

1 **Pangolin homology associated with 2019-nCoV**

2 Tao Zhang^{1*}, Qunfu Wu^{1*}, Zhigang Zhang^{1#}

3 ¹State key Laboratory for Conservation and Utilization of Bio-Resources in Yunnan,

4 School of Life Sciences, Yunnan University, Kunming, Yunnan, 650091, China

5 *These authors contributed equally

6 #Correspondence to: zhangzhigang@ynu.edu.cn (Z.Z.G.)

1 **Abstract**

2 To explore potential intermediate host of a novel coronavirus is vital to rapidly control
3 continuous COVID-19 spread. We found genomic and evolutionary evidences of the
4 occurrence of 2019-nCoV-like coronavirus (named as Pangolin-CoV) from dead Malayan
5 Pangolins. Pangolin-CoV is 91.02% and 90.55% identical at the whole genome level to
6 2019-nCoV and BatCoV RaTG13, respectively. Pangolin-CoV is the lowest common
7 ancestor of 2019-nCoV and RaTG13. The S1 protein of Pangolin-CoV is much more
8 closely related to 2019-nCoV than RaTG13. Five key amino-acid residues involved in the
9 interaction with human ACE2 are completely consistent between Pangolin-CoV and 2019-
10 nCoV but four amino-acid mutations occur in RaTG13. It indicates Pangolin-CoV has
11 similar pathogenic potential to 2019-nCoV, and would be helpful to trace the origin and
12 probable intermediate host of 2019-nCoV.

1 In the late December of 2019, an epidemic pneumonia (the W.H.O. announced it
2 recently as Corona Virus Disease (COVID-19)(1)) outbreak in city of Wuhan in China and
3 soon widely spreads all over the world. According to authoritative statistics, the COVID-
4 19 has caused more than 40,000 laboratory-confirmed infections with more than 1000
5 deaths by 12 February 2020 and is still increasing. It was caused by a novel identified
6 coronavirus 2019-nCoV (the International Committee on Taxonomy of Viruses (ICTV)
7 renamed this virus as severe acute respiratory syndrome coronavirus 2, SARS-CoV-2 (1)).
8 Released complete genomes of 2019-nCoVs (2, 3) have helped rapid identification and
9 diagnosis of the COVID-19. Another key task is to find potential origin of 2019-nCoV(4).
10 Unsurprisingly, like SARS-CoV and MERS-CoV(5), the bat is still a probable origin of the
11 2019-nCoV because the 2019-nCoV shared 96% whole genome identity with a bat
12 coronavirus Bat-CoV-RaTG13 from *Rhinolophus affinis* from Yunnan Province(2).
13 However, SARS-CoV and MERS-CoV usually pass into medium host like civets or camels
14 before leaping to humans(4). It indicates that the 2019n-Cov was probably transmitted to
15 humans by some other animals. Considering the earliest COVID-19 patient reported no
16 exposure at the seafood market(6), finding intermediate host of 2019-nCoV is vital to block
17 its transmission.

18 On 24 October 2019, Liu and his colleagues from the Guangdong Wildlife Rescue
19 Center of China (7) firstly detected the existence of SARS-liked coronavirus from lung
20 samples of two dead Malayan Pangolins with a frothy liquid in lung and pulmonary fibrosis,

1 which is close on the outbreak of COVID-19. From their published results, all virus contigs
2 assembled from 2 lung samples (lung07, lung08) showed not high identities ranging from
3 80.24% to 88.93% with known SARS coronavirus. Hence, we conjectured that dead
4 Malayan pangolin may carry a new coronavirus close to 2019-nCoV.

5 To confirm our assumption, we downloaded raw RNA-seq data (SRA accession
6 number PRJNA573298) of those two lung samples from SRA and conducted consistent
7 quality control and contamination removing as described by Liu's study(7). We found 1882
8 clean reads from lung08 sample mapped upon 2019-nCoV reference genome (GenBank
9 Accession MN908947)(3) with high genome coverage of 76.02%. We performed *de novo*
10 assembly of those reads and totally obtained 36 contigs with length ranging from 287bp to
11 2187bp with mean length of 700bp. Blasting against proteins from 2845 coronavirus
12 reference genomes including RaTG13, 2019-nCoVs and other known coronaviruses, we
13 found 22 contigs can be best matched to 2019-nCoVs (70.6%-100% aa identity; average:
14 95.41%) and 12 contigs matched to Bat SARS-like coronavirus (92.7%-100% aa identity;
15 average: 97.48%) (Table S1). These results indicate Malayan pangolin indeed carries a
16 novel coronavirus (here named as Pangolin-CoV) close to 2019-nCoV.

17 Using reference-guided scaffolding approach, we created Pangolin-CoV draft
18 genome (19,587 bp) based on the above 34 contigs. Remapping 1882 reads against the draft
19 genome resulted in 99.99% genome coverage at a mean depth of 7.71 X (range: 1X-47X)
20 (**Figure 1A**). Based on Simplot analysis, Pangolin-Cov showed highly overall genome

1 sequence identity throughout the genomes to RaTG13 (90.55%) and 2019-nCoV (91.02%)
2 (**Figure 1B**), although there is greatly high identity 96.2% between 2019-nCoV and
3 RaTG13(3). Another SARS-like coronavirus more similar to Pangolin-Cov were Bat
4 SARSr-CoV ZXC21(85.65%) and Bat SARSr-CoV ZC45 (85.01%). These results indicate
5 Pangolin-Cov may be the common origin of 2019-nCoV and RaTG13.

6 The viral genome organization of Pangolin-Cov was characterized by sequence
7 alignment against 2019-nCoV (GenBank Accession MN908947) and RaTG13. The
8 Pangolin-Cov genome consists of six major open reading frames (ORFs) common to
9 coronaviruses and other four accessory genes (**Figure 1C** and **Table S2**). Further analysis
10 indicates that Pangolin-Cov genes covered 2019-nCoV genes with coverage ranging from
11 45.8% to 100% (average coverage 76.9%). Pangolin-Cov genes shared high average
12 nucleotide and amino identity with both 2019-nCoV (MN908947) (93.2% nt/ 94.1% aa)
13 and RaTG13 (92.8% nt/93.5% aa) (**Figure 1C** and **Table S2**). Surprisingly, some of
14 Pangolin-Cov genes showed higher aa sequence identity to 2019-nCoV than RaTG13,
15 including orf1b (73.4/72.8), S-protein (97.5/95.4), orf7a (96.9/93.6), and orf10 (97.3/94.6).
16 High S-protein amino acid identity implies function similarity between Pangolin-Cov and
17 2019-nCoV.

18 To determine the evolutionary relationships among Pangolin-Cov, 2019-nCoV and
19 previously identified coronaviruses, we estimated phylogenetic trees based on the
20 nucleotide sequences of the whole genome sequence, RNA-dependent RNA polymerase

1 gene (RdRp), non-structural protein genes ORF1a and 1b, and the main structural proteins
2 encoded by the S and M genes. In all phylogenies, Pangolin-CoV, RaTG13 and 2019-nCoV
3 were clustered into a well-supported group, here named as “SARS-CoV-2 group” (**Figure**
4 **2 and Figures S1 to S2**). This group represents a novel Beta-coronaviruses group. Within
5 this group, RaTG13 and 2019-nCoV was grouped together, and the Pangolin-CoV was
6 their lowest common ancestor. However, whether the basal position of the SARS-CoV-2
7 group is SARSr-CoV ZXC21 and/or SARSr-CoV ZC45 or not is still in debate. Such
8 debate also occurred in both Wu et al.(3) and Zhou et al. (2)studies. Possible explanation
9 is due to a past history of recombination in Beta-CoV group(3). It is noteworthy that our
10 discovered evolutionary relationships of coronaviruses shown by the whole genome, RdRp
11 gene, and S-gene were highly consistent with that discovered by complete genome
12 information in Zhou et al. study(2). It indicates our Pangolin-CoV draft genome has enough
13 genomic information to trace true evolutionary position of Pangolin-CoV in coronaviruses.

