Detection and characterization of diverse alpha- and betacoronaviruses from bats in China

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Bats have been implicated as important reservoir hosts of alpha- and betacoronaviruses. In this study, diverse coronaviruses (CoVs) were detected in 50 of 951 (positive rate 5.3%) intestinal specimens of eight bat species collected in four provinces and the Tibet Autonomous Region of China by pan-coronavirus RT-PCR screening. Based on 400-nt RNA-dependent RNA polymerase (RdRP) sequence analysis, eight belonged to genus Alphacoronavirus and 42 to Betacoronavirus. Among the 50 positive specimens, thirteen gave rise to CoV full-length RdRP gene amplification for further sequence comparison, of which three divergent sequences (two from a unreported province) were subjected to full genome sequencing. Two complete genomes of betacoronaviruses (JTMC15 and JPDB144) and one nearly-complete genome of alphacoronavirus (JTAC2) were sequenced and their genomic organization predicted. The present study has identified additional numbers of genetically diverse bat-borne coronaviruses with a wide distribution in China. Two new species of bat CoV, identified through sequence comparison and phylogenetic analysis, are proposed.

KEYWORDS bats; Alphacoronavirus; Betacoronavirus; diversity

INTRODUCTION

Coronaviruses (CoVs, family Coronaviridae, subfamily Coronavirinae) are important human and animal pathogens which, according to the latest release of Virus Taxonomy by the International Committee on Taxonomy of Viruses (ICTV, http://www.ictvonline.org/virusTaxonomy.asp?msl_id=26), currently comprise four distinct genera: Alphacoronavirus (αCoV), Betacoronavirus (βCoV), Gammacoronavirus (γCoV) and Deltacoronavirus (δCoV). This large group of viruses has a wide spectrum of hosts, including humans, rodents, carnivores, chiropters and avians, and cause respiratory, enteric, hepatic and neurological diseases (Lai et al, 2007). They include even public threats such as the severe acute respiratory syndrome (SARS) and the current Middle East respiratory syndrome (MERS) (Moratelli et al, 2015). Bats are host animals of diverse αCoVs and βCoVs that may serve as the ancestral origins of mammalian CoVs (Falcon et al, 2011; Woo et al, 2012). In last decade, increasing numbers of bat CoVs with wide molecular diversities have been reported worldwide, particularly in China (Li et al, 2005; Tang et al, 2006; Woo et al, 2007; Chu et al, 2008; Yuan et al, 2010; He et al, 2014), some of
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which likely have the potential ability to cause human diseases (Ge et al, 2013; He et al, 2014; Menachery et al, 2015). These findings indicate that further diverse CoVs circulate in bat populations. China has a nationwide distribution of about 120 bat species, with many roosting regions remaining uninvestigated for harbored mammal viruses. Here, we report a continuing investigation on bat-borne CoVs in some unexplored regions in China, the results of which have revealed more novel CoVs that circulate and evolve in bat populations with great molecular diversity and wide geographic distribution.

MATERIALS AND METHODS

Bat collection and species conformation
A total of 951 bats covering 5 families and 21 species were captured between 2005 and 2013, in Jilin, Liaoning, Yunnan, Guangdong provinces and the Tibet Autonomous Region, China. Bat species were morphologically identified by a trained field biologist and further confirmed by PCR of their mitochondrial cytochrome b gene sequence (Wang et al, 2003). Respiratory and intestinal tissue specimens were collected separately from each bat and stored at –80 °C immediately until further processing.

RNA extraction and detection by RT-PCR
Viral RNA of each specimen was extracted by using the RNeasy Mini Kit (Qiagen, Hilden, Germany), and was immediately reverse-transcribed with the Superscript III Kit (Invitrogen, San Diego, CA) using random primers. Pan-CoV nested PCR primers were used to amplify a 440-nt sequence in the RNA-dependent RNA polymerase (RdRP) gene by our published methods (He et al, 2014) (see Supplementary Table S1 for primer information). Expected PCR amplicons were directly sequenced by the Sanger method in an ABI 3730 sequencer (Comate Bio, Changchun, China).

