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Coronaviruses and Australian bats: a review in the midst of a pandemic

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Abstract. Australia's 81 bat species play vital ecological and economic roles via suppression of insect pests and maintenance of native forests through pollination and seed dispersal. Bats also host a wide diversity of coronaviruses globally, including several viral species that are closely related to SARS-CoV-2 and other emergent human respiratory coronaviruses. Although there are hundreds of studies of bat coronaviruses globally, there are only three studies of bat coronaviruses in Australian bat species, and no systematic studies of drivers of shedding. These limited studies have identified two betacoronaviruses and seven alphacoronaviruses, but less than half of Australian species are included in these studies and further research is therefore needed. There is no current evidence of spillover of coronaviruses from bats to humans in Australia, either directly or indirectly via intermediate hosts. The limited available data are inadequate to determine whether this lack of evidence indicates that spillover does not occur or occurs but is undetected. Conversely, multiple international agencies have flagged the potential transmission of human coronaviruses (including SARS CoV-2) from humans to bats, and the consequent threat to bat conservation and human health. Australia has a long history of bat research across a broad range of ecological and associated disciplines, as well as expertise in viral spillover from bats. This strong foundation is an ideal platform for developing integrative approaches to understanding bat health and sustainable protection of human health.

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Introduction

The global COVID-19 pandemic resulting from the emergence of a novel coronavirus, SARS-CoV-2 in late 2019 (Zhu et al. 2020) has refocussed attention on coronaviruses of bat origin. Bats host a wide diversity of coronaviruses (Cui et al. 2007; Vijaykrishna et al. 2007; Drexler et al. 2014; Anthony et al. 2017a; Lacroix et al. 2017; Wong et al. 2019; Latinne et al. 2020), including viral species linked to emergent human respiratory syndrome coronaviruses (Cui et al. 2019). SARS-CoV-2 has close nucleotide sequence identity with coronaviruses isolated from bats in Yunnan Province in China (Hu et al. 2017; Zhou et al. 2020). The epidemiological link between early COVID-19 cases and a Wuhan wet market (Chen et al. 2020; Li et al. 2020) led to an initial hypothesis that transmission from bats and recombination in an unknown intermediate host sold in the market might be involved (Zhang et al. 2020); however, the links to intermediate hosts are unclear (Boni et al. 2020), and it is possible that the market was the site of a superspreading event after the emergence of SARS-CoV-2 elsewhere. This scenario parallels the emergence of severe acute respiratory syndrome (SARS) in China in 2002 (Ksiazek et al. 2003); the search for the origins of the aetiological coronavirus (SARS-CoV) of that pandemic (Guan et al. 2003; Tu et al. 2004; Kan et al. 2005; Poon et al. 2005; Tang et al. 2006) and the cluster of SARS-related betacoronaviruses subsequently described in Rhinolophus species in China (Lau et al. 2005; Li et al. 2005) provides the background for the current focus on bats in the search for the origins of SARS-CoV-2.

Australia has a diverse bat community, with 81 species representing both suborders and 9 families (Reardon *et al.* 2015). The majority of Australian species are smaller insectivorous bats with little public profile and generally limited public contact opportunity. In contrast, large, arboreal, colonially roosting

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pteropodid bats (commonly known as flying-foxes) are regularly present, and prominent, in eastern Australian cities and towns. This bat community has been the focus of a diverse range of research studies, including their genetic and evolutionary history (e.g. Godthelp *et al.* 1992; Webb and Tidemann 1996), ecology (e.g. McKenzie and Rolfe 1986; Lumsden and Bennett 2005; Eby and Law 2008), physiology (e.g. Hosken and Withers 1997; Welbergen *et al.* 2008), immunology (e.g. Zhang *et al.* 2013; Zhou *et al.* 2016; Schountz *et al.* 2017), and infectious disease emergence (Gould *et al.* 1998; Halpin *et al.* 2000; Barr *et al.* 2012; Kessler *et al.* 2018) and dynamics (e.g. Field *et al.* 2015; Plowright *et al.* 2015; Paez *et al.* 2017; Edson *et al.* 2019; Peel *et al.* 2019).

The vital ecosystem services provided by bats through insect pest control (e.g. Kolkert et al. 2020), pollination (e.g. Law and Lean 1999; Eby and Law 2008) and seed dispersal (e.g. Moran et al. 2009) are often overshadowed in public perception by their potential to act as reservoirs of zoonotic viruses, such as Hendra virus, Australian bat lyssavirus and Menangle virus. Most research on bat-borne viruses in Australia (~200 papers to date) has focussed on these pathogens, with only three coronavirus surveillance studies (Smith et al. 2016; Holz et al. 2018; Prada et al. 2019). Even though none of Australia's bat species have distributions overlapping with species that are known hosts of SARS-related viruses, most of the chiropteran genera in which zoonoticrelated coronaviruses have been detected in China and elsewhere are represented in Australia (e.g. Rhinolophus spp., Hipposideros spp., Chaerophon spp., Pipistrellus spp.) (Tang et al. 2006; Pfefferle et al. 2009; Tong et al. 2009).

Against this background, we first seek to review the current state of knowledge of bat coronaviruses globally and within Australia, identifying key areas in need of further research. Second, we canvass the potential for spillover of endemic Australian bat coronaviruses to humans and the potential for transmission of SARS-CoV-2 from infected humans into Australian bat populations.