14 The coronavirus spike (S) protein consisting of 2 subunits (S1 and S2) mediates
15 infection of receptor-expressing host cells and is a critical target for antiviral neutralizing
16 antibodies. S1 contains a receptor binding domain (RBD) about 193 amino acid fragment,
17 which is responsible for recognizing and binding with the cell surface receptor(8, 9). Zhou
18 et al. had experimentally confirmed that 2019-nCoV is able to use human, Chinese
19 horseshoe bats, civet, and pig ACE2 as an entry receptor in the ACE2-expressing cells(2),
20 suggesting the RBD of 2019-nCoV mediates infection to human and other animals. To gain

1 a sequence-level insight into understanding pathogen potential of Pangolin-Cov, we
2 investigated the amino acid variation pattern of S1 protein from Pangolin-CoV, 2019-
3 nCoV, RaTG13, and other representative SARS-CoVs. The amino acid phylogenetic tree
4 showed the S1 protein of Pangolin-CoV is more closely related to that of 2019-CoV than
5 RaTG13. Within the RBD, we further found Pangolin-CoV and 2019-nCoV was highly
6 conserved with only one amino acid change (500H/500Q) (**Figure 3**) but not belonging to
7 five key residues involved in the interaction with human ACE2(2, 9). In contrast, RaTG13
8 has 17 amino acid residue changes and 4 of them belonged to key amino acid residue
9 (**Figure 3**). These results indicate Pangolin-Cov has similar pathogen potential to 2019-
10 nCoV.

11 The nucleocapsid protein (N-protein) is the most abundant protein in coronavirus.
12 The N-protein is a highly immunogenic phosphoprotein, and it is normally very conserved.
13 The N protein of coronavirus is often used as a marker in diagnostic assays. To gain a
14 further insight into the diagnostic potential for Pangolin-Cov, we investigated the amino
15 acid variation pattern of N-protein from Pangolin-CoV, 2019-nCoV, RaTG13, and other
16 representative SARS-CoV. Phylogenetic analysis based on N protein supports Pangolin-
17 Cov as a sister taxon of 2019-nCoV and RaTG13 (**Figure 4**). We further found seven amino
18 acid mutations can differentiate our defined “SAR-CoV-2 group” (12N, 26 G, 27S, 104D,
19 218A, 335T, 346N, 350Q) from other known SARS-CoVs (12S, 26D, 27N, 104E, 218T,
20 335H, 346Q, 350N). Two amino acid sites (38P and 268Q) are shared by Pangolin-Cov,

1 RaTG13 and SARS-CoVs, which are mutated as 38S and 268A in 2019-nCoV. Only one
2 amino acid residue shared by Pangolin-CoV and other SARS-CoVs (129E) is consistently
3 changed in both 2019-nCoV and RaTG13 (129D). Our observed amino acid changes in N-
4 protein would be useful for developing antigen for much more sensitive serological
5 detection of 2019-nCoV.

6 Based on published metagenomic data, this study provides the first report on a
7 potential closely related kin (Pangolin-CoV) of 2019-nCoV, which was discovered from
8 dead Malayan Pangolins after extensive rescue efforts. Aside from RaTG13, the Pangolin-
9 CoV is the most closely related to 2019-nCoV. Due to original sample unavailable, we did
10 not perform further experiments to confirm our findings, including PCR validation,
11 serological detection, and even the isolation of virus particle etc. However, on 7 February,
12 researchers from the South China Agricultural University in Guangzhou reported pangolin
13 would be the potential candidate host of 2019-nCoV for isolating a virus 99% similar to
14 2019-nCoV in genome (Data unpublished). Our discovered Pangolin-CoV genome showed
15 91.02% nt identity with 2019-nCoV, implying Pangolin-CoV could be different from that
16 unpublished. Whether pangolin species is a good candidate for 2019-nCoV still need to be
17 further investigated. Considering the wide spread of SARSr-CoV in their natural reservoirs,
18 our findings would be meaningful to find novel intermediate hosts of 2019-nCoV for
19 blocking interspecies transmission.

1 **Materials and Methods**

2 Data preparation

3 We downloaded raw data of lung08 and lung07 published by Liu's study(7) from
4 NCBI sequence read archive (SRA) under Bio Project PRJNA573298. Raw reads were
5 first adaptor- and quality-trimmed using the Trimmomatic program (version 0.39)(10). For
6 removing host contamination, Bowtie2 (version 2.3.4.3) (11) was used to map clean reads
7 to the host reference genome of *Manis javanica* (NCBI Project ID: PRJNA256023). Only
8 unmapped reads were mapped to 2019-nCoV reference genome (GenBank Accession
9 MN908947) for identifying virus reads.

10 Read Assembly and construction of consensus sequence

11 Virus-targeted reads were assembled *de novo* using MEGAHIT (v1.1.3)(12). Read
12 remapping to assembled contigs was performed by using Bowtie2(11). Mapping coverage
13 and depth were produced using Samtools (version 1.9)(13). Contigs were taxonomically
14 annotated using BLAST 2.9.0+ against 2845 Coronavirus reference genomes (Table S1).
15 Bat-Cov-RaTG13 genome was downloaded from NGDC database (<https://bigd.big.ac.cn/>)
16 (Accession no. GWHABKP00000000)(2). 2019-nCoV reference genome was downloaded
17 from NCBI (Accession no. MN908947)(3). Other coronavirus genomes were downloaded
18 ViPR database (<https://www.viprbrc.org/brc/home.spg?decorator=corona>) on 6 February
19 2020. We further used reference-guided strategy to construct draft genome based on those
20 contigs taxonomically annotated to 2019-nCoVs, SARS-CoV, and Bat SARS-like CoV.

1 Each contig was aligned against 2019-nCoV reference genome with MUSCLE software
2 (version 3.8.31)(14). Aligned contigs were merged into consensus scaffold with BioEdit
3 version 7.2.5 ([http://www.mybiosoftware.com/bioedit-7-0-9-biological-sequence-](http://www.mybiosoftware.com/bioedit-7-0-9-biological-sequence-alignment-editor.html)
4 [alignment-editor.html](http://www.mybiosoftware.com/bioedit-7-0-9-biological-sequence-alignment-editor.html)) following manually quality checking. Small fragments with less
5 than length 25bp were discarded, if these fragments were not covered by any large
6 fragments. The potential open reading frames (ORFs) of finally obtained draft genome
7 were annotated by the alignment to 2019-nCoV reference genome (Accession no.
8 MN908947).

9 Phylogenetic relationship analysis

10 Sequence alignment was carried out using MUSCLE software(14). Alignment
11 accuracy was checked manually base-by-base. Gblocks(15) was used to process the gap in
12 aligned sequence. Using MegaX (version 10.1.7)(16), we inferred all maximum likelihood
13 (ML) phylogenetic trees under the best-fit DNA/amino acid substitute model with 1000
14 bootstrap replications. Phylogenetic analyses were performed using the nucleotide
15 sequences of various CoV gene data sets: the whole genome and ORF1a, ORF1b,
16 Membrane (M) gene, spike (S) and RNA-dependent RNA polymerase (RdRp) gene. The
17 best model of M is GTR+G and all others are GTR+G+I. Two additional protein-based
18 trees were constructed under WAG+G (S1 subunit of S protein) and JTT+G (N-protein),
19 respectively. Branches with values < 70% bootstrap were hidden in all phylogenetic trees.

