1	Cryo-EM structures of HKU2 and SADS-CoV spike
2	glycoproteins and insights into coronavirus evolution
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19 Abstract

A new porcine coronavirus SADS-CoV was recently identified from suckling piglets 20 with severe diarrhea in southern China and its genome sequence is most identical (~95% 21 22 identity) to that of bat α -coronavirus HKU2. It again indicates bats are the natural reservoir of many coronaviruses that have great potential for cross-species transmission 23 to animals and humans by recombination and/or mutation. Here we report the cryo-EM 24 structures of HKU2 and SADS-CoV spike glycoprotein trimers at 2.38 Å and 2.83 Å 25 resolution, respectively. HKU2 and SADS-CoV spikes exhibit very high structural 26 similarity, with subtle differences mainly distributed in the NTD and CTD of the S1 27 subunit responsible for cell attachment and receptor binding. We systematically 28 analyzed and compared the NTD, CTD, SD1 and SD2 domains of the S1 subunit and 29 the S2 subunit of HKU2 spike with those of α -, β -, γ -, and δ -coronavirus spikes. The 30 results show that the NTD and CTD of HKU2/SADS-CoV are probably the most 31 ancestral in the evolution of spike. Although the S2 subunit mediating membrane fusion 32 is highly conserved, the connecting region after fusion peptide in HKU2/SADS-CoV 33 34 S2 subunit also adopts a conformation distinct from other coronaviruses. These results structurally demonstrate a close evolutionary relationship between HKU2 /SADS-CoV 35 and β -coronavirus spikes and provide new insights into the evolution and cross-species 36 transmission of coronaviruses. 37

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39 Introduction

Coronaviruses, categorized into the order Nidovirales family Coronaviridae and 40 subfamily *Coronavirinae*, are a large group of viral pathogens with a wide host range¹. 41 42 Their infections in humans, other mammals and birds can cause respiratory, hepatic, enteric and neurological diseases with varying severity². The severe acute respiratory 43 syndrome coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus 44 (MERS-CoV) have posed severe threats to human health in the 21st century^{1,3}. In the 45 meantime, coronaviruses infecting domestic animals also bring substantial economic 46 losses⁴. For example, the swine acute diarrhea syndrome coronavirus (SADS-CoV) 47 (also known as SeACoV and PEAV) isolated in 2017 caused outbreaks of severe watery 48 diarrhea of suckling piglets with a mortality up to 90% in several commercial pig farms 49 in Guangdong Province of China⁵⁻¹⁰. SADS-CoV is an α -coronavirus and other 50 representative members in the α -genus are porcine epidemic diarrhea virus 51 (PEDV), porcine transmissible gastroenteritis coronavirus (TGEV), porcine respiratory 52 coronavirus (PRCV), human NL63 and 229E coronaviruses (HCoV-NL63 and HCoV-53 54 229E)¹. Representative members in other three genera include mouse hepatitis coronavirus (MHV), bovine coronavirus (BCoV), SARS-CoV, MERS-CoV, HCoV-55 OC43 and HCoV-HKU1 in the β -genus, avian infectious bronchitis virus (IBV) in the 56 γ -genus and porcine deltacoronavirus (PdCoV) in the δ -genus¹. 57

58 Cross-species transmission promoted by genetic recombination and/or mutations underlies the host range expansion of coronaviruses^{3,11-13}. Bats are the natural reservoir 59 of more than 30 different α - and β -coronaviruses that have great potential for 60 interspecies transmission by recombination and/or mutation^{12,14-16}. Data on genetic 61 evolution, receptor binding and pathogenesis have demonstrated that human SARS-62 CoV and MERS-CoV most likely originate from bats¹. Palm civets and dromedary 63 camels are the intermediate hosts of SARS-CoV and MERS-CoV from bats to humans¹, 64 respectively. The newly identified porcine SADS-CoV isolates are also found to share 65 ~95% sequence identity with Rhinolophus bat coronavirus HKU2, and this further 66 stressed the severe results of coronavirus spillover from bats to domestic animals⁵⁻¹⁰. 67

However, the molecular mechanisms underlying the transmission of SADS-CoV from
bats to pigs are still unknown and need to be further explored. Recently it was shown
that SADS-CoV is able to infect cells from a broad range of species including mouse,
chicken, pig, monkey and human, indicating a high potential of the SADS-CoV for
interspecies transmission¹⁷.

The spike glycoprotein of coronaviruses mediates viral entry by binding host 73 receptor with the S1 subunit and fusing viral and cellular membranes with the S2 74 subunit, thereby determining viral host range and tissue tropism^{18,19}. As a class I viral 75 fusion protein, the spike exists on the envelope of virion as a homotrimer and each 76 monomer contains more than 1000 amino acid residues that can be cleaved into S1 and 77 S2 subunits¹⁸. For most coronaviruses, the N-terminal domain (NTD) of the S1 subunit 78 (NTD) recognizes cell surface carbohydrates, while the C-terminal domain (CTD) 79 specifically binds to cellular protein receptors¹⁸⁻²⁰. SARS-CoV and HCoV-NL63 utilize 80 CTD to bind human receptor ACE2²¹⁻²³; MERS-CoV utilizes CTD to bind human 81 receptor DPP4^{24,25}; TGEV, PRCV and 229E utilize CTD to bind receptor APN²⁶⁻²⁸; 82 HCoV-OC43 utilizes NTD to recognize glycans²⁹; and one exception is MHV, which 83 utilizes the NTD to bind mouse receptor CEACAM1a³⁰. Therefore, the S1 subunit, 84 especially its NTD and CTD, is the most variable region of the spike, and is responsible 85 for different tropisms of coronaviruses. In comparison, the S2 subunit containing the 86 fusion peptide (FP) and heptad repeats (HR1 and HR2) for membrane fusion are more 87 conserved in both sequence and structure^{18,19}. For the SADS-CoV, receptor analysis 88 indicated that none of the known coronavirus protein receptors including ACE2, DPP4 89 and APN are essential for the cell entry^{7,17}. There are also no reports regarding to the 90 recognition of glycans by the NTD of SADS-CoV. 91