RdRP gene amplification and whole genome sequencing
The complete RdRP genes of positive samples were amplified using LA Taq (TaKaRa, Dalian, China). Primers were designed based on RdRp gene sequences of representative αCoV and βCoV strains available in GenBank. Reactions were carried out with a touch-down PCR program: 94 °C for 3 min, then 10 temperature decrement cycles (94 °C for 30 s, 58 °C minus 1 °C per cycle for 30 s, 72 °C for 2 min), followed by 35 normal cycles (94 °C for 30 s, 52 °C for 30 s, 72 °C for 3 min), and a last extension of 72 °C for 10 min.

To obtain the full genomes of the interesting specimens, overlapping amplicons were obtained by the above PCR program following by assembly into contigs. In addition, deep sequencing and genome walking were also undertaken to recover more genomic sequences. The 5′ and 3′ termini were sequenced using a 5′ Full RACE Kit with TAP and a 3′ Full RACE Core Set with PrimeScript RTase (TaKaRa, Dalian, China). Primer sequences for full-length genome amplification are shown in Supplementary Table S1.

Genomic and phylogenetic analyses
Genomic structures of the CoV complete sequences were predicted by the SeqBuilder program of the DNAStar software package and compared with other representatives from GenBank. Nonstructural proteins (nsps) in ORF1a and ORF1b (replicase) of the CoVs were predicted using Z-Curve version 2.0, a CoV-specific gene-finding system (Gao et al, 2003). All 400-bp amplicons (the primer truncation of 440-nt sequences) were aligned with their closest phylogenetic neighbors in GenBank using Clustal W version 2.0. The phylogenetic tree was then constructed by the maximum likelihood method of MEGA 6.06 with 1,000 bootstrap replications. To better understand their evolutionary relationships, the complete RdRP genes were further amplified and used for the analysis.

Nucleotide sequence accession numbers
The partial RdRp sequences obtained from all positive samples and the complete genome or full length RdRp sequences of some specimens were submitted to the GenBank under accession numbers KU182954 to KU183005.

RESULTS

Detection of CoVs
Of 951 bats tested 50 intestinal specimens (5.3%) were CoV positive, but surprisingly all respiratory specimens showed negative amplification. As shown in Table 1, among 181 bats from 6 species in 3 families in Guangdong province, 16.2% (6/37) Rousettus leschenaulti and 27.5% (14/51) Cynopterus sphinx were CoV positive. Among 599 bats from 17 species in 5 families in Yunnan province, 14.0% (14/100) Rousettus leschenaulti, 2.4% (1/41) Megaerops kusnotei, 9.0% (7/78) Rhinolophus sinicus and 5.3% (5/95) Myotis daubentoni were CoV positive. As the first study of this kind in the Tibet Autonomous Region, fifteen Hipposideros cineraceus and five Rhinolophus hipposideros collected in south Tibet were tested and only 6.7% (1/15) Hipposideros cineraceus showed positive amplification. In northeast China, 2 of 97 (2.1%) bats in Jilin province were positive: one from Murina leucogaster and another from Rhinolophus ferrumequinum. In contrast, all 16 Rhinolophus ferrumequinum and 38 Myotis ricketti in Liaoning
province showed negative amplification. These results revealed a higher CoV incidence in three fruit bat species of the family *Pteropodidae* than in the four insectivorous bat families, indicating that fruit bats are more likely to harbor CoVs.

