PREDICT_CoV-93 with -96, and PREDICT_CoV-96 also with -92.

### Table 1. Summary of positives and coronaviruses detected in bats in Myanmar.

| Taxonomic Level | Common Name | Individual Bats | Rectal Swab | Oral Swab | Pooled Guano | Samples | CoVs Detected |
|-----------------|-------------|----------------|-------------|-----------|-------------|---------|---------------|
| Vespertilioniformes |             |                |             |           |             |         |               |
| Vespertilionidae |             |                |             |           |             |         |               |
| Scotophilus heathii | Greater Asiatic yellow house bat | 3/12\(^1\) | 3/12 | 0/12 | 0/0 | 3/24 | PREDICT_CoV-35, 90\(^3\) |
| Scotophilus kuhlii | Lesser Asiatic yellow house bat | 0/1 | 0/1 | 0/1 | 0/0 | 0/2 |               |
| Emballonuridae     |             |                |             |           |             |         |               |
| Taphozous sp.\(^4\) |             | 1/3 | 1/3 | 0/3 | 0/0 | 1/6 | PREDICT_CoV-35\(^2\) |
| Molossidae         |             |                |             |           |             |         |               |
| Chaerephon plicatus | Wrinkle-lipped free-tailed bat | 0/219 | 0/65 | 0/219 | 4/105 | 4/389 | PREDICT_CoV-47,82 |
| Pteropodiformes    |             |                |             |           |             |         |               |
| Hipposideridae     |             |                |             |           |             |         |               |
| Hipposideros armiger | Great Himalayan leaf-nosed bat | 0/17 | 0/0 | 0/17 | 0/0 | 0/17 |               |
| Hipposideros larvatus | Horsfield’s leaf-nosed bat | 3/81 | 3/16 | 1/81 | 36/50 | 40/147 | PREDICT_CoV-92,93,96\(^3\) |
| Craseonycteridae    |             |                |             |           |             |         |               |
| Craseonycteris thonglongyai | Kitti’s hog-nosed bat | 0/31 | 0/9 | 0/31 | 0/0 | 0/40 |               |
| Pteropodidae       |             |                |             |           |             |         |               |
| Eonycteris spelaea | Lesser dawn bat | 0/33 | 0/0 | 0/33 | 0/0 | 0/33 |               |
| Cynopterus sphinx   | Greater short-nosed fruit bat | 0/38 | 0/5 | 0/38 | 0/0 | 0/43 |               |
| Pteropus giganteus  | Indian flying fox | 0/29 | 0/29 | 0/29 | 0/0 | 0/58 |               |
| Total              |             | 7/464 | 7/140 | 1/464 | 40/155 | 48/759 |               |

\(^1\)Includes Scotophilus cf. heathii based on 95–97% shared nt identity with reference sequences.

\(^2\)Virus previously discovered during PREDICT-1 surveillance activities.

\(^3\)Indicates at least one instance of co-infection.

\(^4\)Did not meet the 95% nt identity threshold for identification to the taxonomic level of species.

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Discussion

Three new alphacoronaviruses, three new betacoronaviruses, and one previously described alphacoronavirus were detected in bats in Myanmar. None of the viruses appeared to be closely related to SARS-CoV, MERS-CoV, or SARS-CoV-2. Guano samples accounted for the majority of positives, suggestive of an important transmission route for CoV shedding from bats [29,28,29] and a possible risk to people during the act of guano harvesting [37,38]. Viral detection in guano also has implications for future surveillance, as our study demonstrates the value of non-invasive collection of guano for viral surveillance, potentially obviating the need for handling individual bats for coronaviral detection. Our findings supplement those of He et al., who profiled the virome of insectivorous bats from northern Myanmar but did not detect coronaviruses in that study [40].

A difference was found in positives for CoV by species, as samples from *H. larvatus* represented 83% of positives. A wide diversity of CoVs has been found in Hipposiderid bats [28,34,39,41], and our study is consistent with those findings. Four CoVs detected in our surveillance study were found in a single host species each: PREDICT_CoV-90 was found only in *S. heathii*; and PREDICT_CoV-92, -93, and -96 were found only in *H. larvatus* (Table 1). These findings may possibly suggest limited host-switching and viral sharing for certain viruses within our study populations, a pattern consistent with prior observations that viral groups are likely significantly associated with host taxa at the family level [28]. However, further evidence is needed to elucidate host-viral relationships and ecology in the region.