Structural studies of the spike and its binding with glycans and protein receptors have provided important insights into the origin, evolution and interspecies transmission of coronaviruses. Cryo-EM structures of spike trimer from all four coronavirus genera have been reported: the α -coronavirus spike structures are determined for HCoV-NL63³¹, HCoV-229E²⁷ and PEDV³²; the β -coronavirus spike structures are determined for MHV^{33,34}, HCoV-HKU1³⁵, HCoV-OC43²⁹, SARS- 98 $CoV^{21,36-38}$ and MERS-Co $V^{36,39,40}$; the γ -coronavirus spike structure is determined for 99 IBV⁴¹ and the δ -coronavirus spike structure is determined for PdCoV^{42,43}. The cryo-100 EM structures of bat coronavirus spike trimers have not been reported, and only crystal 101 structures of the CTD from HKU4⁴⁴, HKU5⁴⁵ and HKU9⁴⁶ were determined.

The spikes of SADS-CoV (1130 amino acid residues) and HKU2 (1128 amino 102 acid residues) are the shortest among all known coronavirus spike glycoproteins and 103 their amino acid identities to other known coronavirus spikes are lower than 28%, 104 indicating the spikes of HKU2 and SADS-CoV are unique^{5-10,47}. In this study, we report 105 the cryo-EM structures of the SADS-CoV and HKU2 spike trimers at 2.83 Å and 2.38 106 Å resolution, respectively. The HKU2 spike trimer structure is the first one from bat 107 coronavirus. We analyzed the HKU2 and SADS-CoV trimer structures and also 108 compared the NTD, CTD, SD1 and SD2 domains of the S1 subunit and the S2 subunit 109 of HKU2 with other spikes from α -, β -, γ -, and δ -coronaviruses. Our results strongly 110 support that HKU2 and SADS-CoV preserve primitive structural features in their spikes 111 that have a close evolutionary relationship with β -coronavirus spikes and provide new 112 113 insights into the evolution and cross-species transmission of coronaviruses.

114

115 **Results**

116 **Protein expression and structure determination**

The cDNAs encoding HKU2 spike (YP 001552236) and SADS-CoV spike 117 (AVM41569.1) were synthesized with codons optimized for insect cell expression. 118 HKU2 ectodomains (residues 1-1066) and SADS-CoV ectodomains (residues 1-1068) 119 were separately cloned into pFastBac-Dual vector (Invitrogen) with C-terminal foldon 120 tag and Strep tag. After expression in Hi5 insect cells and purification to homogeneity, 121 the cryo-EM images on these two spike ectodomains were recorded using FEI Titan 122 Krios microscope operating at 300 KV with a Gatan K2 Summit direct electron detector 123 (Supplementary Fig. 1). About 1,400,000 particles for HKU2 spike and 900,000 124 particles for SADS-CoV spike were subjected to 2D classification, and a total of 125 421,490 particles of HKU2 spike and 152,334 particles of SADS-CoV spike were 126

selected and subjected to 3D refinement with C3 symmetry to generate density maps 127 (Supplementary Fig. 2). The overall density maps were solved to 2.38 Å for HKU2 128 spike and 2.83 Å for SADS-CoV spike (gold-standard Fourier shell correlation = 0.143) 129 (Supplementary Fig. 1 and Supplementary Fig. 2). The atomic-resolution density map 130 enabled us to build nearly all residues of HKU2 spike ectodomains (residues 17-995) 131 except for a few breaks (residues 129-141 and 204-204), as well as 48 N-linked glycans 132 (Supplementary Fig. 3a and Supplementary Fig. 4a). The final refined model of SADS-133 134 CoV spike contains residues 19-998 with some short breaks (residues 134-143 and 488-490) and 45 N-linked glycans (Supplementary Fig. 3b and Supplementary Fig. 4b). 135 Data collection and refinement statistics for these two structures are listed in 136 Supplementary Table 1. 137

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139 Overall structures of HKU2 and SADS-CoV spikes

The overall structures of HKU2 and SADS-CoV spikes we determined here resemble 140 the previously reported pre-fusion structures of coronaviruses spikes. Both spike 141 trimers have a mushroom-like shape (\sim 150 Å in height and \sim 115 Å in width) (Fig. 1a), 142 consisting of a cap mainly formed by β -sheets of the S1 subunit, a central stalk mainly 143 formed by α -helices of the S2 subunit, and a root formed by twisted β -sheets and loops 144 of the S2 subunit (Fig. 1a). In each trimer there is a C3 axis along the central stalk (Fig. 145 1a). The amino acid identity between HKU2 and SADS-CoV spikes is 86%, and these 146 two spike structures are quite similar with the root mean square deviation (r.m.s.d.) 147 being 0.53 Å for 962 aligned Ca atoms of the monomer and 0.56 Å for 2886 aligned 148 $C\alpha$ atoms of the trimer. Due to the high structural similarity, we will use the HKU2 149 150 structure to present the features of both spikes in the subsequent description, whereas significant differences between them will be pointed out only when necessary. 151

The S1 subunit of the HKU2 spike comprises two major domains, NTD and CTD, which are followed by two subdomains SD1 and SD2 connecting them to the S2 subunit (Fig. 1b and Fig. 1c). The S1 subunits from three monomers form the cap of the spike, in which the three CTDs in the inner part are at the apex sitting on top of the central stalk and the three NTDs are located outside the CTDs surrounding the central stalk

(Fig. 1a). The NTD, CTD, SD1 and SD2 of the S1 subunit are all mainly composed of 157 β strands regarding to the secondary structure feature (Fig. 1c). In contrast, the upstream 158 helix (UH), fusion peptide (FP), connecting region (CR), heptad repeat 1 (HR1) and 159 central helix (CH) of the S2 subunit are mainly composed of helices, whereas the β -160 hairpin (BH) and subdomain 3 (SD3) at the bottom part of the S2 subunit mainly consist 161 of β stands and loops (Fig. 1c). Moreover, the residues after the SD3, which contain the 162 heptad repeat 2 (HR2), are not resolved in the HKU2 and SADS-CoV spike structures, 163 164 as well as in all other reported coronavirus spike structures in the pre-fusion state.