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1 **Figure legends**

2 **Figure 1.** Genome-related analysis. **(A)** Sequence depth of mapped reads remapping to
3 Pangolin-Cov. **(B)** Similarity plot based on the full-length genome sequence of Pangolin-
4 Cov. Full-length genome sequences of 2019-nCoV (Beta-CoV/Wuhan-Hu-1), BatCov-
5 RaTG13, Bat SARSr-CoV 21, Bat SARSr-CoV45, Bat SARSr-CoV WIV1, and SARS-
6 CoV BJ01 were used as reference sequences. **(C)** Comparison of common genome
7 organization similarity among 2019-nCoV, Pangolin-Cov and BatCov-RaTG13 linked to
8 **Table S2.**

9 **Figure 2.** Phylogenetic relationship of coronavirus based on whole genome and RdRp gene
10 nucleotide sequences. Red text denotes the Malayan Pangolin-CoV. Pink text denotes
11 2019-nCoV (SARS-CoV-2). Green text denotes a bat coronavirus having 96% similarity
12 at genome level to SARS-CoV-2. Blue text denotes reference coronaviruses used in
13 **Figure1B.** Detailed information can be found in **Materials and Methods.**

14 **Figure 3.** Amino acid sequence alignment of the S1 protein and its phylogeny. The
15 receptor-binding motif of SARS-CoV and the homologous region of other coronaviruses
16 are indicated by the grey box. The key amino acid residues involved in the interaction with
17 human ACE2 are marked with the orange box. Bat SARS-like CoVs had been reported not
18 to use ACE2, had amino acid deletions at two motifs marked by the yellow box. Detailed
19 information can be found in **Materials and Methods.**

- 1 **Figure 4.** Amino acid sequence alignment of N protein and its phylogeny. Highly-
- 2 conserved amino acid residues in N-protein marked by colors have diagnostic potential.
- 3 Detailed information can be found in **Materials and Methods**.

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Figure 1

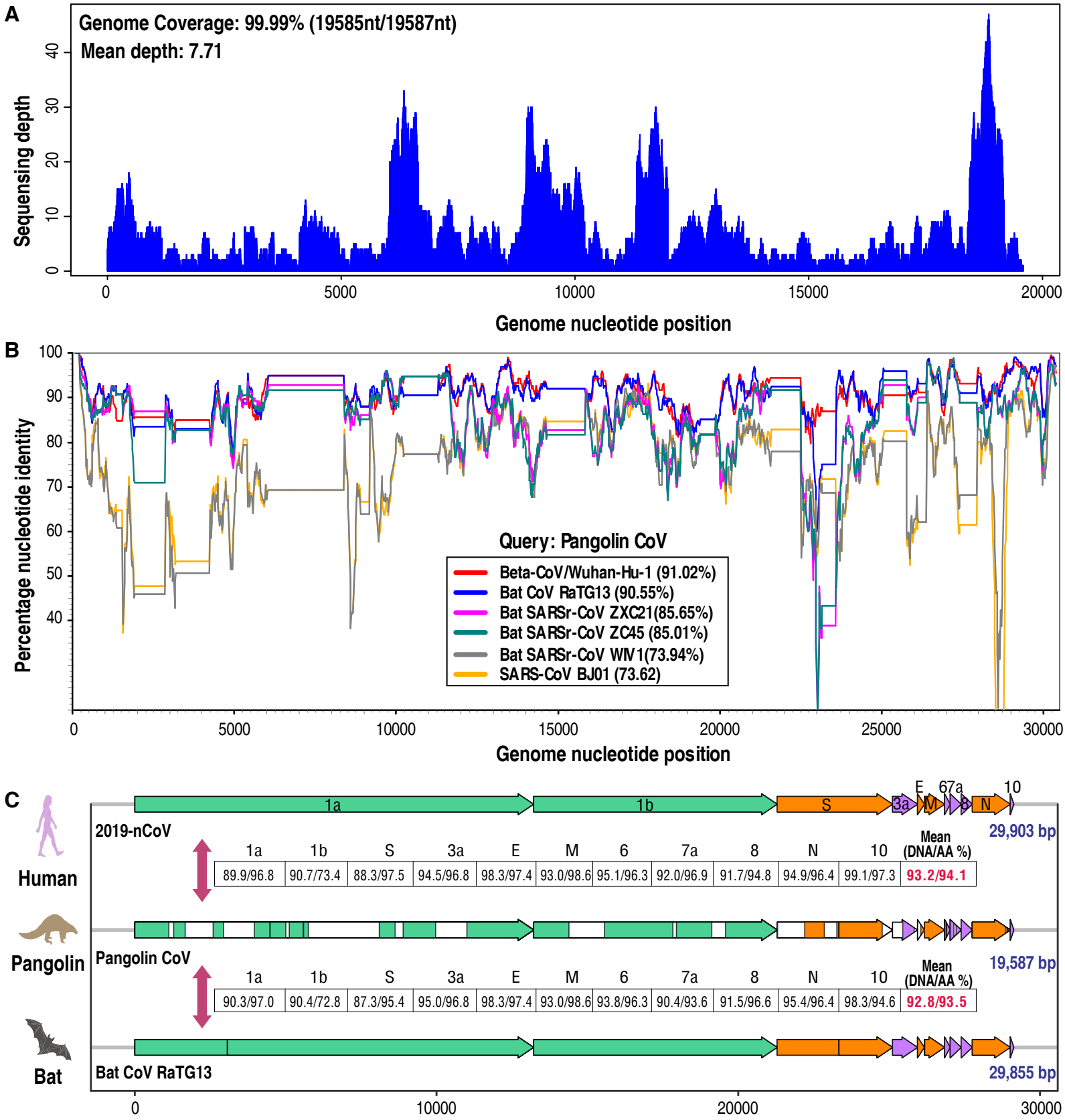


Figure 1. Genome-related analysis. (A) Sequence depth of mapped reads remapping to Pangolin-Cov. (B) Similarity plot based on the full-length genome sequence of Pangolin-Cov. Full-length genome sequences of 2019-nCoV (Beta-CoV/Wuhan-Hu-1), BatCov-RaTG13, Bat SARSr-CoV 21, Bat SARSr-CoV45, Bat SARSr-CoV WIV1, and SARS-CoV BJ01 were used as reference sequences. (C) Comparison of common genome organization similarity among 2019-nCoV, Pangolin-Cov and BatCov-RaTG13 I linked to **Table S2**.