**Phylogenetic analysis**

To describe the genetic relationships among the 50 sequences obtained in this study and previously known CoVs, 400-nt *RdRP* sequences were obtained from the primer truncation of 440-nt sequences and phylogenetically analyzed. Results showed that 8 sequences grouped into 3 clusters within the genus *αCoV* (Figure 1A). YDB5C is the first reported bat-borne CoV (*Hipposideros cineraceus*) in Tibet and clustered closely with MLHJC4, a CoV from *Rhinolophus sinicus* in Yunnan, both sharing 94% nt identity with previously reported strain HKU2/GD/430/2006 from Guangdong (Lau et al, 2007). JTAC2

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**Table 1. Bat sample collection and coronavirus detection.**

| Family            | Species          | Bat² | CoV | Bat | CoV | Bat | CoV | Bat | CoV | Bat | CoV |
|-------------------|------------------|------|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| *Pteropodidae*    | *Rousettus*      | 6/37 (16.2) | β4  | 14/100 (14.0) | β4 |
|                   | *leschenaulti*   | 14/51 (27.5) | β4  |     |     |     |     |     |     |     |     |
|                   | *Cynopterus*     | 1/41 (2.4)  | β4  |     |     |     |     |     |     |     |     |
|                   | *kusnotei*       | 0/9   |     | 1/15 (6.7)  | α  |
| *Hipposideridae*  | *Hipposideros*   | 0/9   |     |     |     |     |     |     |     |     |     |
|                   | *cineraceus*     | 0/9   |     |     |     |     |     |     |     |     |     |
|                   | *pomona*         | 0/84  |     |     |     |     |     |     |     |     |     |
|                   | *larvatus*       | 0/68  |     | 0/2  |     |     |     |     |     |     |     |
|                   | *armiger*        | 0/11  |     | 0/18 |     |     |     |     |     |     |     |
|                   | *Aselliscus*     | 0/17  |     |     |     |     |     |     |     |     |     |
|                   | *stoliczkanus*   | 0/33  |     |     |     |     |     |     |     |     |     |
| *Rhinolophidae*   | *Rhinolophus*    | 0/42  |     | 0/16 |     | 1/30 (3.3) | β2 |
|                   | *ferrumequinum*  | 7/78 (9.0) | α, β2 |     |     |     |     |     |     |     |     |
|                   | *sinicus*        | 0/5   |     | 0/6  |     |     |     |     |     |     |     |
|                   | *pusillus*       | 0/3   |     |     |     |     |     |     |     |     |     |
|                   | *affinis*        | 0/37  |     | 0/5  |     |     |     |     |     |     |     |
| *Vespertilionidae*| *Myotis*         | 5/95 (5.3) | β3  |     |     |     |     |     |     |     |     |
|                   | *daubentonii*    | 0/8   |     |     |     |     |     |     |     |     |     |
|                   | *laniger*        | 0/3   |     |     |     |     |     |     |     |     |     |
|                   | *chinensis*      | 0/40  |     |     |     |     |     |     |     |     |     |
|                   | *capaccinii*     | 0/38  |     | 0/27 |     |     |     |     |     |     |     |
|                   | *ricketti*       | 0/8   |     |     |     |     |     |     |     |     |     |
|                   | *schreibersi*    | 1/40 (2.5) | α  |     |     |     |     |     |     |     |     |
|                   | *leucogaster*    | 0/1   |     |     |     |     |     |     |     |     |     |
| *Megadermatidae*  | *Megaderma*      | 0/1   |     |     |     |     |     |     |     |     |     |
|                   | *lyra*           |       |     |     |     |     |     |     |     |     |     |