Our findings also likely reflect a bias in our sampling effort. Although *H. larvatus* samples accounted for the most positives, these were largely detected in guano samples collected from the environment, as individuals were not frequently caught by mist net. Overall in our study, the numbers of individual bats handled and sampled per species were relatively low, ranging from one to 218 (Table 1). Viral prevalence may vary widely with the species of host and pathogen. Anthony et al. suggested a sample size of at least 154 individuals per species in order to maximize our ability to detect CoVs. Targeting more host species, specific taxa (Hipposideridae), and larger sample sizes might have improved our detection rate in the species where no CoVs were found [28,29].

Currently, active pathogen surveillance at human-wildlife interfaces in Myanmar is limited. Despite relatively small sample sizes, our study detected several coronaviruses in insectivorous bats, suggesting that more may remain to be uncovered. Given the potential consequences for public health in light of expanding human activity, continued surveillance for coronaviruses is warranted, especially in other species and human-wildlife interfaces. Anthony et al. estimated that over 3,200 CoVs occur in bats, most of which remain undiscovered [28]. Enhancing our sampling effort to incorporate more diverse bat families and larger sample sizes may enable us to identify more CoVs in bats in Myanmar. Additionally, because only short fragments of the conserved RdRp gene (328 bp and 434 bp) were amplified in this study, protein sequence and phylogenetic analyses were not pursued, and identification of recombination events was not possible. While this is an inherent shortcoming of our methodology, the purpose of this study was not to fully characterize specific viruses, but to broadly screen for viruses in bats living in proximity to human communities to better understand potential sources of zoonotic transmission in the context of these human-wildlife interfaces. Further studies may consider complete genomic sequencing for more comprehensive profiling of the bat viromes in this ecosystem. In particular, evaluation of the spike gene sequences may provide insights into host range, including potential viral host-sharing or host-switching events [42].

Land use change will likely continue bringing people into closer proximity with bats, raising encounter rates and opportunities for spillover, facilitating the emergence of zoonotic viruses,
and supporting the need for surveillance [12,43]. Historically, human activities have arguably played a significant role in interspecies transmission events. Following the SARS outbreak, coronaviruses have since been detected in numerous bat species globally, including in Asia, Africa, Europe, the Americas, and the Australasian region [28,44–49]. Mounting evidence supports the role of bats in the transmission of viruses of public health concern—including SARS-CoV and MERS-CoV—and the zoonotic potential of unknown bat-borne coronaviruses warrants vigilant, continued surveillance [10]. Understanding their ecology and prevalence in their natural hosts can improve our ability to detect, prevent, and respond to potential public health threats. Finally, given the essential ecosystem services provided by bats, public health efforts should advocate for preventative measures to protect people against disease transmission while enabling human communities and bats to coexist on a shared landscape.

Supporting information

S1 Table. Final edited sequences and genbank accession numbers. (XLSX)