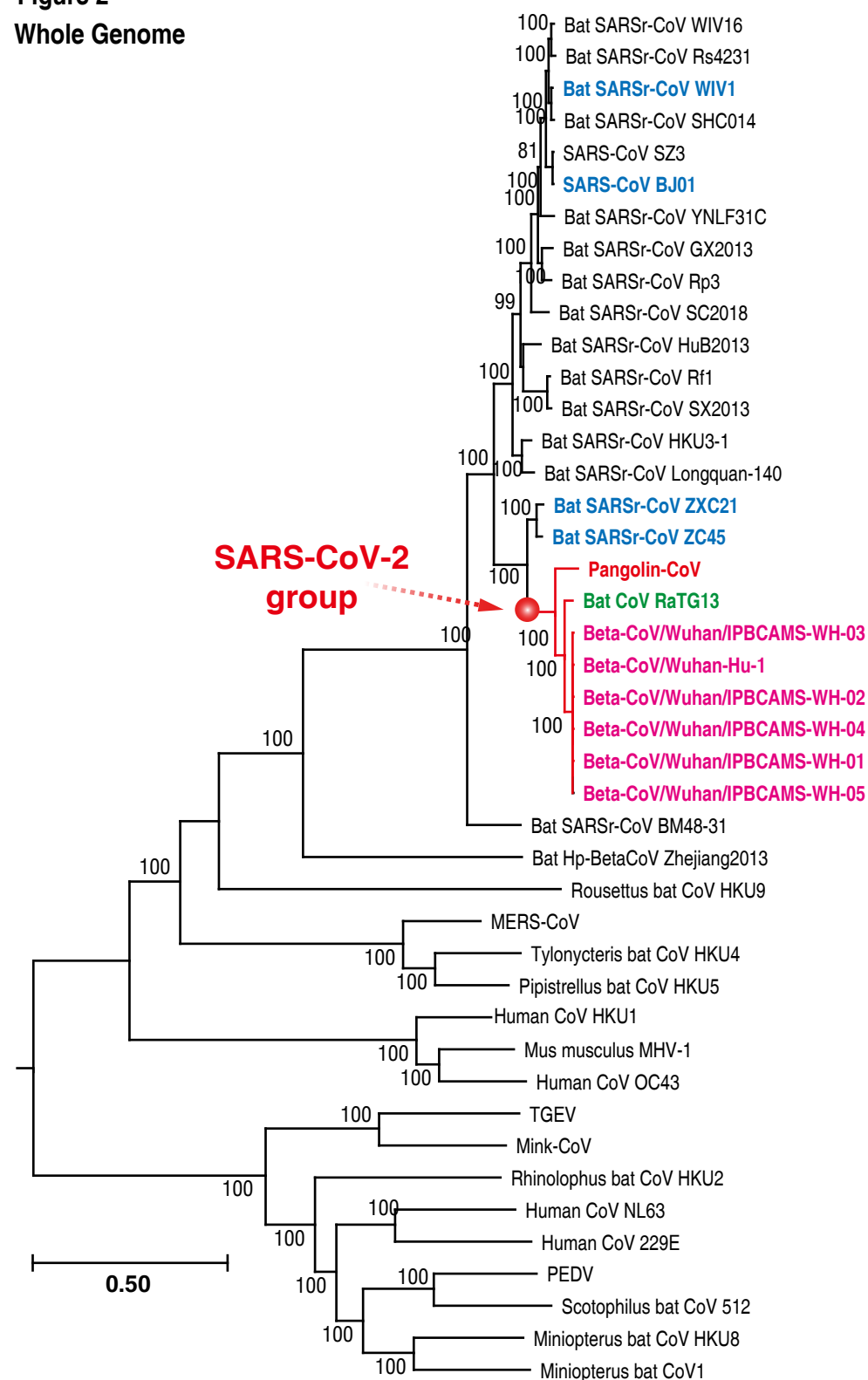
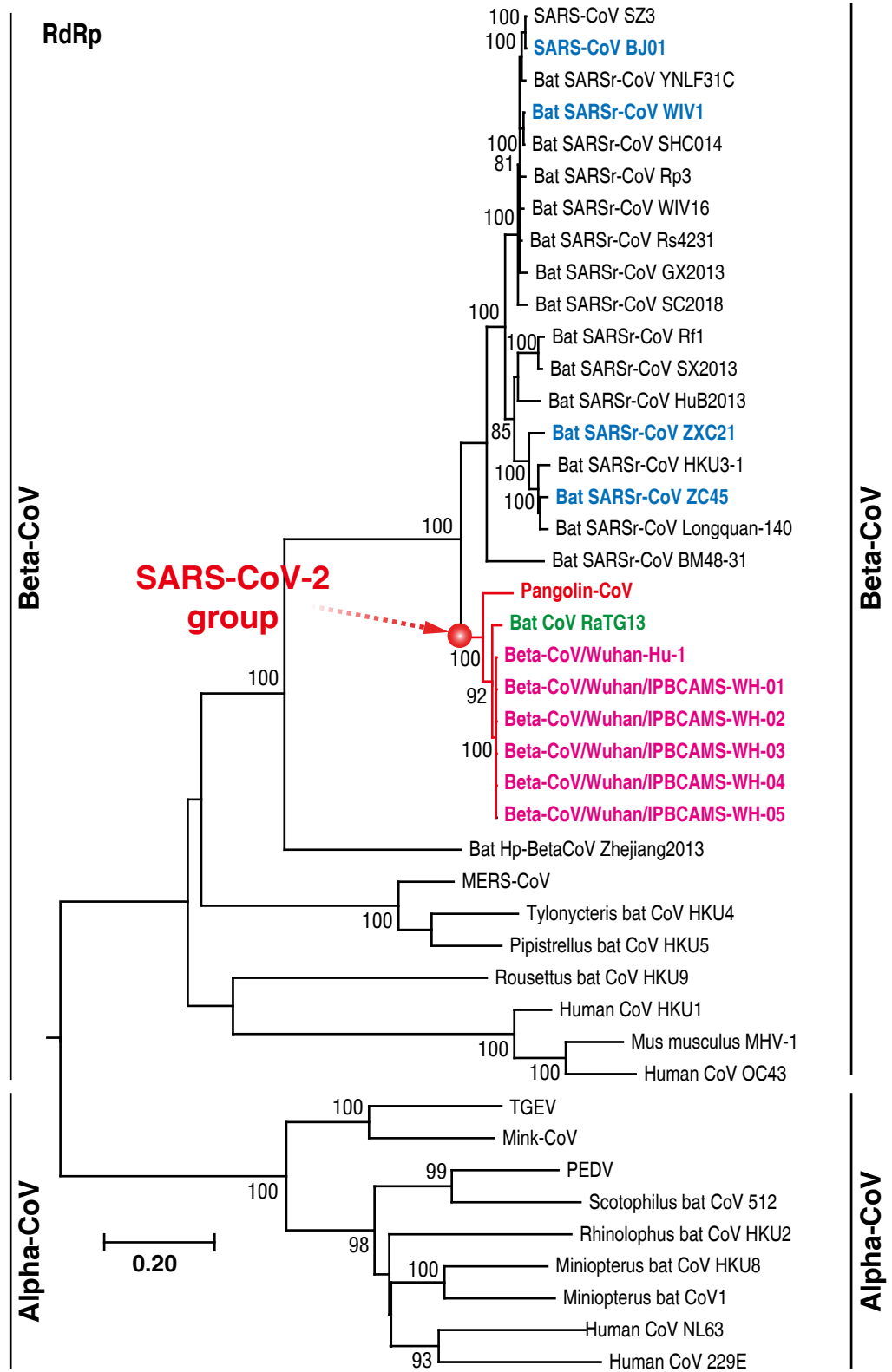
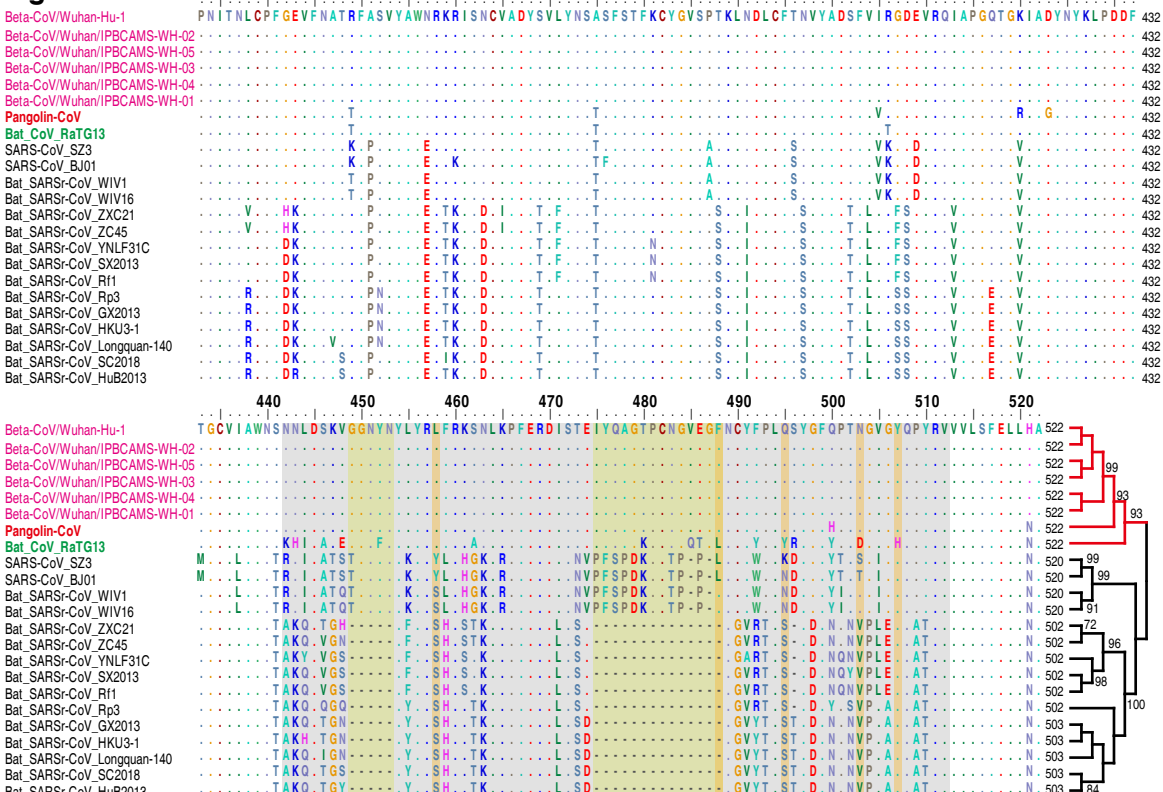
Figure 2**Whole Genome****RdRp**