The SD1 and SD2 of the S1 subunit and the S2 subunit are highly similar in amino 165 acid sequence (85%, 84% and 95% identities) and structure (C α r.m.s.d. less than 0.5 166 Å) between HKU2 and SADS-CoV spikes (Supplementary Fig. 5). The NTD has the 167 lowest sequence identity of 70% among all domains. Structural superimposition also 168 gave a Cα r.m.s.d. of 1.2 Å between these two NTDs, and conformational variations 169 reside in the loops, although the core β sheet structure is structurally conserved 170 (Supplementary Fig. 5a). The CTD sequence identity between HKU2 and SADS-CoV 171 spikes is 82%, but the C α r.m.s.d. between these two CTDs is 1.1 Å, also indicating 172 structural variations in the CTD in comparison with the highly conserved SD1, SD2 173 and S2 subunit. The NTD and CTD of the S1 subunit are commonly utilized by 174 coronaviruses for binding cell-surface carbohydrates or protein receptors for cell 175 attachment²⁰. Therefore, the sequence and/or structural variations indicate that HKU2 176 and SADS-CoV would also bind different host receptors by NTD and/or CTD of the 177 S1 subunit, although their receptors in bat and pig are unknown and the receptor-178 binding sites on spike have not been defined. 179

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181 NTD structure and comparisons

The NTD of HKU2 has three layers of antiparallel β -sheet with the top one consisting of six strands, the middle one consisting of five strands and the bottom one consisting of three strands. Below the bottom sheet is a short α -helix (Fig. 2a). The top and middle β -sheets form a galectin-like β -sandwich fold, which is inserted between two stands of the bottom sheet (Fig. 2a). To supplement, three disulfide bonds are detected in the 187 HKU2 NTD structure: C^{17} - C^{56} connecting the N-terminus of the NTD to its upper loop, 188 C^{124} - C^{149} connecting $\beta 6$ and $\beta 7$ stands in the top sheet and C^{234} - C^{244} connecting the 189 bottom helix to the bottom sheet (Fig. 2a).

Although the NTDs of all coronaviruses adopt a similar overall architecture, the 190 NTD of HKU2 has the highest structural similarity with the NTD1 (named domain 0 in 191 previous reports) of α-coronavirus HCoV-NL63 with an r.m.s.d. of 2.7 Å for 186 192 aligned Ca atoms (Fig. 2b and Supplementary Fig. 7d). The NTD1 of a-coronavirus 193 PEDV is not completely modeled in the spike trimer structure, however the partial 194 model still fits well with the NTD of HKU2 with an r.m.s.d. of 2.3 Å for 73 aligned Ca 195 atoms. Both HCoV-NL63 and PEDV have a second NTD (NTD2, also named domain 196 A in previous reports), and the NTD of HKU2 is structurally less similar to the NTD2 197 with an r.m.s.d. of 4.3 Å against HCoV-NL63 NTD2 and of 4.1 Å against PEDV NTD2 198 (Fig. 2b and Supplementary Fig. 7d). Recent structural determination showed that 199 another α -coronavirus, HCoV-229E, also has one NTD, which is more structurally 200 similar to the NTD2 (Ca r.m.s.d. of 2.0 Å) than the NTD1 (Ca r.m.s.d. of 3.7 Å) of 201 HCoV-NL63 (Fig. 2b). All above comparisons indicate that there are two subtypes of 202 NTD in the α -coronaviruses: the subtype I represented by the NTDs of HKU2 and 203 SADS-CoV and the NTD1s of HCoV-NL63 and PEDV, and the subtype II represented 204 by the NTD of HCoV-229E and the NTD2s of HCoV-NL63 and PEDV (Fig. 2b). 205 206 Although sharing an overall architecture, these two NTD subtypes have a structural difference in the galectin-like β -sandwich fold containing the top and middle sheets 207 stacked together through hydrophobic interactions. These two β sheets are well aligned 208 in the galectin-like domain of subtype I, whereas there is an alignment shift in the 209 galectin-like domain of subtype II (Fig. 2b). The other notable difference is the 210 distribution of signature disulfide bonds. A signature disulfide bond C¹²⁴-C¹⁴⁹ 211 (numbered in HKU2 and connecting $\beta 6$ and $\beta 7$ stands in the top sheet) is conserved in 212 all subtype I NTDs (Fig. 2b), and the subtype II NTDs have two signature disulfide 213 bonds: the first one C^{145} - C^{168} (numbered in HCoV-229E) connecting β 5 and β 6 strands 214 and the second one C^{81} - C^{105} (numbered in HCoV-229E) connecting $\beta 2$ and neighbor 215

loop (Fig. 2b). Besides, C^{234} - C^{244} (numbered in HKU2 and connecting the bottom helix to the bottom sheet) is conserved in both subtype I and subtype II NTDs (Fig. 2b).

The NTDs of β -coronaviruses including BCoV, HCoV-HKU1, HCoV-OC43, 218 MERS-CoV, SARS-CoV and MHV resemble the subtype I, rather than the subtype II 219 NTD in the topology and distribution of the disulfide bonds (Fig. 2c). These β -220 coronavirus NTDs have additional loops in the N-terminus, between $\beta 1$ and $\beta 2$ strands, 221 and between $\beta6$ and $\beta7$ stands (numbered in HKU2 structure), forming an extensive 222 223 ceiling-like structure on top of the galectin-like fold (Fig. 2c). It has been found that the evolvement of this ceiling-like structure has functional outcomes such as immune 224 evasion or receptor binding⁴¹. The NTD of γ -coronavirus IBV also resemble the subtype 225 I NTD in the topology, although its disulfide bond positions are not conserved as in the 226 227 subtype I NTD (Fig. 2d). To be note, the NTD of δ -coronavirus PdCoV resemble the subtype II NTD in the topology and distribution of the disulfide bonds (Fig. 2e). Both 228 IBV and PdCoV NTDs also have additional insertions including loops and short helices 229 in the galectin-like fold compared to the two subtypes of NTD in α -coronaviruses (Fig. 230 231 2d and Fig. 2e).