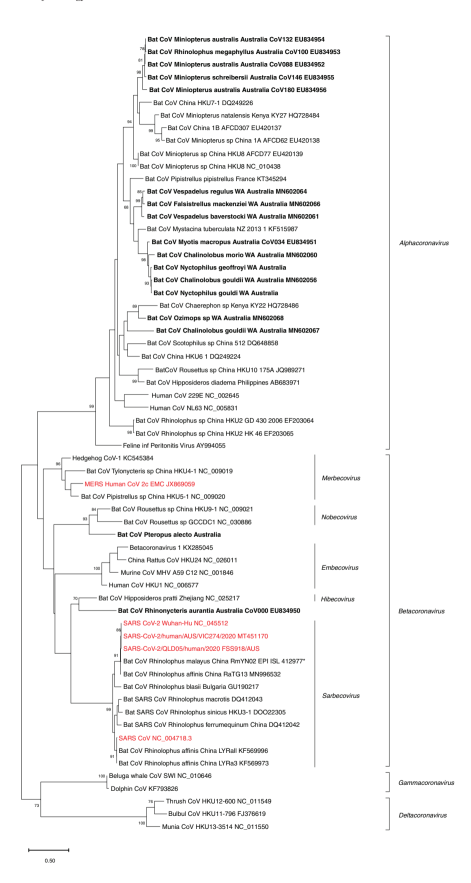
Coronaviruses in bats worldwide

Bats naturally host viruses from two of the four genera of coronaviruses: alpha- and betacoronaviruses – the other two (delta- and gammacoronaviruses) are predominantly found in birds (Woo *et al.* 2009, 2010; Su *et al.* 2016) (Fig. 1). The diversity of coronaviruses circulating in bat populations is the highest detected to date in any mammalian host group (Drexler *et al.* 2014). Empirical detections and estimates suggest the existence of 2–5 unique coronaviruses per bat species, and potentially up to 7000 coronaviruses worldwide when ~1400 bats species are considered (Anthony *et al.* 2017*a*; Willoughby *et al.* 2017; Latinne *et al.* 2020; Simmons and Cirranello 2020).

All of the known zoonotic coronaviruses with origins in bats are betacoronaviruses, belonging either to the Sarbecovirus subgenus (i.e. the SARS-related coronaviruses [SARSr-CoV]: SARS-CoV-1, SARS-CoV-2), or the Merbecovirus subgenus (MERS-CoV) (Fig. 1). Although the specific conditions resulting in the spillover of SARS-

CoV-1. SARS-CoV-2 and MERS-CoV remain unresolved, the closest evolutionary relatives to these viruses have been identified in bat hosts. Viruses with 88-91% nucleotide identity to SARS-CoV-1 have been amplified from horseshoe bat species (Rhinolophus sinicus, R. ferrumequinum, R. macrotis and R. pearsoni), supporting Rhinolophus bats as the ancestral reservoir origin of SARS (Lau et al. 2005; Li et al. 2005; He et al. 2014; Zhou et al. 2020; Zhou et al. 2020a). The viruses most closely related to SARS-CoV-2 identified so far include lineages of sarbecoviruses circulating in bat populations in China: RaTG13 CoV isolated from Rhinolophus affinis (Zhou et al. 2020b) and RmYN02 CoV detected in Rhinolophus malayanus (Zhou et al. 2020). Beyond Rhinolophidae, SARS-related coronaviruses have also been detected in bats from the Hipposideridae and Vespertilionidae in Europe and Asia, and the Molossidae in Asia (Drexler et al. 2014; Fan et al. 2019; Wong et al. 2019). At least three bat families have been identified as hosts of Merbecoviruses to date, though most detections have been in bats from the Vespertilionidae (reviewed in Wong et al. 2019).

Existing bat coronavirus studies demonstrate several factors with apparent influence on coronavirus detectability and diversity. Coronavirus detection rates are statistically significantly higher in specimens collected from bats in humanaltered landscapes (e.g. particularly associated with animal usage in hunting, trade, etc), compared with those from undisturbed environments, suggesting the influence of environmental conditions on bat-coronavirus interactions (Anthony et al. 2017a). Coronavirus diversity is higher where bat diversity is also high, suggesting an intimate and long evolutionary history of interactions. This is evident at the continental scale (bat coronavirus diversity is highest in South-east Asia, northern Latin America, and south-west, equatorial and east Africa) and also within continents (Anthony et al. 2017a; Wong et al. 2019; Joffrin et al. 2020). Phylogenetic analyses of bat coronavirus genetic sequences show that closely related coronaviruses are commonly shared among closely related bat families (Drexler et al. 2014). For instance, in Mexico, coronaviruses in vespertilionid bats cluster together with coronaviruses in molossid bats (Anthony et al. 2013). This relationship supports an influence of cospeciation and host-switching in the ecology and evolution of bats and their coronaviruses (Cui et al. 2007; Joffrin et al. 2020; Latinne et al. 2020). Both alpha- and betacoronaviruses have been detected across all continents occupied by bats, yet in North America, betacoronaviruses have been detected in Mexico but are notably absent in bats in the USA and Canada (Anthony et al. 2013; Wong et al. 2019). It is important to note that sampling efforts are not globally uniform, and therefore such absences may not necessarily represent true diversity and abundance of circulating coronaviruses (Anthony et al. 2017a). This may also affect analyses that indicate that the number of coronavirus species detected per bat family varies by continent in Africa and Asia (with high rates of detection in Pteropodidae in Africa; high rates of detection in Miniopteridae, Vespertilionidae and Pteropodidae in Asia), but bat family did not influence rate of detection in Latin America (Anthony et al. 2017a).



Coronaviruses in Australian bats

Coronavirus diversity and antibody prevalence in Australian bats is known from two cross-sectional studies on bat faeces or rectal swabs, encompassing 35 Australian bat species across 17 genera and 8 families (Smith et al. 2016; Prada et al. 2019) (Table 1, Fig. 2). Based on PCR amplification and sequencing of a 396-440 bp fragment of the RNA-dependent RNA polymerase (RdRp) gene, Smith et al. (2016) and Prada et al. (2019) detected two betacoronaviruses (exclusively from pteropodiformes) and seven alphacoronaviruses (mainly from vespertilionid bats) across 15 Australian bat species. Full genome sequences from four of the seven alphaviruses amplified from host species have been subsequently obtained and made publicly available (Prada et al. 2019). To date, there is no evidence of any SARS-like viruses (Sarbecoviruses) or MERS-like viruses (Merbecoviruses) in any Australian bat species, though cross-reactive antibodies have been detected (Smith et al. 2016; W. Boardman, pers. comm.) and further surveillance is currently underway.

In agreement with other studies globally (Drexler et al. 2014; Lacroix et al. 2017), strains amplified from Australian bats have strong host associations, with sequences clustering by host taxonomic affinity instead of sampling site (Fig. 1). For example, betacoronaviruses amplified from Pteropus alecto and Rhinonicteris aurantia were more similar to other sequences amplified from Asian bats of the same taxonomic order than to each other.