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**References**

1. Morens DM, Folkers GF, Fauci AS. The challenge of emerging and reemerging infectious diseases. Nature. 2004; 468: 647–652. [https://doi.org/10.1038/nature02759]
2. Morse SS. Factors in the emergence of infectious diseases. Emerg Infect Dis. 1995; 1: 7–15. [https://doi.org/10.3201/eid0101.950102] PMID: 8903148
3. Woolhouse MEJ, Gowtage-Sequeria S. Host range and emerging and reemerging pathogens. Emerg Infect Dis. 2005; 11: 1842–1847. [https://doi.org/10.3201/eid1112.050997] PMID: 16485468
4. Jones KE, Patel NG, Levy MA, Storeygard A, Balk D, Gittleman JL, et al. Global trends in emerging infectious diseases. Nature. 2008; 451: 990–993. [https://doi.org/10.1038/nature06536] PMID: 18288193
5. Taylor LH, Latham SM, Woolhouse MEJ. Risk factors for human disease emergence. Phil Trans R Soc Lond B. 2001; 356: 983–989. [https://doi.org/10.1098/rstb.2001.0888] PMID: 11516376
6. Murray KA, Daszak P. Human ecology in pathogenic landscapes: two hypotheses on how land use change drives viral emergence. Curr Opin Virol. 2013; 3: 79–83. [https://doi.org/10.1016/j.coiviro.2013.01.006] PMID: 23415415
7. Karesh WB, Dobson A, Lloyd-Smith JO, Lubroth J, Dixon MA, Bennett M, et al. Ecology of zoonoses; natural and unnatural histories. Lancet. 2012; 380: 1936–1945. [https://doi.org/10.1016/S0140-6736(12)6178-X] PMID: 23200502
8. Keesing F, Belden LK, Daszak P, Dobson A, Harvell CD, Holt RD, et al. Impacts of biodiversity on the emergence and transmission of infectious diseases. Nature. 2010; 468: 647–652. [https://doi.org/10.1038/nature09575] PMID: 21124449
9. Cleaveland S, Laurenson MK, Taylor LH. Diseases of humans and their domestic mammals: pathogen characteristics, host range and the risk of emergence. Phil Trans R Soc Lond B. 2001; 356: 991–999. [https://doi.org/10.1098/rstb.2001.0889] PMID: 11516377
10. Hu B, Ge X, Wang L, Shi Z. Bat origin of human coronaviruses. Virol J. 2015; 12: 1–8. [https://doi.org/10.1186/s12985-014-0235-7]
11. Calisher CH, Childs JE, Field HE, Holmes KV, Schountz T. Bats: Important reservoir hosts of emerging viruses. Clin Microbiol Rev. 2006; 19: 531–545. [https://doi.org/10.1128/CMR.00017-06] PMID: 16847084
12. Olival KJ, Hosseini PR, Zambrana-Torrelio C, Ross N, Bogich TL, Daszak P. Host and viral traits predict zoonotic spillover from mammals. Nature. 2017; 546: 646–650. [https://doi.org/10.1038/nature22975] PMID: 28636590
13. Han HJ, Wen HL, Zhou CM, Chen FF, Luo LM, Liu JW, et al. Bats as reservoirs of severe emerging infectious diseases. Virus Res. 2015; 205: 1–6. [https://doi.org/10.1016/j.virusres.2015.05.006] PMID: 25997928
14. Ge XY, Li JL, Yang XL, Chmura AA, Zhu G, Epstein JH, et al. Isolation and characterization of a bat SARS-like coronavirus that uses the ACE2 receptor. Nature. 2013; 503: 535–538. [https://doi.org/10.1038/nature12711] PMID: 24172901
15. Li W, Shi Z, Yu M, Ren W, Smith C, Epstein JH, et al. Bats are natural reservoirs of SARS-like coronaviruses. Science. 2005; 310: 676–679. [https://doi.org/10.1126/science.1118391] PMID: 16195424
16. Lau SK, Woo PC, Li KS, Huang Y, Tsoi HW, Wong BH, et al. Severe acute respiratory syndrome coronavirus-like virus in Chinese horseshoe bats. Proc Natl Acad Sci USA. 2005; 102: 14040–14045. [https://doi.org/10.1073/pnas.0506735102] PMID: 16169905
17. Memish ZA, Mishra N, Olival KJ, Fagbo SF, Kapoor V, Epstein JH, et al. Middle East respiratory syndrome coronavirus in bats, Saudi Arabia. Emerg Infect Dis. 2013; 19: 1819–1823. [https://doi.org/10.3201/eid1911.131172] PMID: 24206838
18. Anthony SJ, Gilardi K, Menachery VD, Goldstein T, Ssebide B, Mbabazi R, et al. Further evidence for bats as the evolutionary source of Middle East respiratory syndrome coronavirus. mBio. 2017; 8: 1–13. [https://doi.org/10.1128/mBio.00373-17] PMID: 28377531
19. Zhou P, Yang X-L, Wang X-G, Hu B, Zhang L, Zhang W, et al. A pneumonia outbreak associated with a new coronavirus of probable bat origin. Nature. 2020. [https://doi.org/10.1038/s41586-020-2012-7] PMID: 32015507
20. Wong ACP, Li X, Lau SKP, Woo PCY. Global epidemiology of bat coronaviruses. Viruses. 2019; 11: 174. [https://doi.org/10.3390/v11020174] PMID: 30791586