Figure 2. Phylogenetic relationship of coronavirus based on whole genome and RdRp gene nucleotide sequences. Red text denotes the Malayan Pangolin-CoV. Pink text denotes 2019-nCoV (SARS-CoV-2). Green text denotes a bat coronavirus having 96% similarity at genome level to SARS-CoV-2. Blue text denotes reference coronaviruses used in Figure1B. Detailed information can be found in **Materials and Methods**.

Figure 3.**Figure 3. Amino acid sequence alignment of the S1 protein and its phylogeny.**

The receptor-binding motif of SARS-CoV and the homologous region of other coronaviruses are indicated by the grey box. The key amino acid residues involved in the interaction with human ACE2 are marked with the orange box. Bat SARS-like CoVs had been reported not to use ACE2, had amino acid deletions at two motifs marked by the yellow box. Detailed information can be found in **Materials and Methods**.

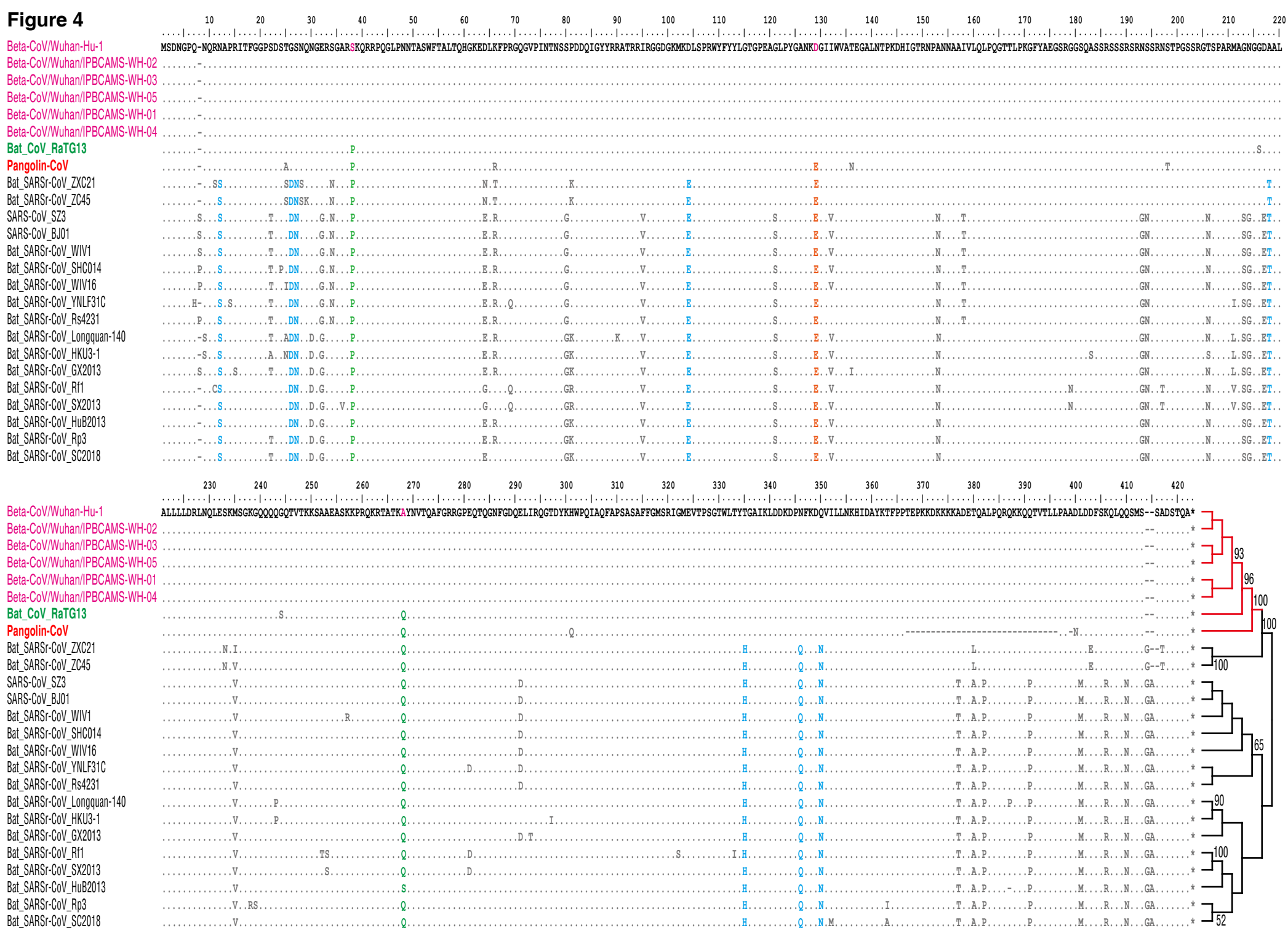


Figure 4. Amino acid sequence alignment of N protein and its phylogeny. Highly-conserved amino acid residues in N-protein marked by colors have diagnostic potential. Detailed information can be found in **Materials and Methods**.