232

233 **CTD structure and comparisons**

The CTD of HKU2 has a twisted five-stranded antiparallel β sheet as the core with 234 connecting loops between the stands (Fig. 3a). It contains four disulfide bonds: C²⁷⁷-235 C³⁰⁰ and C²⁸⁵-C²⁹⁰ at the N-terminus, C³⁴¹-C³⁹⁷ at the C-terminus and the last one C³³¹-236 C^{369} connecting the $\beta 2$ and $\beta 5$ strands in the core β sheet (Fig. 3a). Interestingly, the 237 CTD core of HKU2 is of high structural similarity with the conserved CTD core of β-238 coronaviruses and the disulfide bonds in the CTD of HKU2 except for C²⁸⁵-C²⁹⁰ are 239 also detected in all β-coronavirus CTDs (Fig. 3b). These CTDs have the core of one 240 twisted β -sheet and here we name them as one-layer CTD subtype (Fig. 3a and Fig. 3b). 241 The β -coronavirus CTDs always have an insertion consisting of loops and/or stands 242 between the β 5 and β 6 strands of the core (Fig. 3b). SARS-CoV, MERS-CoV, HKU4 243 244 and HKU5 have receptor-binding motif (RBM) in this insertion region responsible for binding their respective protein receptors⁴⁵. In the CTD of HKU2, there is only one short loop between the β 5 and β 6 strands of the core twisted β -sheet (Fig. 3a).

Although as members in the α -genus, HKU2 and SADS-CoV CTD structures are 247 significantly different from those of other α -coronaviruses HCoV-NL63, HCoV-229E, 248 PEDV, TGEV and PRCV that contain two layers of β -sheets (Fig. 3c). And we named 249 these CTDs as two-layer CTD subtype. All available two-layer CTD structures can be 250 well aligned with C α r.m.s.d. in the range of 1.0-3.4 Å, except for the CTDs of HKU2 251 and SADS-CoV. These two-layer CTDs contain two highly conserved disulfide bonds: 252 C⁵⁴⁰-C⁵⁸⁶ and C⁵⁶⁹-C⁵⁹⁶ (numbered in PEDV CTD) (Fig. 3c). The C⁵⁶⁹-C⁵⁹⁶ is conserved 253 among all coronaviruses, whereas the C^{540} - C^{586} is conserved in all α -coronaviruses 254 (except for HKU2 and SADS) and δ -coronavirus PdCoV (Fig. 3c). 255

The CTD of δ -coronavirus PdCoV have a core of two β -sheets, belonging to the two-layer CTD subtype (Fig. 3c). As for the γ -coronavirus IBV, the core of its CTD is also similar to the typical two-layer CTD (Fig. 3c). However, several β strands are replaced by loops and the disulfide bonds are in different positions from the two-layer CTD (Fig. 3c). IBV CTD also has an extra region of loops, reminiscent of the extra domain in the CTDs of β -coronaviruses (Fig. 3c).

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263 SD1 and SD2 structures and comparisons

The SD1 and SD2 are two subdomains following the CTD in the S1 subunit, linking 264 the CTD to the S2 subunit. The HKU2 SD1 is a partial β barrel consisting of five β 265 strands and a disulfide bond (C^{409} - C^{458}) connecting its C-terminus to the β 1 strand (Fig. 266 4a). This five-stranded β barrel and the linking disulfide bond are conserved among all 267 four genera of coronavirus (Fig. 4a). The HKU2 SD2 has a structure of two layers of 268 β -sheet with an additional short α -helix over the top sheet (Fig. 4b). The additional α -269 helix and the top sheet is linked by a disulfide bond (C^{482} - C^{509}), and another disulfide 270 bond $(C^{524}-C^{533})$ links the C-terminal loop to the bottom sheet (Fig. 4b). The two-layer 271 272 core structure and the second disulfide bond are conserved among all genera of coronavirus, however, the additional α helix and the first linking disulfide bond is a 273

distinct feature of β -coronaviruses plus α -coronavirus HKU2 and SADS-CoV (Fig. 4b).

275 This additional helix appears to be an insertion between the primitive $\beta 2$ and $\beta 3$ strands

- of the SD2, and is retained during evolution of β -coronaviruses.
- 277

278 Quaternary packing of the NTD and CTD in the spike

It has been observed that coronaviruses have two types of quaternary packing mode of 279 the S1 subunits in the trimer: intra-subunit packing and cross-subunit packing⁴¹. 280 Actually, this is mainly caused by different positioning and interaction between NTD 281 and CTD in the spike monomer. The HKU2 S1 subunit, similar to those in α -282 coronaviruses HCoV-NL63, HCoV-226E and PEDV and δ-coronavirus PdCoV, have 283 an "inward" CTD which contacts with the NTD (Fig. 5a). The three structural "NTD-284 CTD" modules in the cap region of these spikes are composed of NTD and CTD from 285 the same monomer, forming the intra-subunit packing in the spike trimer (Fig. 5a). The 286 S1 subunits of other coronaviruses in the β - and γ -genera including MHV, SARS-CoV, 287 MERS-CoV, HCoV-OC43 and IBV have an "outward" CTD that is far away from the 288 289 NTD (Fig. 5b). Therefore, the three structural "NTD-CTD" modules in the cap region of these spikes have the NTD from one monomer and the CTD from the adjacent 290 monomer, forming the cross-subunit packing in the spike trimer (Fig. 5b). Interestingly, 291 we found that the "outward" CTDs always have an insertion in the core structure, such 292 as β -coronavirus CTDs and γ -coronavirus IBV CTD (Fig. 5b). In contrast, all "inward" 293 CTDs only have the one-layer or two-layer core structure without obvious inserted 294 region. 295

296

297 Conserved S2 subunit and a distinct CR

Sequence analysis suggested that the S1/S2 protease cleavage site at the boundary
between the S1 and S2 subunits is R544-M545 in HKU2 spike and R546-M547 in
SADS-CoV spike^{5,8,47}. Compared to the S1 subunit, the topology and structure of S2
subunit are highly conserved in all coronavirus spikes. The HKU2 S2 subunit contains
a 3-helix upstream helix (UH) (residues 589-639), a fusion peptide helix (FP) (residues
672-684), a connecting region (CR) (residues 689-747), a 4-helix heptad repeat 1 (HR1)