Note: ²positive/total bats; numbers in brackets indicate the coronavirus positive percentage. ³CoV, α: αCoV; β: unclassified βCoV; β2: βCoV lineage 2; β3: βCoV lineage 3; β4: βCoV lineage 4.
identifying in *Murina leucogaster* in Jilin province diverged considerably from known CoVs, showing the highest nt identity of only 83% to bat-borne coronavirus Neixiang-14 and Neixiang-52 detected also in *Murina leucogaster*, and followed by 78% nt identity with some pandemic porcine epidemic diarrhea virus (PEDV) strains that have emerged recently in China, USA and Japan (Vlasova et al, 2014; Sun et al, 2015; Suzuki et al, 2015). Five other αCoVs (MLHJC1, MLHJC6, MLHJC8, MLHJC22, MLHJC34) identified from *Rhinolophus sinicus* in Yunnan formed a new group with MLHJC8 being slightly more divergent, showing highest nt identities (75%–89%) with the previously reported BtCoV/860/2005 (Tang et al, 2006). The remaining 42 bat CoV sequences were classified as βCoV and fell into 5 clusters (Figure 1B). Twenty identified in Guangdong fell into lineage β4, which showed the geographical relationship and was further divided into two distinct clusters, one with 6 sequences sharing 99% highest nt identity with HKU9-10-1 (Lau et al, 2010), while another including 14 sequences sharing the closest relationship with BtCoV/BRT55629/H.lek/CK/Tha/05/2012 detected in *Hippoposideros lekaguli* in Thailand (Wacharapluesadee et al, 2015). The 21 βCoVs identified in Yunnan province exhibited considerable genetic diversity and were distributed among lineages β2, β3 and β4. Fifteen fell into β4 and further divided into 2 lineages, fourteen sequences showing closest relationship to previously reported BtCoV/BRT55629/H.lek/CK/Tha/05/2012 (Wacharapluesadee et al, 2015), while another (ML92C) grouped with the
above Guangdong sequences. Five sequences detected from *Myotis daubentoni* clustered within lineage β3, sharing > 91% nt identities with previously reported HKU4-4 from *Tylonycteris pachypus* (Woo et al, 2007). This group showed about 80% nt identity with MERS-CoVs recently isolated in China (Lu et al, 2015) and Korea (Kim et al, 2015). The remaining Yunnan bat CoV sequence, MLHJC35, detected in *Rhinolophus sinicus*, was 97% identical with SARSr-BatCoV Cp/Yunnan2011 previously isolated in Yunnan province (Yang et al, 2011), while JTMC15 shared 99% identity with SARSr-BatCoV Rf1 found in *Rhinolophus ferrumequinum*, Hubei province (Li et al, 2005).

To obtain more precise analysis, representative specimens of the 8 phylogenetic clusters were subjected to full RdRP gene amplification. Complete RdRP sequences were obtained with 13 specimens belonging to 6 clusters comprised of 3 αCoVs and 10 βCoVs. Phylogenetic analysis based on the full RdRP gene sequences was highly consistent with Figure 1 (Phylogenetic tree of the full RdRP gene sequences is not shown).

**Full genomic sequences characterization**

Full genomic sequencing was successful in 2 of the above 13 specimens: JTMC15 from *Rhinolophus ferrumequinum*, Jilin province, and JPDB144 from *Myotis daubentoni*, Yunnan province, with a nearly complete genome sequence obtained of JTAC2 from *Murina leucogaster*. The full genomes of JTMC15 and JPDB144 (including complete terminal sequences of 5′ end and 3′-poly A) and near-complete genome of JTAC2 were 28,761 nt, 30,321 nt and 25,719 nt in size respectively, with G+C contents of 38.1%, 41.0% and 43.4%. It is proved that two proteinases, papain-like proteinase (PLPro) encoded by nsp3 gene and main proteinase (MPro) encoded by nsp5 gene in ORF1a of CoVs are able to cleave the complex of ORF1a and ORF1b (replicase) into 16 mature nonstructural proteins (nsp5) (Neuman et al, 2008). Our analysis of the nsp5 in ORF1ab revealed that all the three bat CoV genomic sequences contain 16 nsps (nsp1–nsp16) in ORF1ab, but the cleavage sites are different for nsp3 or nsp5 in different CoVs. The length of deduced amino acids of putative nsps, their first-last residue and position in replicase are shown in Supplementary Table S2.