Notable differences are also evident in Australian bat coronavirus studies; for example, despite high viral prevalences of betacoronaviruses in rhinolophid bats in China (e.g. 39%: Lau et al. 2005) and elsewhere (e.g. 32%: Drexler et al. 2010), no betacoronaviruses have been detected in Australian rhinolophids. Despite an extensive sampling effort of Rhinolophus megaphyllus (n = 506), only one bat tested positive by PCR and cross-regional seroprevalence was low (6.7%; 95% CI = 4-9%) (Smith et al. 2016). The single positive result was an alphacoronavirus with 99% nucleotide sequence identity with Miniopterus bat coronavirus HKU8 and which was widespread in concurrently sampled Miniopterus australis and M. schreibersii (now renamed M. orianae), suggesting that R. megaphyllus may not be the primary reservoir host for this viral species. Similarly, there has been no detection of betacoronaviruses in Australian Hipposideros species, even though SARSr-CoV have been amplified from this genera in Africa (Pfefferle et al. 2009; Quan et al. 2010). In Australia, no active viral shedding was detected in *Hipposideros ater* (n = 56), and seroprevalence was very low (1.5%; 95% CI = 0.2-8.3%). Equally surprising as the lack of representative Australian betacoronavirus strains within these genera is the small number of strains and lack of host-specific alphacoronaviruses detected within the broader Pteropodiformes group, since alphacoronaviruses generally have greater detection rates and a widespread global distribution (Wong *et al.* 2019).

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Sample sizes and the employed primer set are congruent with global studies (Tsuda et al. 2012; Moreira-Soto et al. 2015; Lacroix et al. 2017; Joffrin et al. 2020), but broader sampling would help determine how representative these existing studies are, and how sample sizes, sample type, number of sampling sites, sampling season, number of bat species surveyed, or the implemented PCR assay have affected detection rates. For example, an additional study focussed in Australian populations of Miniopterus orianae yielded no coronavirus detections from oropharyngeal swabs (Holz et al. 2018). This lack of detection is likely due to decreased sensitivity of coronavirus detection in oropharyngeal swabs, because the route of coronavirus shedding in bats is predominantly through faeces (Wacharapluesadee et al. 2013; Wong et al. 2019). By comparison, rectal swabs collected from the same species in south-east Queensland and the Northern Territory were positive for coronavirus RNA in 26.5% and 10.2% of individuals, respectively (Smith et al. 2016) (Table 1). Only small numbers of individuals of several Australian bat species have been screened for coronaviruses, and an additional 46 species have not yet been screened at all (Table 2). A meta-analysis of bat coronavirus studies globally identified that a sampling effort of ~400 individuals per species is required to capture the full diversity of coronaviruses present in each species (Anthony et al. 2017a). Only one Australian species has been sampled to this intensity (R. megaphyllus, n = 506), and the single detection in this species suggests that this target sample number must be replicated across sites and seasons before firm conclusions are drawn on a species' viral diversity.

Spillover potential of Australian bat-borne coronaviruses

Various processes must align in space and time for viruses to successfully spill over from a natural reservoir host to alternative hosts (Plowright *et al.* 2017). The viruses must be circulating within reservoir host populations, be shed and survive within the environment at a time and place where novel hosts may be exposed, and, finally, overcome any structural or immune barriers to infection and replication within that host. These cross-species transmission events can involve one-directional transmission from one species to another (e.g. bat to civet to human, as is suspected for SARS-CoV-1), or circular transmission between two or more host species (e.g. a shared virus between two bat host species; conceptually illustrated in Fig. 3). While coronaviruses are known to be circulating within Australian bat populations, there have been no reports of their spillover to humans, either directly or via an

Fig. 1. Maximum-likelihood coronavirus consensus tree based on a 394-bp fragment of the RdRp gene. Representative coronavirus strains amplified from Australian bats are shown in bold and coronavirus sequences associated with emergent pandemics (MERS, SARS-CoV-1, SARS-CoV-2) are highlighted in red in the online version of this article. GenBank accession numbers are shown at the end of each sequence; the sequence indicated with an asterisk shows its GISAID accession reference. The tree was constructed with MegaX (Kumar *et al.* 2018) using a General Time Reversible evolutionary model, a Gamma distribution and 1000 bootstrap replication. Bootstrap values above 70% are indicated on the tree branches. Branch lengths reflect the number of substitutions per site. Aligned sequences are included as a supplementary material file.

Table 1. Evidence of coronavirus infection in Australian bats across existing studies, and antibodies cross-reactive with SARS-coronavirus antigen Geographic location of sampling: FNQ, far-north Queensland; CQ, central Queensland; SEQ, south-east Queensland; NT, Northern Territory; WA, Western Australia; SE-AUST, south-eastern Australia; SW-WA, south-west Western Australia; SA, South Australia; Vic., Victoria. PCR and Antibody results are represented as: no. of positive individuals/no. tested (proportion). Blank cells = not tested