(residues 748-836), a central helix (CH) (residues 837-887), a twisted β-hairpin (BH) 304 (residues 888-929) and a β -sandwich like SD3 (residues 930-995) (Fig. 1b and Fig. 6a). 305 Like in other coronavirus spikes in the prefusion state, the model of HR2 after SD3 was 306 not built in the structure due to poor density. Five disulfide bonds in S2 are detected. 307 Two of them (C^{590} - C^{612} and C^{595} - C^{601}) stabilize the folded helices of UH, C^{696} - C^{706} 308 bends the CR, C⁸⁸⁴-C⁸⁹⁵ links the CH and the BH, and C⁹³⁴-C⁹⁴³ is within the SD3 (Fig. 309 6a). The first four disulfide bonds are conserved in all coronaviruses, and the last one 310 311 in the SD3 has different positions in different spikes. Specifically, it links the β 2 and β 3 stands of SD3 in the spikes of HKU2, SADS-CoV and MERS-CoV (numbered in 312 MERS-CoV), and in other coronavirus spikes it links the β 2 stand to the C-terminal 313 loop of SD3 (numbered in MERS-CoV) (Supplementary Fig. 6). 314

All coronavirus spikes have the S2' protease site upstream from the FP in the S2 315 subunit, which is essential for proteolytic fusion activation of the spike. Receptor 316 binding and cleavage at the S2' site promote large-scale conformational changes of the 317 FP, CR, HR1 and HR2, allowing the insertion of FP into host cell membrane and the 318 319 formation of six-helix bundle. The FP and CR, which are often not well and totally resolved in other coronavirus spike structures, can be clearly modeled in the HKU2 320 spike due to the atomic resolution of the map (Supplementary Fig. 4). The typical CR 321 in the S2 subunit contains three helices and one short strand, with a disulfide bond 322 bending the first and second helix to form a turning (Fig. 6b). In HKU2, the second 323 helix is replaced by a short strand (713-716) and the third helix is replaced by a loop 324 (721-741), therefore there are two short strands and only one helix in HKU2 CR (Fig. 325 6c). The conserved disulfide bond C^{696} - C^{706} makes the first helix of CR in HKU2 spike 326 turn upside down. The S2' cleavage site (between R671 and S672) is then covered by 327 the reversed CR helix and loops, and R671 interacts with E723 in the loop and K697 328 and K698 in helix 1 (Fig. 6c). In other coronaviruses, taking the MHV S2 for example, 329 the helix 1 does not cover the S2' site (between R869 and S870), and R869 only loosely 330 interacts with T929 in helix 3 (Fig. 6b). After the dissociation of the S1 subunit triggered 331 by receptor binding, the exposure of the S2' site for cleavage is a prerequisite for the 332 proteolytic activation of the coronavirus spike to mediate membrane fusion. The buried 333

S2' site indicates that HKU2 spike, compared to other coronavirus spikes, would
require more conformational changes around the S2' site for the exposure.

336

337 **Discussion**

A new porcine coronavirus SADS-CoV (also named as SeACoV and PEAV in other 338 reports) was recently identified from suckling piglets with diarrhea in southern China, 339 and its genome sequence was most identical (~95% identity) to that of Rhinolophus bat 340 α -coronavirus HKU2⁵⁻¹⁰. The SADS-CoV and HKU2 are phylogenetically located in a 341 sub-lineage closely related to the proposed α -coronavirus group-1b lineage at the 342 complete genome level⁵⁻¹⁰. However, phylogenetic analysis based on the spike 343 glycoprotein indicated that they are members of a separate lineage clustered within β-344 coronavirus (Supplementary Fig. 7a), suggesting that HKU2 and SADS-CoV probably 345 resulted from recombination of an α -coronavirus with an unrecognized β -coronavirus 346 S gene⁵⁻¹⁰. These results, together with the lower than 28% amino acid identities to 347 other known coronavirus glycoproteins, strongly indicate that the spike glycoproteins 348 of HKU2 and SADS-CoV are unique⁵⁻¹⁰. In this study, we determined the cryo-EM 349 structures of HKU2 and SADS-CoV spike glycoproteins at atomic resolutions. Pairwise 350 comparisons demonstrated nearly identical overall structures, and differences mainly 351 locate in the loops of NTD and CTD of the S1 subunit between the spikes of HKU2 and 352 SADS-CoV (Supplementary Fig. 5). A series of structural analysis and comparisons 353 were also performed at the domain level between HKU2 spike with other coronavirus 354 spikes with determined structures. Our results show that HKU2 and SADS-CoV spikes 355 356 maintain primitive structural features, especially in the NTD and CTD, and provide more insights into the evolution of coronaviruses. 357

The HKU2 and SADS-CoV have one NTD in the S1 subunit, and their structures are more similar to the NTD1 than the NTD2 of α -coronaviruses HCoV-NL63 and PEDV, whereas the only NTD of HCoV-229E is structurally more similar to the NTD2 than the NTD1 (Fig. 2). Therefore, we suggest that α -coronaviruses have two subtypes of NTD. The evolution relationship between them are not clear yet. It was once