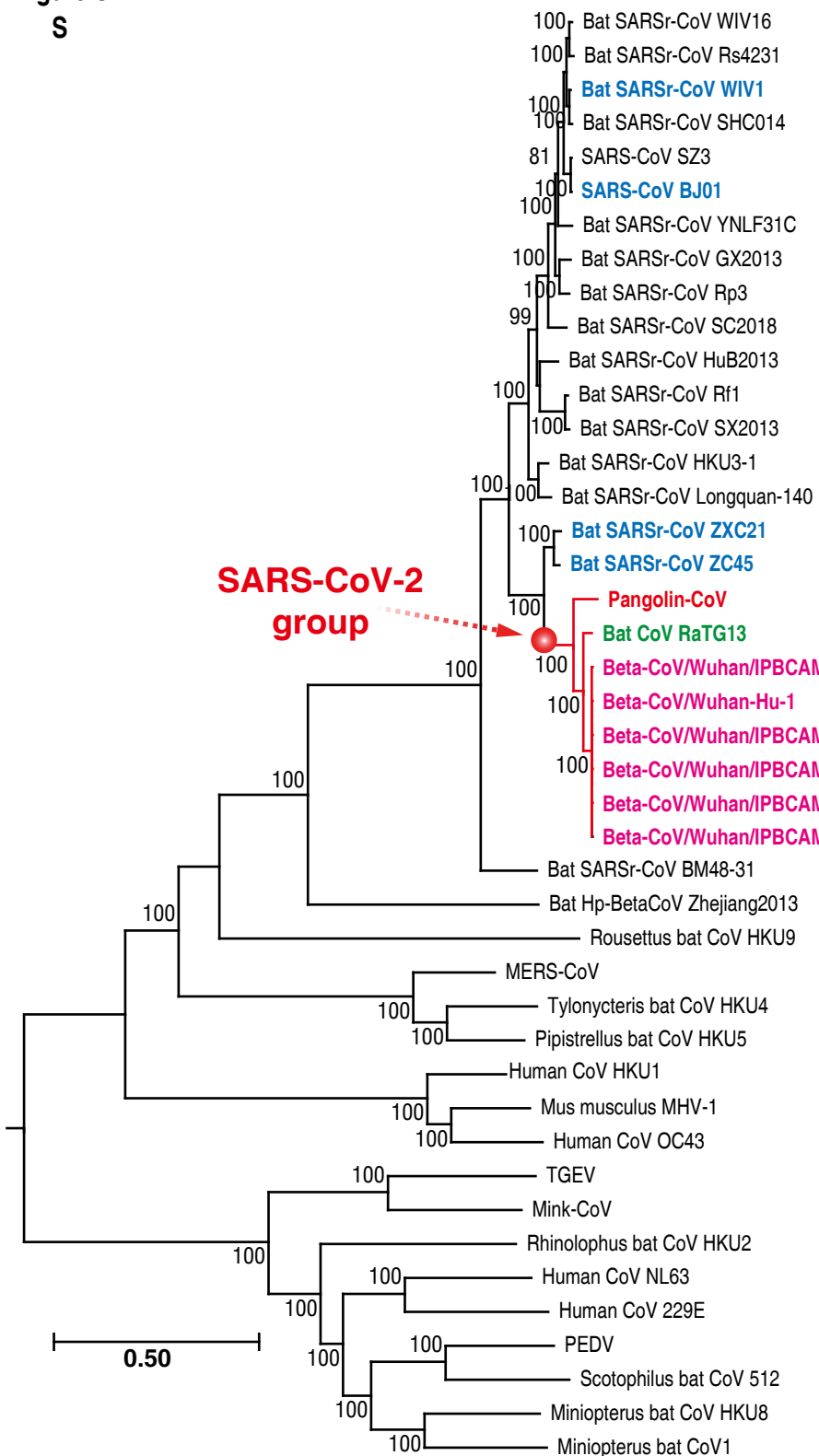
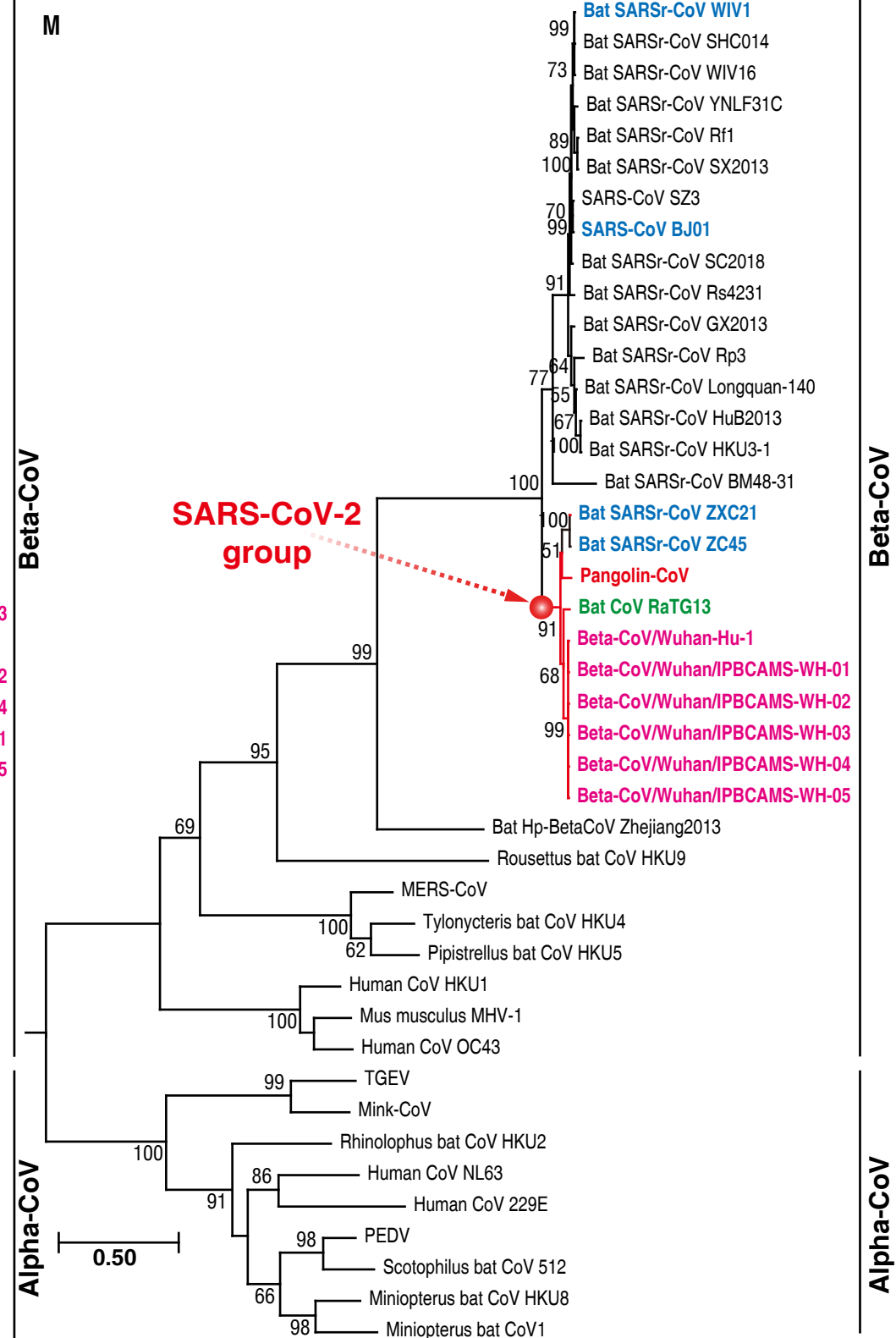
Figure S1**S****M****Figure S1.** Phylogenetic relationship of coronavirus based on ORF1a gene (A) and ORF1b gene (B) nucleotide sequences.