Genus and higher taxa	Species	Common name	Location	PCR ^A : +/n (%)	Antibody ^B : +/n (%)	Study
Pteropodiformes						
Hipposideridae						
Hipposideros	ater	Dusky leaf-nosed bat	FNQ	0/29 (0%)	0/29 (0%)	Smith et al. (2016)
			NT	0/27 (0%)	0/4 ^C	Smith et al. (2016)
			WA		0/31 (0%)	Smith et al. (2016)
Rhinonicteris	aurantia	Pilbara leaf-nosed bat	NT	1/126 (1%)	0/105 (0%)	Smith et al. (2016)
Megadermatidae			3.700		4164 (4.600)	
Macroderma	gigas	Ghost bat	NT	0/57 (0%)	1/63 (1.6%)	Smith et al. (2016)
D: 11.1			WA		17/21 (81%)	Smith et al. (2016)
Pteropodidae	1 .	DI I d ' C	GEO.	4/22 (120/)	0/24 (26 50/)	0.34 (1.0010
Pteropus	alecto	Black flying-fox	SEQ	4/33 (12%)	9/34 (26.5%)	Smith et al. (2016)
Pteropus Pteropus	conspicillatus	Spectacled flying-fox	FNQ SEQ	0/27 (0%)	6/40 (15%) 12/73 (16.4%)	Smith et al. (2016) Smith et al. (2016)
Pteropus Pteropus	poliocephalus scapulatus	Grey-headed flying-fox Little red flying-fox	NT	0/27 (0%)	3/40 (7.5%)	Smith et al. (2016)
Rhinolophidae	scapulalus	Little fed flyffig-fox	INI		3/40 (7.570)	Silitii et at. (2010)
Rhinolophus	megaphyllus	Smaller horseshoe bat	FNQ	1/58 (1.7%)	5/61 (8.2%)	Smith et al. (2016)
иногорниз	megapnymus	Smarter norseshoe but	SEQ	0/448 (0%)	13/399 (3.3%)	Smith et al. (2016)
Vespertilioniformes			DEQ	0/110 (0/0)	13/377 (3.370)	Sintil Ci ui. (2010)
Emballonuridae						
Saccolaimus	flaviventris	Yellow-bellied sheath-tailed bat	WA		0/18 ^C	Smith et al. (2016)
Taphozous	spp.	Tomb bats	WA		8/38 (38%)	Smith et al. (2016)
Miniopteridae	11				` '	` ′
Miniopterus	australis	Little bent-wing bat	CQ	1/20 (5%)	15/30 (50%)	Smith et al. (2016)
1			FNQ	14/30 (46.7%)	16/30 (53.3%)	Smith et al. (2016)
			SEQ	38/154 (24.7%)	80/124 (64.5%)	Smith et al. (2016)
			WA		1/1 ^C	Smith et al. (2016)
Miniopterus	$orianae^{D}$	Northern bent-wing bat (Miniopterus	NT	6/59 (10.2%)	25/56 (44.6%)	Smith et al. (2016)
		orianae orianae)				
			SEQ	63/238 (26.5%)	145/211 (68.7%)	Smith et al. (2016)
		Eastern/large bent-wing bat (Miniopterus	Vic.	0/26 (0%)		Holz et al. (2018)
		orianae oceanensis)				
		Southern bent-wing bat (Miniopterus	Vic.	0/32 (0%)		Holz et al. (2018)
		orianae bassanii)	~ .	0.4 = = .00.13		***
		Southern bent-wing bat (Miniopterus	SA	0/155 (0%)		Holz et al. (2018)
M 1 - 11		orianae bassanii)				
Molossidae	1:-	White stained fore tailed had	SW-WA	0/9 ^C	1/11 ^C	Des de et el (2010)
Astronomus Chaerephon	australis jobensis	White-striped free-tailed bat Northern free-tailed bat	WA	0/9	2/4 ^C	Prada et al. (2019) Smith et al. (2016)
Ozimops ^E	lumsdenae ^F	Northern free-tailed bat	SEQ	0/3 ^C	40/41 (97.6%)	Smith et al. (2016)
Ozimops ^E	norfolkensis	East coast free-tailed bat	SEQ	0/3 0/1 ^C	40/41 (97.070)	Smith et al. (2016)
Ozimops ^E	spp.	Molossid bats	SW-WA	1/3 ^C	0/1 ^C	Prada et al. (2019)
Vespertilionidae	эрр.	Holossia bats	5W W11	1/3	0/1	1 rada er ar. (2015)
Chalinolobus	spp.	Long-tailed bats	WA		2/4 ^C	Smith et al. (2016)
Chalinolobus	gouldii	Gould's wattled bat	SW-WA	59/232 (25.4%)	2/62 (3.0%)	Prada et al. (2019)
Chalinolobus	morio	Chocolate wattled bat	SW-WA	17/45 (37.8%)	4/59 (6.8%)	Prada et al. (2019)
Myotis	macropus	Southern myotis	FNQ	0/31 (0%)	18/31 (58.1%)	Smith et al. (2016)
•	•	•	SEQ	13/64 (20.3%)		Smith et al. (2016)
Nyctophilus	bifax	Eastern long-eared bat	SEQ	0/6 ^C		Smith et al. (2016)
Nyctophilus	gouldi	Gould's long-eared bat	SEQ	0/7 ^C		Smith et al. (2016)
			SW-WA	2/56 (3.5%)	7/67 (10.4%)	Prada et al. (2019)
Nyctophilus	geoffroyi	Lesser long-eared bat	SW-WA	2/51 (3.9%)	0/27 (0%)	Prada et al. (2019)
Nyctophilus	major	Western long-eared bat	SW-WA	0/10 ^C	2/5 ^C	Prada et al. (2019)
Scotorepens	greyii	Little broad-nosed bat	SEQ	0/1 ^C		Smith et al. (2016)
Scotorepens	rueppellii	Greater broad-nosed bat	SEQ	0/1 ^C		Smith et al. (2016)
Scotorepens	spp.	Broad-nosed bats	SEQ		24/24 (100%)	Smith et al. (2016)
			WA	o :- C	0/1 ^C	Smith et al. (2016)
Scotorepens	balstoni	Inland broad-nosed bat	SW-WA	0/9 ^C	0/2 ^C	Prada et al. (2019)
Vespadelus	finlaysoni	Finlayson's cave bat	WA	1	0/1 ^C	Smith et al. (2016)
Vespadelus	pumilus	Eastern forest bat	SEQ	1/4 ^C	5/01 /15 10 C	Smith et al. (2016)
Vespadelus	troughtoni	Eastern cave bat	FNQ	0/31 (0%)	5/31 (16.1%)	Smith et al. (2016)
Vespadelus	baverstocki	Inland forest bat	SW-WA	2/4 ^C	0/1 ^C	Prada et al. (2019)
Vespadelus	regulus	Southern forest bat	SW-WA	17/141 (12.1%)	9/150 (6.0%)	Prada et al. (2019)
Falsistrellus	mackenziei	Western false pipistrelle	SW-WA	2/11 ^C	0/7 ^C	Prada et al. (2019)

APCR results are based on detection of viral RNA via Rdrp PCR on faecal samples or rectal swabs, except for results from Miniopterus orianae oceanensis and Miniopterus orianae bassanii, which were based on oropharyngeal swabs.