21. Banerjee A, Kulcsar K, Misra V, Frieman M, Mossman K. Bats and coronaviruses. Viruses. 2019; 11: 41. [https://doi.org/10.3390/v11010041] PMID: 30633496
22. Woo PCY, Lau SKP, Lam CF, Lau CCY, Tsang AKL, Lau JHN, et al. Discovery of seven novel mammalian and avian coronaviruses in the genus Deltacoronavirus supports bat coronaviruses as the gene source of Alphacoronavirus and Betacoronavirus and avian coronaviruses as the gene source of Gamma-coronavirus and Deltacoronavirus. J Virol. 2012; 86: 3995–4008. [https://doi.org/10.1128/JVI.06540-11] PMID: 22278237
23. Zhou P, Fan H, Lan T, Yang XL, Shi WF, Zhang YZ, et al. Fatal swine acute diarrhea syndrome caused by an HKU2-related coronavirus of bat origin. Nature. 2018; 556: 255–258. [https://doi.org/10.1038/s41586-018-0010-9] PMID: 29618817
24. Gong L, Li J, Zhou Q, Xu Z, Chen L, Zhang Y, et al. A new bat-HKU2-like coronavirus in swine, China, 2017. Emerg Infect Dis. 2017; 23: 1607–1609. [https://doi.org/10.3201/eid2309.170915] PMID: 28654418
25. Tang XC, Zhang JZ, Zhang SY, Wang P, Fan XH, Li LF, et al. Prevalence and genetic diversity of coronaviruses in bats from China. J Virol. 2006; 80: 7481–7490. [https://doi.org/10.1128/JVI.00697-06] PMID: 16840328
26. Marques Simas PV, de Souza Barnabe AC, Durães-Carvalho R, de Lima Neto DF, Cardia Caserta L, Artacho L, et al. Bat coronavirus in Brazil related to Appalachian Ridge and porcine epidemic diarrhea viruses. Emerg Infect Dis. 2015; 21: 729–731. [https://doi.org/10.3201/eid2104.141783] PMID: 25811911
27. Kunz TH, Braun de Torrez E, Bauer D, Lobova T, Fleming TH. Ecosystem services provided by bats. Ann N.Y. Acad Sci. 2011; 1223:1–38. [https://doi.org/10.1111/j.1749-6632.2011.06004.x] PMID: 21449963
28. Anthony SJ, Johnson CK, Greig DJ, Kramer S, Che X, Wells H, et al. Global patterns in coronavirus diversity. Virus Evol. 2017; 3: 1–15. [https://doi.org/10.1093/verx/vex012] PMID: 28630747
29. Young CCW, Olival KJ. Optimizing viral discovery in bats. PLoS One 2016; 11: 1–18. [https://doi.org/10.1371/journal.pone.0149237] PMID: 2867024
30. Allen T, Murray KA, Zambrana-Torrelo C, Morse SS, Rondinini C, Di Marco M, et al. Global hotspots and correlates of emerging zoonotic diseases. Nat Commun. 2017; 8: 110. [https://doi.org/10.1038/s41586-017-00114-5] PMID: 28377531
31. Coker RJ, Hunter BM, Rudge JW, Liverani M, Hanvoravongchai P. Emerging infectious diseases in southeast Asia: regional challenges to control. Lancet. 2011; 377: 599–609. [https://doi.org/10.1016/S0140-6736(10)62004-1] PMID: 21269678
32. Bordier M, Roger F. Zoonoses in South-East Asia: a regional burden, a global threat. Anim Health Res Rev. 2013; 14: 40–67. [https://doi.org/10.1017/S1466252313000017] PMID: 23506700
33. Watanabe S, Masangkay JS, Nagata N, Morikawa S, Mizutani T, Fukushi S, et al. Bat coronaviruses and experimental infection of bats, the Philippines. Emerg Infect Dis. 2010; 16: 1217–1223. [https://doi.org/10.3201/eid1608.100208] PMID: 20678314
34. Quan PL, Firth C, Street C, Henriquez JA, Petrosov A, Tashmukhamedova A, et al. Identification of a severe acute respiratory syndrome coronavirus-like virus in a leaf-nosed bat in Nigeria. mBio. 2010; 1: 1–9. [https://doi.org/10.1128/mBio.00208-10] PMID: 21063474
35. Townzen JS, Brower AVZ, Judd DD. Identification of mosquito bloodmeals using mitochondrial cytochrome oxidase subunit I and cytochrome b gene sequences. Med Vet Entomol. 2008; 22: 286–292. [https://doi.org/10.1111/j.1365-2915.2008.00760.x] PMID: 19120966
36. USAID PREDICT. PREDICT 1 & 2 Test Data [cited 2019 Oct 30]. HealthMap Database [Internet]. Available from: <www.data.predict.global>.
37. Wacharapluesadee S, Sintunawa C, Kaewpom T, Khongnomnan K, Olival KJ, Epstein JH, et al. Group C betacoronavirus in bat guano fertilizer, Thailand. Emerg Infect Dis. 2013; 19: 1349–1351. [https://doi.org/10.3201/eid1908.130119] PMID: 23880503
38. Kudagamanna HDWS, Thevanesam V, Chu DKW, Eriyagama NB, Peiris JSM, Noordeen F. Coronaviruses in guano from Pteropus medius bats in Peradeniya, Sri Lanka. Transbound Emerg Dis. 2018; 65: 1122–1124. [https://doi.org/10.1111/tbed.12851] PMID: 29498228