suggested that the presence of two NTDs in HCoV-NL63 is a result of gene 363 duplication³¹. However, the sequence identity between these two NTDs is only 15.7% 364 in HCoV-NL63 and 12.9% in PEDV. Considering that HKU2 (SADS-CoV) and 365 HCoV-229E have one NTD belonging to either subtype I or subtype II, a more plausible 366 evolution way of the NTD in α -coronaviruses is the recombination of two separate 367 primitive domains into the genome, resulting in the presence of two NTDs in the S1 368 subunit α-coronaviruses including HCoV-NL63 and PEDV. To be note, these two NTD 369 370 subtypes may represent primitive structures that could be the evolutionary ancestors of NTDs of other genera coronaviruses. For example, in the current available spike 371 structures, the NTDs of β -coronavirus are similar to the HKU2 NTD representing the 372 subtype I in both architecture and disulfide bond positions. These β-coronavirus NTDs 373 also have additional loop ceiling over the top sheet, functionally facilitating immune 374 evasion or binding protein receptor such as in MHV³⁰. The NTD of γ -coronavirus IBV 375 is also architecturally similar to the subtype I, although the disulfide bond positions are 376 not conserved (Fig. 2d). In contrast, the δ -coronavirus PdCoV NTD is similar to the 377 378 HCoV-229E NTD representing the subtype II in both architecture and disulfide bond positions (Fig. 2e). A previous study of the IBV spike proposed that α -coronavirus 379 NTDs are probably the most ancestral and the NTDs of the four genera form an 380 evolutionary spectrum in the order of α -, δ -, γ -, and β -genus⁴¹. Our proposal here is 381 similar to the previous one in the point that two NTD subtypes in α -coronaviruses may 382 represent primitive structures that could be the evolutionary ancestors of NTDs. 383 However, we argue that the evolution pathways may not be in the order of α -, δ -, γ -, 384 and β -genus. A more plausible pathway is that the β -, γ - and δ -coronavirus NTDs may 385 evolve independently and parallelly from subtype I (β - and γ -coronavirus NTDs) or 386 subtype II (δ -coronavirus NTDs) (Supplementary Fig. 7b). 387

The HKU2 and SADS-CoV CTD structure have a one-layer core consisting of a twisted five-stranded antiparallel β sheet. Interestingly, β -coronavirus CTDs also have the similar one-layer core structure and three strictly conserved disulfide bonds are also present in the core of HKU2 CTD. Currently, all identified receptor-binding motif of β coronavirus CTDs are within an inserted domain between two stands of the core sheet,

and this insertion responsible for receptor binding of β -coronaviruses is replaced by a 393 short loop in HKU2 CTD. This result firstly indicates HKU2 CTD represent a primitive 394 structure in the one-layer CTD family, while the inserted domain in β -coronaviruses 395 results from a recombinant event during evolution (Supplementary Fig. 7c). The second 396 indication is that HKU2 and SADS-CoV may not utilize the CTD to bind protein 397 receptors that have not been identified yet, and their different receptor usage may be 398 determined by the NTD that harbors almost 50% of residue difference between them. 399 400 To be note, the CTDs from other α -coronaviruses, γ -coronavirus IBV and δ -coronavirus PdCoV all belong to the two-layer subtype consisting of two layers β-sheets, although 401 with structural variations in different viruses. These results further confirmed the 402 previous phylogenetic analysis suggesting that HKU2 and SADS-CoV probably 403 resulted from a recombination of an α -coronavirus genomic backbone with an 404 unrecognized β -coronavirus spike gene^{5,8,47}. 405

In contrast with the NTD and CTD having unique structural feathers, the SD1 and 406 SD2 domains of the S1 subunit and the S2 subunits of HKU2 and SADS-CoV are 407 408 structurally conserved as those of other coronaviruses. In the evolutionary aspect, it is not surprising because this region either connects the CTD to the S2 subunit (SD1 and 409 SD2) or mediates the membrane fusion, whereas the NTD and CTD are key factors 410 determining tissue tropism and host range of coronaviruses²⁰. Even highly conserved 411 in overall structure, the S2 subunit in HKU2 and SADS-CoV still have a secondary 412 structure arrangement in the connecting region (CR) after the fusion peptide (FP), 413 resulting in a more buried S2' cleavage site (Fig. 6c). It indicates that although the 414 membrane fusion mechanism is highly conserved, the dynamic fusion procedures of 415 416 HKU2 and SADS-CoV may still have their unique features that need to be addressed 417 in the future.

Since the outbreak of SARS epidemic in 2002-2003, new coronaviruses including MERS-CoV, HCoV-HKU1, HCoV-NL63 in human and PdCoV and SADS-CoV in livestock has been identified^{1,16}. Extensive studies also revealed a diversity of coronaviruses in bats that could be the source for emergence of zoonotic epidemics such as SARS in the past and new one in the future^{1,12,16}. The spikes of bat coronavirus 423 HKU2 and porcine coronavirus SADS-CoV (1128 and 1130 residues) are the shortest among all current known coronavirus spike glycoproteins and their NTDs and CTDs 424 may also represent evolutionary ancestors^{5-9,47}. In comparing the structures of HKU2 425 and SADS-CoV spikes with other coronavirus spikes, we observed that additional 426 NTDs could be recombined, ceiling loops could be inserted into NTD core, and extra 427 domain containing receptor-binding motif could be inserted into the CTD core structure. 428 These phenomena indicate the subdomains are gradually recruited into the S1 subunit 429 430 during evolution, and the recruitments are required for cross-species transmission, adapting to different host range, and responding to the updating of host immune system, 431 which provides a vivid example for the co-evolution of virus and host. 432

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441 Author contributions

J.Y. carried out protein expression, purification, electron microscopy sample
preparation, data collection, image processing and model building with the help of S.Q.
X.W. conceived, designed and directed the study. X.W., J.Y. and R.G analyzed the
structure, made the figures and wrote the manuscript.