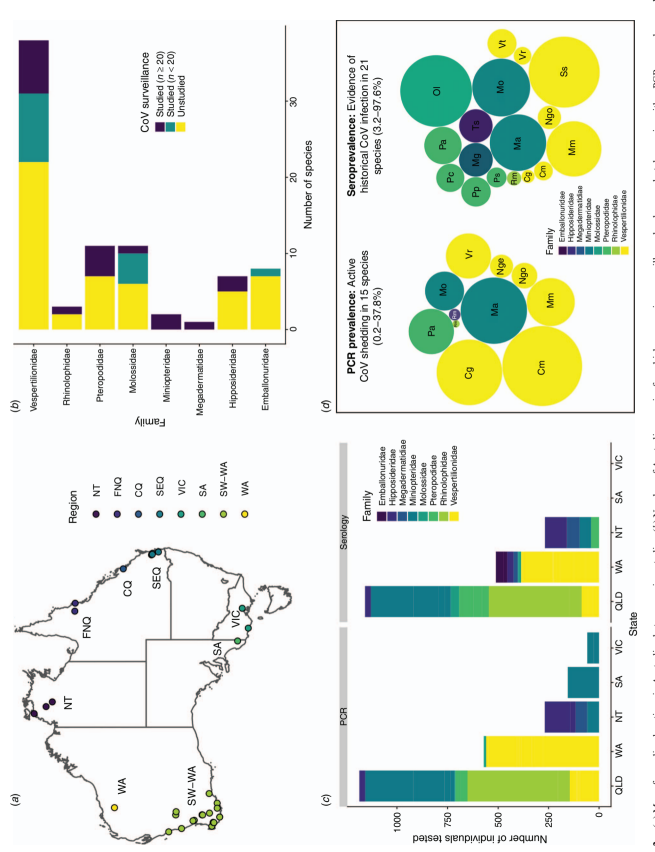
^BAntibody results reflect detection of antibodies cross-reactive with SARS-coronavirus antigen via ELISA or microbead assay (refer to original studies for details).

^CPrevalence not calculated (for species/locations with total sample sizes below 20 individuals).

^DThe species *Miniopterus orianae* was formerly recognised as *Miniopterus schreibersii* (identified as *M. schreibersii* in Smith et al. 2016).

EThe genus Ozimops was formerly recognised as Mormopterus (as identified in Smith et al. 2016).

FThe species Ozimops lumsdenae was formerly recognised as Mormopterus beccarii (identified as M. beccarii in Smith et al. 2016).



Queensland; WA, Western Australia; NT, Northern Territory; SA, South Australia; Vic., Victoria). (d) Overall PCR prevalence and seroprevalence by species (n > 20), where circle area is proportional to Nyctophilus gouldi, Ol, Ozimops lumsdenae; Pa, Pteropus alecto; Pc, Pteropus conspicillatus; Pp, Pteropus poliocephalus; Ps, Pteropus scapulatus; Rm, Rhinolophus megaphyllus; Ra, Rhinonicteris (a) Map of sampling locations in Australian bat coronavirus studies. (b) Number of Australian species for which coronavirus surveillance has been undertaken, using either PCR or serology and prevalence (Cg, Chalinolobus gouldii; Cm, Chalinolobus morio; Mg, Macroderma gigas; Ma, Miniopterus australis; Mo, Miniopterus orianae; Mm, Myotis macropus; Nge, Nyctophilus geoffroyi; Ngo, noting whether data were derived from more or less than 20 individuals of each species. (c) Number of individuals tested by PCR or serology within each bat family, faceted by Australian state (Qld, aurantia; Ss, Scotorepens spp.; Ts, Taphozous spp.; Vr, Vespadelus regulus; Vt, Vespadelus troughtoni.) A colour version of this figure is available online.

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Table 2. List of Australian bat species (by family), including common names and inclusion in bat coronavirus surveillance studies

n.a., no data available (species not included in any previous coronavirus studies). ID, Identified. Smith *et al.* (2016) were not able to differentiate between the two species of *Taphozous* that have an over-lapping range at the Western Australian sampling location (*T. georgianus* and *T. hilli*), but 38 serum samples from *Taphozous* spp. were tested for CoV antibodies