39. Wacharapluesadee S, Duengkae P, Rodpan A, Kaewpom T, Maneenorn P, Kanchasakha B, et al. Diversity of coronavirus in bats from Eastern Thailand. Virol J. 2015; 12: 1–7. [https://doi.org/10.1186/s12985-014-0235-7]

40. He B, Li Z, Yang F, Zheng J, Feng Y, Guo H, et al. Virome profiling of bats from Myanmar by metagenomic analysis of tissue samples reveals more novel mammalian viruses. PLoS One. 2013; 8: e61950. [https://doi.org/10.1371/journal.pone.0061950] PMID: 23630620

41. Bourgarel M, Pfukenyi DM, Boué V, Talignani L, Chiweshe N, Diop F, et al. Circulation of Alphacoronavirus, Betacoronavirus and Paramyxovirus in Hipposideros bat species in Zimbabwe. Infect Genet Evol. 2018; 58: 253–257. [https://doi.org/10.1016/j.meegid.2018.01.007] PMID: 29331670

42. Graham RL, Bric RS. Recombination, reservoirs, and the modular spike: mechanisms of coronavirus cross-species transmission. J Virol. 2010; 84: 3134–3146. [https://doi.org/10.1128/JVI.01394-09] PMID: 19906932

43. Afelt A, Frutos R, Devaux C. Bats, coronaviruses, and deforestation: Toward the emergence of novel infectious diseases? Front Microbiol. 2018; 9: 1–5. [https://doi.org/10.3389/fmicb.2018.00001]

44. Dominguez SR, O'Shea TJ, Oko LM, Homes KV. Detection of group 1 coronaviruses in bats in North America. Emerg Infect Dis. 2007; 13: 1295–1300. [https://doi.org/10.3201/eid1309.070491] PMID: 18252098

45. Tong S, Conrardy C, Ruone S, Kuzmin IV, Guo X, Tao Y, et al. Detection of novel SARS-like and other coronaviruses in bats from Kenya. Emerg Infect Dis. 2009; 15: 482–485. [https://doi.org/10.3201/eid1503.081013] PMID: 19239771

46. Gloza-Rausch F, Ipsen A, Seebens A, Göttliche M, Panning M, Drexlter JF, et al. Detection and prevalence patterns of group I coronaviruses in bats, Northern Germany. Emerg Infect Dis. 2008; 14: 626–631. [https://doi.org/10.3201/eid1404.071439] PMID: 18400147

47. Smith CS, de Jong CE, Meers J, Henning J, Wang LF, Field HE. Coronavirus infection and diversity in bats in the Australasian region. EcoHealth. 2016; 13: 7282. [https://doi.org/10.1007/s10393-016-1116-x] PMID: 27048154

48. Anthony SJ, Ojeda-Flores R, Rico-Chávez O, Navarrete-Marcias I, Zambrana-Torrello CM, Rostal MK, et al. Coronaviruses in bats from Mexico. J Gen Virol. 2013; 94: 1028–1038. [https://doi.org/10.1099/vir.0.049759-0] PMID: 23364191

49. Lelli D, Papetti A, Sabelli C, Rosti E, Moreno A, Boniotti MB. Detection of coronaviruses in bats of various species in Italy. Viruses. 2013; 5: 2679–2689. [https://doi.org/10.3390/v5112679] PMID: 24184965