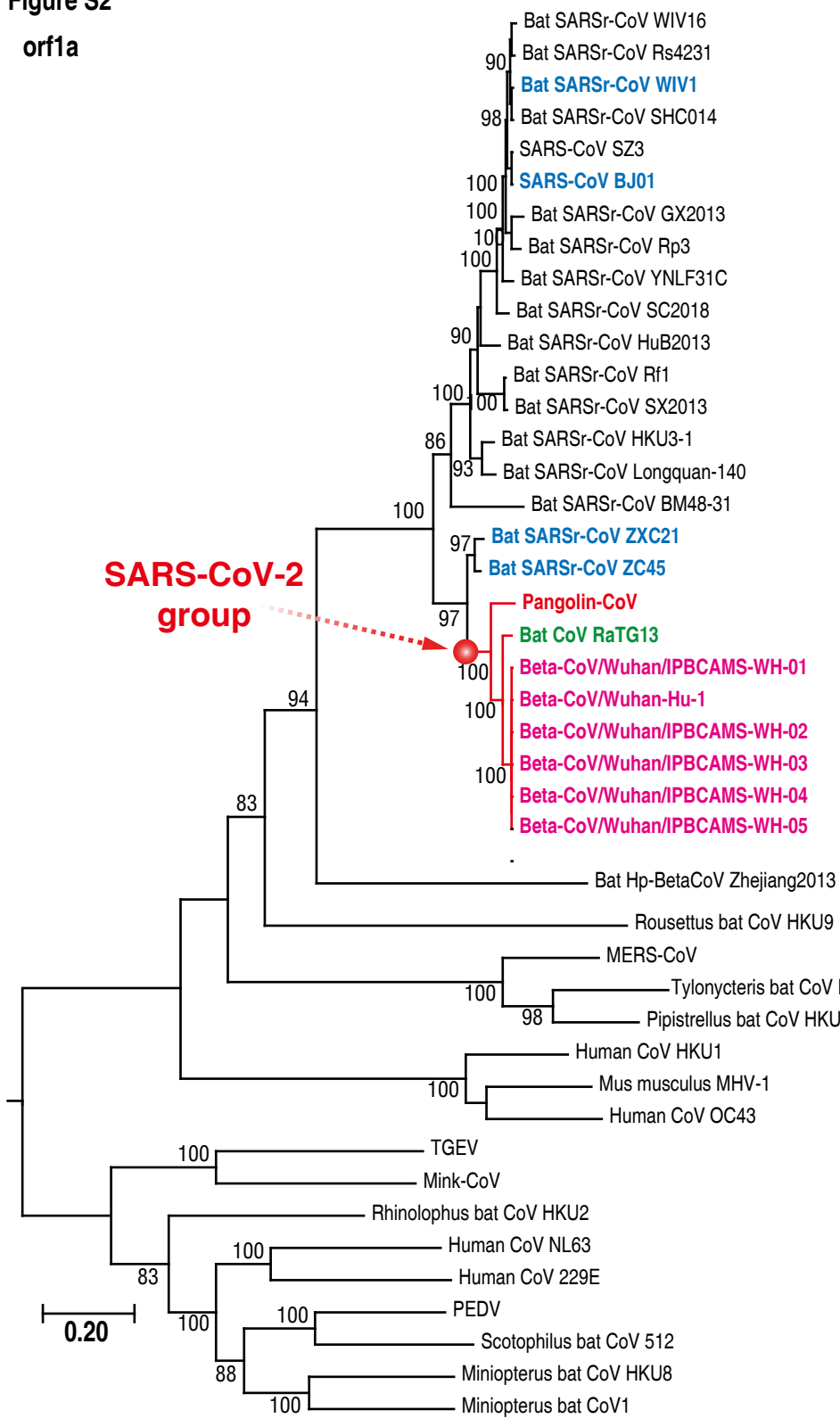
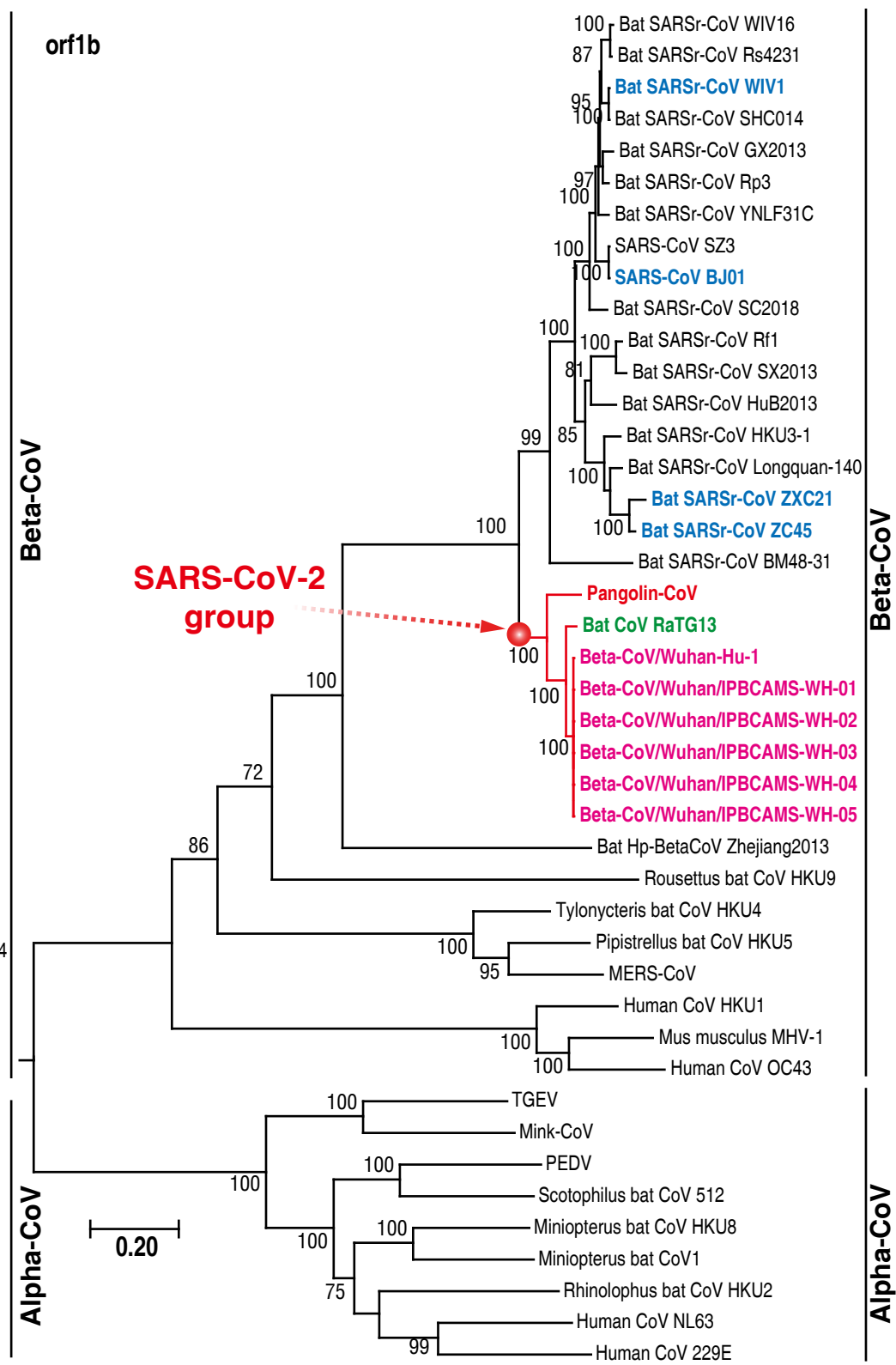
Figure S2**orf1a****orf1b****Figure S2.** Phylogenetic relationship of coronavirus based on S gene (A) and M gene (B) nucleotide sequences.