Taxa	Common name	CoV surveillance
Emballonuridae		
Saccolaimus flaviventris	Yellow-bellied sheath-tailed bat	Smith et al. (2016)
Saccolaimus mixtus	Cape York sheath-tailed bat	n.a.
Saccolaimus saccolaimus	Bare-rumped sheath-tailed bat	n.a.
Taphozous australis	Coastal sheath-tailed bat	n.a.
Taphozous georgianus	Common sheath-tailed bat	n.a.
Taphozous hilli	Hill's sheath-tailed bat	n.a.
Taphozous kapalgensis	Arnhem sheath-tailed bat	n.a.
Taphozous troughtoni	Troughton's sheath-tailed bat	n.a.
Hipposideridae		
Hipposideros ater	Dusky leaf-nosed bat	Smith et al. (2016)
Hipposideros cervinus	Fawn leaf-nosed bat	n.a.
Hipposideros diadema	Diadem leaf-nosed bat	n.a.
Hipposideros inornatus	Arnhem leaf-nosed bat	n.a.
Hipposideros semoni	Semon's leaf-nosed bat	n.a.
Hipposideros stenotis	Northern leaf-nosed bat	n.a.
Megadermatidae		
Macroderma gigas	Ghost bat	Smith et al. (2016)
Miniopteridae		
Miniopterus australis	Little bent-winged bat	Smith et al. (2016)
Miniopterus orianae ^A	Large bent-winged bat	Smith et al. (2016) and
	-	Holz et al. (2018)
Molossidae		
Austronomus australis	White-striped free-tailed bat	Prada et al. (2019)
Chaerephon jobensis	Greater northern free-tailed bat	Smith et al. (2016)
Ozimops cobourgianus	Northern coastal free-tailed bat	n.a.
Ozimops eleryi	Bristle-faced free-tailed bat	n.a.
Ozimops halli	Cape York free-tailed bat	n.a.
Ozimops kitcheneri ^B	Western free-tailed bat	Prada et al. (2019)
Ozimops lumsdenae ^C	Northern free-tailed bat	Smith et al. (2016)
Ozimops norfolkensis	Eastern coastal free-tailed bat	Smith et al. (2016)
Ozimops petersi	Inland free-tailed bat	n.a.
Ozimops planiceps	Southern free-tailed bat	n.a.
Ozimops ridei	Ride's free-tailed bat	n.a.
Pteropodidae		
Dobsonia magna	Bare-backed fruit bat	n.a.
Macroglossus minimus	Northern blossom bat	n.a.
Nyctimene robinsoni	Eastern tube-nosed bat	n.a.
Pteropus alecto	Black flying-fox	Smith et al. (2016)
Pteropus brunneus	Percy island flying-fox	n.a.
Pteropus conspicillatus	Spectacled flying-fox	Smith et al. (2016)
Pteropus macrotis	Large-eared flying-fox	n.a.
Pteropus natalis	Christmas island flying-fox	n.a.
Pteropus poliocephalus	Grey-headed flying-fox	Smith et al. (2016)
Pteropus scapulatus	Little red flying-fox	Smith et al. (2016)
Syconycteris australis	Eastern blossom bat	n.a.
Rhinolophidae		
Rhinolophus megaphyllus	Eastern horseshoe bat	Smith et al. (2016)
Rhinolophus robertsi	large-eared horseshoe bat	n.a.
Rhinolophus sp.	Intermediate horseshoe bat	n.a.
Rhinonycteridae		
Rhinonicteris aurantia	Orange leaf-nosed bat	Smith et al. (2016)
Vespertilionidae		
Chalinolobus dwyeri	Large-eared pied bat	n.a.
Chalinolobus gouldii	Gould's wattled bat	Prada et al. (2019)
Chalinolobus morio	Chocolate wattled bat	Prada et al. (2019)
Chalinolobus nigrogriseus	Hoary wattled bat	n.a.
Chalinolobus picatus	Little pied bat	n.a.
Falsistrellus mackenziei	Western falsistrelle	Prada et al. (2019)
Falsistrellus tasmaniensis	Eastern falsistrelle	n.a.
Murina florium	Flute-nosed Bat	n.a.
	Large-footed myotis	Smith et al. (2016)
Myotis macronus		
Myotis macropus Nyctophilus arnhemensis	Arnhem long-eared bat	n.a.

Table 2. (continued) (continued next page)

axa	Common name	CoV surveillance	
Nyctophilus corbeni	Corben's long-eared bat	n.a.	
Nyctophilus daedalus	Pallid long-eared bat	n.a.	
Nyctophilus geoffroyi	Lesser long-eared bat	Prada et al. (2019)	
Nyctophilus gouldi	Gould's long-eared bat	Smith <i>et al.</i> (2016) and Prada <i>et al.</i> (2019)	
Nyctophilus howensis	Lord Howe long-eared bat	n.a.	
Nyctophilus major	Greater long-eared bat	Prada et al. (2019)	
Nyctophilus sherrini	Tasmanian long-eared bat	n.a.	
Nyctophilus walkeri	Pygmy long-eared bat	n.a.	
Phoniscus papuensis	Golden-tipped bat	n.a.	
Pipistrellus adamsi	Forest pipistrelle	n.a.	
Pipistrellus murrayi	Christmas island pipistrelle	n.a.	
Pipistrellus westralis	Northern pipistrelle	n.a.	
Scoteanax rueppellii	Greater broad-nosed bat	Smith et al. (2016)	
Scotorepens balstoni	Inland broad-nosed bat	Prada et al. (2019)	
Scotorepens greyii	Little broad-nosed bat	Smith et al. (2016)	
Scotorepens orion	Eastern broad-nosed bat	n.a.	
Scotorepens sanborni	Northern broad-nosed bat	n.a.	
Scotorepens sp.	Parnaby's broad-nosed bat	n.a.	
Vespadelus baverstocki	Inland forest bat	Prada et al. (2019)	
Vespadelus caurinus	Northern cave bat	n.a.	
Vespadelus darlingtoni	Large forest bat	n.a.	
Vespadelus douglasorum	Yellow-lipped cave bat	n.a.	
Vespadelus finlaysoni	Finlayson's cave bat	Smith et al. (2016)	
Vespadelus pumilus	Eastern forest bat	Smith et al. (2016)	
Vespadelus regulus	Southern forest bat	Prada et al. (2019)	
Vespadelus troughtoni	Eastern cave bat	Smith et al. (2016)	
Vespadelus vulturnus	Little forest bat	n.a.	

^AThe species *Miniopterus orianae* was formerly recognised as *Miniopterus schreibersii* (identified as *M. schreibersii* in Smith *et al.* 2016).

intermediate host. Some evidence exists for bat-bat crossspecies transmission events between sympatric species within the studied communities (in fact, they appear to be somewhat common) (Fig. 3), yet sustained infection within the new host and further transmission within populations may be limited. For instance, high nucleotide similarity (100%) was detected between sequences amplified from M. australis and R. megaphyllus, yet marked differences existed between the species in the proportion of active infections (RNA detection; 46.7% versus 1.7%, respectively) and historical infections (coronavirus antibodies) (53% versus 8.2%, respectively) (Smith et al. 2016). These data are indicative of a crossinfection event from the vespertilionid bat M. australis into the pteropodiforme R. megaphyllus and warrant further investigation. Similarly, cross-infections were also detected in south-west Western Australian bat communities, with tree hollows proposed as a source of transmission between sympatric bat species (Prada et al. 2019). One alphacoronavirus strain detected in Chalinolobus gouldii had a spike protein with a higher sequence similarity to the coronavirus that causes Porcine Endemic Diarrhoea in pigs (Jung and Saif 2015) than to any other virus of bat origin (Prada et al. 2019). Since the spike protein determines host tropism, the ability of the spike protein to enter mammalian cells other than bat cells is currently under investigation (Haynes et al., unpubl. data). In general, however, the

spatiotemporal shedding dynamics of Australian bat-borne coronaviruses and their capacity to infect cells and overcome immune systems of human or other animal species are unknown, making it challenging to assess whether spillover does not occur, or does not cause disease, or occurs but is simply not detected.