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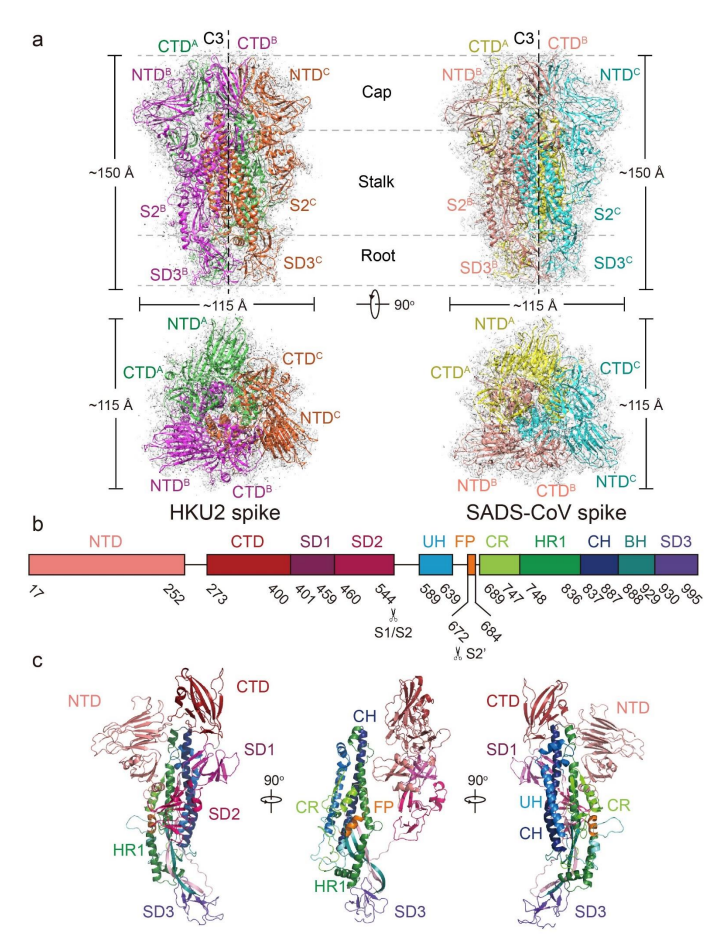
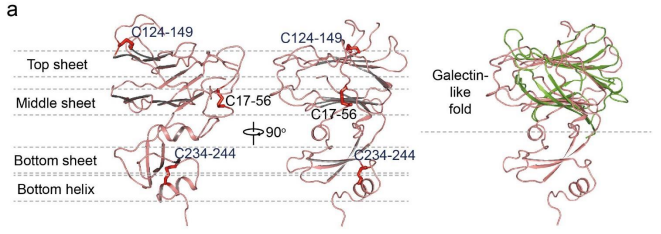


Fig. 1 Overall structures of HKU2 and SADS-CoV spike glycoproteins. (a) Overall structures of HKU2 and SADS-CoV spike glycoproteins shown in side view (upper panel) and top view (lower panel). Three monomers of HKU2 spike are colored magenta, green, and orange, respectively; three monomers of SADS-CoV spike are colored pink, yellow, and cyan, respectively. The cryo-EM maps are shown as semitransparent surface and contoured at 2.6 RMS and 3 RMS for HKU2 and

SADS-CoV spikes, respectively. The trigonal axes are shown as black dashed lines. Visible segments of each monomer are labeled accordingly. The cap, stalk and root parts are partitioned by gray dashed lines. (b) Segmentation of HKU2 monomer. The segments of HKU2 are shown as boxes with the width related to the length of amino acid sequence. The start and end amino acids of each segment are labeled. The position of S1/S2, and S2' cleavage sites are indicated. NTD, N-terminal domain; CTD, C-terminal domain; SD1, subdomain 1; SD2, subdomain 2; UH, upstream helix; FP, fusion peptide; CR, connecting region; HR1, heptad repeat 1; CH, central helix; BH, β -hairpin; SD3, subdomain 3. (c) Overall structure of HKU2 monomer. Side views of HKU2 monomer shown in three directions. The segments are colored the same as in b.



HKU2 NTD

HKU2 NTD Galectin3

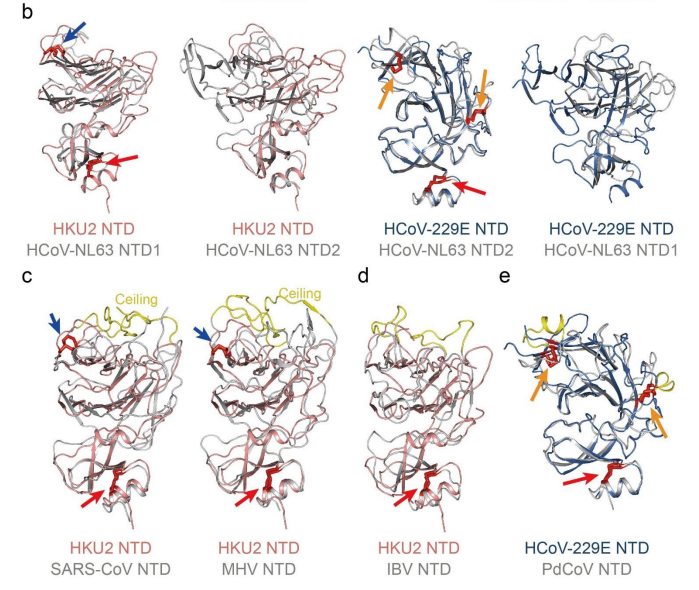


Fig. 2 Structure of HKU2 NTD and comparisons. (a) Structure of HKU2 NTD. Side views of HKU2 NTD are shown in two orthogonal directions. Disulfide bonds are shown as red sticks. Conserved disulfide bonds are labeled blue; other disulfide bonds are labeled black. Top sheet, middle sheet, bottom sheet, and bottom helix are partitioned by gray dashed lines. Comparison of HKU2 NTD and galectin3 are shown in the right panel. HKU2 NTD is colored salmon; galectin3 is colored green. PDB code: galectin3, 1A3K. (b) Two subtypes of α -coronavirus NTD. Structural alignments of HKU2 NTD with HCoV-NL63 NTD1, and with HCoV-NL63 NTD2 are shown in the left two panels; structural alignments of HCoV-229E NTD with HCoV-NL63 NTD1, and with HCoV-NL63 NTD2 are shown in the right two panels. HKU2 NTD is colored salmon;