Table S1. Contigs taxonomically annotated by using BLASTx against 2845 Coronavirus reference genomes.

Contig_ID	Contig Length (nt)	Subject id	aa identity (%)	alignment length	q. start	q. end	s. start	s. end	e-value	bit score
Con_01	1306	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	86.8	423	37	1305	1	422	1.40E-221	767.3
Con_02	383	AVP78041 non-structural polyprotein 1ab [Bat SARS-like coronavirus]	96.1	127	3	383	431	557	7.50E-64	241.5
Con_03	398	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	70.6	102	2	307	867	968	6.50E-34	142.1
Con_04	541	AVP78030 non-structural polyprotein 1ab [Bat SARS-like coronavirus]	95.5	179	4	540	1311	1489	9.90E-94	341.3
Con_05	525	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	93.1	174	2	523	1495	1668	1.90E-94	343.6
Con_06	611	AVP78041 non-structural polyprotein 1ab [Bat SARS-like coronavirus]	92.7	165	60	554	1671	1835	6.30E-89	325.5
Con_07	307	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	96.2	53	2	160	1866	1918	1.90E-25	113.6
Con_08	721	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	95.5	179	167	703	2700	2878	9.10E-95	345.1
Con_09	1078	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	98.6	359	2	1078	2966	3324	1.80E-208	723.4
Con_10	987	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	99.7	329	1	987	3670	3998	8.20E-179	624.8
Con_11	751	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	100	202	144	749	3960	4161	3.00E-80	297
Con_12	2138	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	96.4	413	859	2094	4402	4814	5.10E-242	835.9
Con_13	562	AIA62319 ORF1ab polyprotein [BIRs-BetaCoV/GX2013]	100	186	3	560	5165	5350	5.80E-113	405.2
Con_14	512	ARI44798 orf1ab polyprotein [Bat coronavirus]	100	170	2	511	5307	5476	3.50E-101	365.9
Con_15	296	ATO98191 non-structural polyprotein 1ab [Bat SARS-like coronavirus]	100	46	157	294	5307	5352	6.50E-23	105.1
Con_16	983	ARI44798 orf1ab polyprotein [Bat coronavirus]	100	288	3	866	5431	5718	3.50E-166	582.8
Con_17	646	ATO98167 non-structural polyprotein 1ab [Bat SARS-like coronavirus]	98.1	215	2	646	5708	5922	2.10E-122	436.8
Con_18	640	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	98.6	213	2	640	5985	6197	9.50E-128	454.5
Con_19	832	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	92.2	244	1	723	6151	6394	2.20E-140	496.9
Con_20	1830	QH060603 orf1ab polyprotein [Wuhan seafood market pneumonia virus]	98.6	570	2	1711	6527	7096	0.00E+00	1145.2
Con_21	496	QH060594 surface glycoprotein [Wuhan seafood market pneumonia virus]	95.8	118	143	496	307	424	1.70E-63	240.7
Con_22	440	QH060594 surface glycoprotein [Wuhan seafood market pneumonia virus]	96.6	145	1	435	378	522	9.80E-84	307.8
Con_23	381	QH060594 surface glycoprotein [Wuhan seafood market pneumonia virus]	93.1	131	1	381	658	788	1.20E-61	234.2
Con_24	503	QH060594 surface glycoprotein [Wuhan seafood market pneumonia virus]	100	157	3	473	785	941	7.00E-86	315.1
Con_25	646	ABD75323 spike protein [Bat SARS CoV Rf1/2004]	95.1	183	96	644	946	1128	3.20E-99	359.8
Con_26	381	QH060595 orf3a protein [Wuhan seafood market pneumonia virus]	96.9	127	1	381	114	240	4.40E-72	268.9
Con_27	900	AVP78045 membrane protein [Bat SARS-like coronavirus]	99.5	222	170	835	1	222	2.40E-121	433.7
Con_28	359	QH060599 orf7a protein [Wuhan seafood market pneumonia virus]	100	42	131	256	1	42	3.40E-18	89.7
Con_29	989	AVP78048 hypothetical protein [Bat SARS-like coronavirus]	95	121	461	823	1	121	1.40E-66	251.9
Con_30	488	QH060601 nucleocapsid phosphoprotein [Wuhan seafood market pneumonia virus]	96.9	162	1	486	3	164	3.20E-91	332.8
Con_31	287	AAU04658 nucleocapsid protein [SARS coronavirus civet010]	97.8	46	2	139	258	303	7.70E-21	98.2
Con_32	555	QH060601 nucleocapsid phosphoprotein [Wuhan seafood market pneumonia virus]	97.8	184	2	553	119	302	7.80E-78	288.5
Con_33	461	QH060601 nucleocapsid phosphoprotein [Wuhan seafood market pneumonia virus]	98.2	109	2	328	257	365	1.50E-58	224.2
Con_34	375	QH060602 orf10 protein [Wuhan seafood market pneumonia virus]	97.4	38	90	203	1	38	4.90E-15	79.3

Table S2. Comparing nt and aa sequence identity difference of ten genes among Pangolin-Cov, 2019-nCoV, and BatCov-RaTG13

Genes	2019-nCoV (nt)	RaTG13 (nt)	Pangolin-CoV (nt)
orf1ab	21290	21287	13669 (partial)
orf1a	13128	13215	7342 (partial)
orf1b	8072	8072	6327 (partial)
S	3822	3810	2156 (partial)
3a	828	828	379 (partial)
E	228	228	120 (partial)
M	669	666	669 (Complete)
orf6	186	186	175 (partial)
orf7a	366	366	296 (partial)
orf8	366	366	366 (Complete)
N	1260	1260	1192 (partial)
orf10	117	117	117 (Complete)
Nucleotide	2019-nCoV vs Pangolin-CoV	2019-nCoV vs RaTG13	Pangolin-CoV vs RaTG13
orf1ab	90.3%	96.6%	90.3%
orf1a	89.9%	96.2%	90.3%
orf1b	90.7%	97.0%	90.4%
S	88.3%	91.5%	87.3%
3a	94.5%	98.6%	95.0%
E	98.3%	100.0%	98.3%
M	93.0%	95.8%	93.0%
orf6	95.1%	98.8%	93.8%
orf7a	92.0%	95.1%	90.4%
orf8	91.7%	96.9%	91.5%
N	94.9%	96.7%	95.4%
orf10	99.1%	99.1%	98.3%
Average	93.2%	96.9%	92.8%
Amino Acid	2019-nCoV vs Pangolin-CoV	2019-nCoV vs RaTG13	Pangolin-CoV vs RaTG13
orf1ab	87.1%	95.8%	86.9%
orf1a	96.8%	98.6%	97.0%
orf1b	73.4%	92.2%	72.8%
S	97.5%	96.7%	95.4%
3a	96.8%	100.0%	96.8%
E	97.4%	100.0%	97.4%
M	98.6%	100.0%	98.6%
orf6	96.3%	100.0%	96.3%
orf7a	96.9%	96.9%	93.6%
orf8	94.8%	94.8%	96.6%
N	96.4%	99.0%	96.4%
orf10	97.3%	97.3%	94.6%
Average	94.1%	97.6%	93.5%