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Insights from other bat-viral systems and geographic regions will be key in targeting further research in Australia. The origin of SARS-CoV-2, the chain of transmission to humans, and direct evidence for the involvement of an intermediate host are yet to be established. Pathogen transmission between species cannot occur without every condition, in a hierarchical series of events, lining up (Plowright et al. 2017). Hence, if intermediate hosts are required, then even if diverse viruses are present in bats and peaks of shedding generate pathogen pressure for bat-borne coronaviruses, spillover may not be able to occur without the presence of these competent intermediate hosts from which human exposure occurs. There are multiple reports of transmission of bat-borne coronaviruses to various laboratory animal species in experimental settings (Gretebeck and Subbarao 2015), domestic and wild felids across a range of settings (Shi et al. 2020; US Department of Agriculture Animal and Plant Health Inspection Service 2020), and to nonbat wildlife species utilised in wildlife farming and trade (Shi and Hu 2008; Gretebeck and Subbarao 2015). There are also

^BLikely species identified as 'Ozimops sp.' in Prada et al. (2019).

^CThe species *Ozimops lumsdenae* was formerly recognised as *Mormopterus beccarii* (identified as *M. beccarii* in Smith *et al.* 2016).

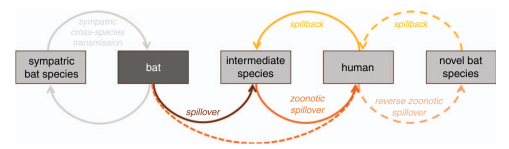


Fig. 3. Conceptual schematic representing terms for various forms of cross-species transmission of viruses associated with bats. Demonstrated transmission pathways for bat coronaviruses are shown in solid lines, hypothesised pathways are shown in dashed lines.

examples of spillover of coronaviruses from bats to livestock or farmed domestic species: in camels, associated with the emergence of MERS in 2012 in Saudi Arabia (Anthony et al. 2017b), and in pigs, associated with the emergence of Swine Acute Diarrhoea Syndrome (SADS) in 2016 in southern China that resulted in the deaths of ~25 000 pigs across four farms (Zhou et al. 2018). The novel SADS coronavirus detected in pigs that caused this outbreak shared 96-98% sequence identity with SADS-related bat coronaviruses present in ~10% of Rhinolophus spp. bats in the same region. This event was preceded by at least three other examples of bat-pig viral spillover: transmission of Nipah virus from Pteropus spp. flying foxes to pigs in Malaysia (Chua et al. 2000), Reston ebolavirus in pigs in the Philippines and China (Jayme et al. 2015), and Menangle virus from Pteropus poliocephalus to pigs in Australia (Philbey et al. 1998). The latter example highlights that bat-pig viral transmission events are feasible within an Australian context. Spillover events are more likely to succeed where susceptible intermediate hosts are stressed and immunocompromised and where ongoing transmission is facilitated by high-density cohousing and poor on-farm biosecurity (Lindahl and Grace 2015).

The potential for direct bat-human transmission of coronaviruses remains unclear (Fig. 3). There is no molecular evidence for direct spillover; however, Wang *et al.* (2018) found serological evidence of human exposure to SARS-related coronaviruses in 6 of 218 (2.7%) rural people living close to caves in which SARS-related coronaviruses had been identified in bats. None of the positive individuals had a history of exposure to SARS-CoV-1 virus during the 2002–03 outbreak and none could recall any clinical symptoms consistent with SARS-CoV-1 infection in the past 12 months, suggesting the possibility of subclinical or mild clinical infection. Uncertainty remains as to whether these findings could represent transmission via an intermediate host, but they are sufficient to warrant inclusion of direct bat-human transmission routes in coronavirus spillover risk assessments.

Several human—bat interaction scenarios could theoretically present the opportunity for spillover of bat coronaviruses to humans in Australia, including where people are occupationally exposed to bats and bat body fluids. Bat rescuers and rehabilitators would likely be at high risk of exposure because of the number of animals they handle, the length of time that animals are frequently in care, and the often physically close nature of contact between carer and bat

(particularly orphan flying-foxes). Indeed, following the emergence of Hendra virus in 1994 (which spilled from flying-foxes to horses to humans), health authorities recognised that bat carers and others working with bats were an ideal sentinel group to screen for serological evidence of exposure to Hendra virus; however, no evidence of exposure was found (Selvey *et al.* 1996).

Given the tangible threats to human health and the unknown risks that emergent coronaviruses pose to wild bat populations, future surveillance efforts on bat taxa traditionally associated with zoonotic strains should prioritise species common to the human/urban interface. Additionally, the commonality and marked variability of alphacoronavirus infections among insectivorous bats, provide an opportunity to further investigate drivers of cross-species spillover, as well as pathogen persistence, viral shedding, transmission routes and their associations to the host ecology and health (Jeong *et al.* 2017; Smith 2015).

Potential establishment of novel coronaviruses in Australian bat populations following human-to-bat transmission

SARS-CoV-2 is expected to persist as the world's fifth endemically circulating human coronavirus (Kissler et al. 2020), and it is unlikely to be the last novel coronavirus linked to bat origins. Widespread transmission within human populations of viruses that are either bat-borne, or whose progenitors originate in bats, raises the possibility of reverse zoonotic spillover (Fig. 3) of these viruses from humans into novel bat or other animal hosts that have no evolutionary association with the circulating strains. Bat organisations internationally have responded to this threat by advocating, at a minimum, for increased biosecurity protocols when handling bats for research or rehabilitation (IUCN Bat Specialist Group 2020; Olival et al. 2020; Runge et al. 2020). In some cases, this has extended to blanket bans on handling wild bats. Establishment of SARS-CoV-2, or other endemic human coronaviruses from people into Australian bat populations, could have serious public health implications, via establishment of an adjunct source of human infection (via 'spillback': Fig. 3) or via recombination events with endemic bat coronavirus strains. Bat welfare, conservation, and management, already contentious issues in Australia (Degeling and Kerridge 2012; Kung et al. 2015), would likely

be further compromised. The likelihood of human-to-bat transmission of SARS-CoV-2 in Australia is likely to reflect levels of community transmission (at the time of writing (4 June 2020), there had been a total of ~7240 cases of COVID-19 in Australia (~28 per 100 000 individuals), and detection rates for new cases had dropped to an average of 13 per day: Australian Government Department of Health 2020). However, the likelihood of transmission would be expected to increase proportionately with increases in human cases, which has subsequently been observed in a 'second wave' in the State of Victoria (Australian Government Department of Health 2020). Given the dynamic nature of the COVID-19 pandemic, and the potentially serious consequences of a human-bat transmission event, now is an ideal time to be discussing practices to mitigate the risk of human-to-bat transmission of endemic and exotic coronaviruses more broadly.