Base on the nearly complete genomic sequence obtained, JTAC2 possesses the same genome structure as PEDVs with 7 genes in the order: 5′-ORF1a, 1b, spike (S), 3a, envelope (E), membrane (M) and nudeocapsid (N)-3′ (Figure 2A). JTAC2 showed the nearest relationship (87.9% in ORF1a and 92.8% in ORF1b) with Lushi M1 bat CoV isolates Neixiang-14 and Neixiang-52, but the latter two have very limited sequences available for further analysis. The recent PEDV-1C isolated from a piglet with diarrhea and vomiting (Sun et al, 2015) was therefore used for sequence comparison and genomic organization analysis, since it has been fully sequenced and shares high identity with JTAC2 (Figure 1A and Supplementary Table S3). The aa identity comparison shown in Supplementary Table S3 suggests that JTAC2 is a novel αCoV.

JTMC15 is a SARSr-BatCoV having the same genome organization as other SARSr-BatCoVs (e.g., Rf1), but sequence deletions were observed in ORF1a and N, and between genes 7b–8. A 579-nt deletion in ORF1a of JTMC15 was also observed in SARSr-BatCoV Rs672 from a *Rhinolophus sinicus* bat (Yuan et al, 2010) and a human SARS-CoV ShanghaiQXC2 from the late phase of the 2003 epidemic (GenBank #AY463060). This 579-nt deletion results in a 193-aa deletion of nsp3 in ORF1a, from residues 1059 to 1251 in the nucleic acid-binding (NAB) domain (Serrano et al, 2009). A second deletion in the N gene of JTMC15 (1156–1158 nt, one residue Q108) was also found in 3 SARSr-BatCoV strains, Rp/Shaanxi2011 (Yang et al, 2013), Rm1 (Li et al, 2005) and 279/2005 (Tang et al, 2006). Interestingly, four discontinuous deletions were identified in JTMC15 between genes 7b and 8, which is unique in JTMC15, resulting in an ORF shift and elimination of gene 8 (Figure 2B). Similar to known CoVs, extensive S gene variations were also observed in JTMC15, resulting in low aa identities with other SARSr-BatCoV strains (the highest being 86.1% to Rf1) as compared with other gene fragments in the genome. Receptor-binding motif (RBM) is an extended loop that lies on the surface of the receptor binding domain (RBD) of the spike protein, and is the most important domain for SARSr-BatCoV to recognize its host receptor, angiotensin-converting enzyme 2 (ACE2) (Ren et al, 2008; Baez-Santos et al, 2015). Further alignment of the deduced amino acid sequences of RBM (55 aa) showed a closer relationship of JTMC15 to SARSr-BatCoVs than to human or civet SARS-CoVs (Supplementary Figure S1). Taking the above altogether, as shown in Figure 2A and Table 2, there are 13 genes predicted in JTMC15: 5′-ORF1a, 1b, S, 3a, 3b, E, M, 6, 7a, 7b, N, 9a, 9b-3′. Apart from gene 7b (83.0%) and S (86.1%) all ORFs of JTMC15 had high aa identities to Rf1, ranging from 94.4% (9b gene) to 99.1% (M gene), indicating that JTMC15 is a new variant within the SARSr-BatCoV Rf1 species.

For JPDB144, the genome organization is almost the same as HKU4-4, with 10 genes in the order: 5′-ORF1a, 1b, S, 3a, 3b, 3c, 3d, E, M, N-3′ (Figure 2A). However, two differences were observed in the nsp2 of JPDB144:
a 12-nt insertion (residues 1143 to 1146 of \(1a\)) and a 3-nt deletion (residue 1155 of \(1a\)). Other JPDB144 ORFs were the same as HKU4-4 in length, sharing aa identities of between 88.8% (\(3c\) gene) and 98.8% (\(E\) gene); however, an aa sequence comparison of JPDB144 ORFs to those of HKU5 and MERS-CoV strains in the Betacoronavirus lineage 3 showed rather low similarities (Supplementary Table S4).

**DISCUSSION**

As shown in Figure 1, diverse αCoVs and βCoVs have been identified in the present study from different bats sampled at 25 locations in 4 provinces and the Tibet Autonomous Region, demonstrating the wide distribution of CoVs among a range of bat species. Of 8 αCoVs identified, YDB5C is the first bat-borne CoV identified in the Tibet and Himalayan area, detected in 1 of 15 *Hipposideros cineraceus* bats collected in Yadong county of Tibet, located at the southern edge of the Himalayas bordering on Bhutan and India. Another newly identified CoV, MLHJC4, was detected in *Rhinolophus sinicus* in Yunnan province, which phylogenetically clustered closely with YDB5C, both showing 94% nt identity to HKU2/GD/430/2006 identified in Guangdong (Lau et al, 2007), indicating that this type of αCoV has a wide range of bat reservoirs and geo-distribution in south-west China and perhaps neighboring regions. In addition, six other αCoV sequences JTAC2, MLHJC2, MLHJC6, MLHJC8, MLHJC22 and MLHJC34, found in this study clustered as two novel CoV groups. Although not novel, the βCoVs identified here showed abundance in genetic and geographical diversities. It is interesting to note that SARSr- and MERS-like CoVs were identified, particularly JTMC15 isolated in Jilin province – the first SARSr-Bat CoV to be discovered in Northeast China.

In consideration of bat species, sampling locations and CoVs diversities, four pathogen/host/environment situations can be proposed. First, a single bat species at one location (even a single cave) harboring different CoV species (e.g., *Rhinolophus sinicus* collected at the same site in Menglian county, Yunnan province, harboring three CoV species: HKU2-like, SARSr- and new αCoVs). Second, a single bat species roosting at different locations harboring the same CoVs (e.g., *Rousettus leschenaulti* collected in Mengla county (south Yunnan) and Wand-...
ing county (west Yunnan) harboring the same βCoVs). Third, multiple bat species sampled at the same site harboring the same CoVs (e.g., *Rousettus leschenaulti* and *Megaerops kusnotei* sampled at the same location in Wanding county harboring the same βCoVs). Different bat species collected at different locations may even harbor the same CoVs: e.g., a *Hipposideros cineraceus* in Yadong, Tibet and a *Rhinolophus sinicus* in Menglian, Yunnan harbored HKU2-like viruses, and another *Rhinolophus sinicus* in Menglian and a *Rhinolophus ferrumequinum* in Tonghua county, Jilin province, harbored SARSr CoVs. Altogether, the data provided further evidence for the wide distribution of CoVs among bat populations in China, and for the suggestion that different CoVs employ different bat species as reservoirs.

The present study has identified genetically diverse bat-borne CoVs, which were detected from intestinal tissue specimens of different bat species of wide geographic distribution. But we failed to detect any CoV sequence from the respiratory specimens that probably due to the low virus load in the lung or specific intestinetropism of CoV in bats. Bats are considered the gene source of *Alphacoronavirus* and *Betacoronavirus* (Woo et al, 2012), especially of pathogenic CoVs that cause public threats. In last decade, increasing number of CoVs have been identified in bats, in which the viral genes were presumably originated and evolved with high mutation and recombination rates (Woo et al, 2007). It is apparent that large numbers of circulating CoVs remain unidentified, and are evolving worldwide within the bat population. Investigations in unexplored regions are therefore urgently needed to gain further insights into CoV diversity and evolutionary dynamics.

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**COMPLIANCE WITH ETHICS GUIDELINES**

The authors declared that they have no conflict of interest. The whole study was approved by the Administrative Committee on Animal Welfare of the Institute of Military Veterinary, Academy of Military Medical Sciences, China (Laboratory Animal Care and Use Committee Authorization, permit number JSY-DW-2010-02). All institutional and national guidelines for the care and use of laboratory animals were followed.

**AUTHOR CONTRIBUTIONS**

CT conceived the study and LX carried it out with BH’s
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guidance. FZ, WY, TJ, GL, TH, GC, YF, YZ, QF, JF and HZ were responsible for field investigation and bat sampling. TJ and GL identified bat species morphologically. XL took part in samples screening and CoVs detection. LX wrote the paper, CT and BH then revised it. All authors read and approved the final manuscript.

Supplementary figures/tables are available on the website of Virologica Sinica: www.virosin.org; link.springer.com/journal/12250.

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