The risk of human-to-bat transmission will vary with the emergence of each new human-transmitted coronavirus, and also over space and time. Important human factors to consider in assessing this risk include prevalence in the human population and the proportion of asymptomatic carriers, as well as the likely transmission route, infectious period, and virus survival characteristics (Olival et al. 2020). Conditional on exposure of a bat to an infected human, successful humanto-bat transmission will depend on individual bat and species susceptibility, which may be based on cross-reactive immunity to circulating bat coronaviruses, receptor sequence similarity across species, or alternative transmission mechanisms yet to be elucidated (Olival et al. 2020; Wildlife Health Australia, unpubl. data). Finally, the frequency and patterns of contact among people with different occupational or recreational exposure to bats (e.g. researchers, ecological consultants, bat rehabilitators, veterinary staff, and recreational cavers) and the different bat species they interact with, will determine groups at greatest risk of initiating or receiving spillover. Regardless, direct human-to-bat contact might not be necessary for spillover: since SARS-CoV-2 can be shed in faeces and may not be wholly inactivated by wastewater treatment processes, Franklin and Bevins (2020) have hypothesised that wastewater might be a potential route of transmission when human case numbers are high.

Discussion

Australian bats are not known to host Merbecoviruses or Sarbecoviruses with demonstrated zoonotic and pandemic potential. Nevertheless, the patchy coverage of coronavirus studies to date should alert us to the possibility that related viruses might be detected with increased sampling intensity. The wide variation in both viral RNA (0–46%) and antibody (0–100%) prevalence detected across species, space and time from existing cross-sectional studies also suggests that point-estimates of prevalence should be interpreted with care (Becker *et al.* 2019). Such dynamic systems can be understood only through detailed spatiotemporal surveillance of coronaviruses in Australasian bat populations (Plowright *et al.* 2019).

Globally, coronaviruses are widely distributed across bat species and populations. Few studies, however, include sufficient spatial or temporal sampling design or host ecological factors to infer the drivers of transmission. Point prevalence of coronaviruses in wild bat populations varies considerably among sites, seasons, age categories and physiological stages of bats, with values of up to 60% of infected individuals (Montecino-Latorre et al. 2020). For instance, in a study in Pteropus lylei in Thailand, Wacharapluesadee et al. (2018) report higher prevalence of coronaviruses in juveniles than adults, with detections only in adults in January, only in juveniles in April and no differences among both age categories between May and October. Smith (2015) identified significantly higher anti-coronavirus antibody prevalence in M. schreibersii (now M. orianae) and R. megaphyllus in summer and in female adult bats. Comparable, but non-significant, trends were observed for PCR detection of coronaviruses. Similar age and seasonal effects have been reported for coronavirus shedding in other bat species: in some studies, juveniles present up to 16 times higher probability of shedding virus than adults, with a peak during the weaning season (Montecino-Latorre et al. 2020). These age, seasonal and physiologically related infection status and shedding patterns of coronaviruses show similarities to the dynamic of infections by Henipaviruses in Pteropus bats with seasons and life stages of high physiological stress playing a role in infection and shedding risk (Plowright et al. 2008; Field et al. 2015; Paez et al. 2017; Brook et al. 2019; Edson et al. 2019). These extensive data on the ecological drivers of Hendra virus shedding in Australia could provide a foundation upon which to base deeper investigations into the drivers of coronavirus shedding. An increase in coronavirus studies in Australian bats would provide an indication of the broad viral diversity, the potential for novel spillover risks, as well as a baseline comparison for efforts to detect human-bat transmission of CoV.

The current limited information and uncertainty about potentially zoonotic viruses, including coronaviruses, in Australian bats leaves openings for misinformation and fear. For example, in April 2020, a Victorian state member of Parliament called for the grey-headed flying-fox (P. poliocephalus) colony in Yarra Bend National Park to be removed because of a perceived disease threat, including COVID-19 (Rimmer 2020). Further investigations of the diversity, distribution, and zoonotic potential of coronaviruses in Australian bats are needed to allow a rational assessment of the risk, if any, of potential future spillover to humans or livestock, or spillback of SARS-CoV-2 to endemic bats.

Investigations of spillover potential of coronaviruses between Australian bats and humans should be framed in a broader ecological context. Describing the viruses that infect Australian bats is a first step, but without the ecological conditions that promote pathogen shedding, contact with susceptible hosts (intermediate or otherwise), and human behaviours that allow cross-species transmission, spillover may be unlikely. Landscape change is a key factor modulating all of these risk factors for disease emergence (Kessler *et al.* 2018; Johnson *et al.* 2020). The ecological integrity of landscapes affects bat distribution and health, and hence contact with other species and pathogen shedding dynamics. With increasing rates of emergence of pathogens from bats

associated with rapid environmental change (Johnson *et al.* 2020) and amplified stressors as a result of climate change, the links between environmental stress and spillover need more attention. Environmental changes not only drive bat virus emergence, they can decrease the ecosystem services that bats provide that directly benefit humans. Investigations of disease risk need to consider the holistic interaction of bats, their infections, and their environment.

Conclusion

Australia has a unique global advantage with its long history of bat research across a broad range of ecological and associated disciplines, as well as its expertise in viral spillover from bats. We must utilise this foundation to develop integrative approaches to understanding bat health and sustainable protection of human health.

Conflicts of interest

The authors declare no conflicts of interest.

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