HCoV-229E NTD is colored marine; HCoV-NL63 NTD1 and NTD2 are colored gray. Disulfide bonds are shown as red sticks. Disulfide bonds conserved in both types of NTDs are indicated by red arrows; disulfide bonds conserved in subtype I NTD are indicated by blue arrows; disulfide bonds conserved in subtype II NTD are indicated by orange arrows. PDB codes: HCoV-229E, 6U7H; HCoV-NL63, 5SZS. (c) β -coronavirus NTDs resemble subtype I. Structural alignments of HKU2 NTD with SARS-CoV NTD, and with MHV NTD are shown. Disulfide bonds are shown and labeled the same as in b. Ceilings in β -coronavirus NTDs are shown in yellow. PDB codes: SARS, 5XLR; MHV, 3JCL. (d) γ -coronavirus IBV NTD resembles subtype I. Structural alignment of HKU2 NTD with IBV NTD is shown. Disulfide bonds are shown and labeled the same as in b. The additional loops in IBV NTD is shown in yellow. PDB code: IBV, 6CV0. (e) δ -coronavirus PdCoV NTD resembles subtype II. Structural alignment of HCoV-229E NTD with PdCoV NTD is shown. Disulfide bonds are shown and labeled the same as in b. The additional loops in IBV NTD is shown in yellow. PDB code: IBV, 6CV0. (e) δ -coronavirus PdCoV NTD resembles subtype II. Structural alignment of HCoV-229E NTD with PdCoV NTD is shown. Disulfide bonds are shown and labeled the same as in b. The additional helices in PdCoV NTD are shown in yellow. PDB codes: HCoV-229E, 6U7H; PdCoV, 6B7N.

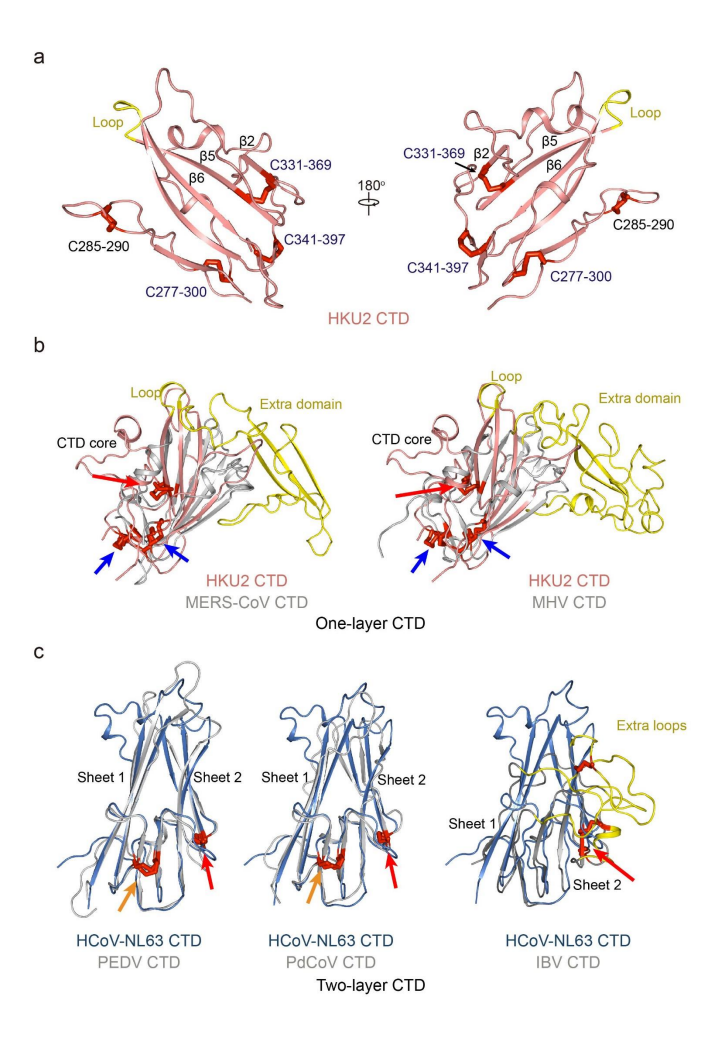


Fig. 3 Structure of HKU2 CTD and comparisons. (a) HKU2 CTD shown in two opposite directions. Strands mentioned in the main text are labeled. The loop replaced by extra domain in β -coronavirus CTDs is shown yellow. Disulfide bonds are shown as red sticks. Conserved disulfide bonds are labeled blue; other disulfide bonds are labeled black. **(b)** β -coronavirus CTDs belong to one-layer CTD. Structural alignments of HKU2 CTD with MERS-CoV CTD, and with MHV CTD are shown. Extra domains in β -coronavirus CTDs are colored yellow. Disulfide bonds are shown as red sticks. Disulfide bonds conserved in both one-layer CTD and two-layer CTD are indicated by red arrows. Disulfide bonds only conserved in one-layer CTD are indicated by blue arrows. PDB codes: MHV, 3JCL; MERS-CoV, 6Q05. **(c)** α -coronavirus (except HKU2 and SADS), γ -coronavirus, and δ -coronavirus CTDs belong to two-layer CTD. Structural alignments of β sheets are labeled. Extra loops in IBV CTD are colored yellow. Disulfide bonds conserved in both one-layer CTD and two-layer CTD are shown. Two layers of β sheets are labeled. Extra loops in IBV CTD are colored yellow. Disulfide bonds conserved in both one-layer CTD and two-layer CTD are shown. Two layers CTD are indicated by orange arrows. PDB codes: HCoV-NL63, 5SZS; PEDV, 6U7K; IBV, 6CV0; PdCoV, 6B7N.

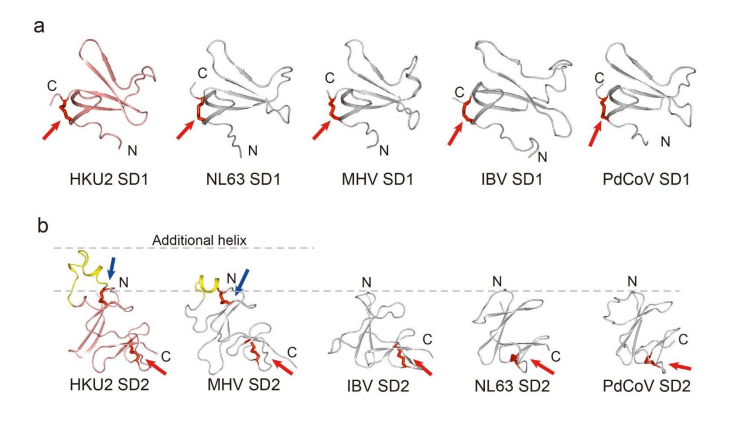


Fig. 4 Structures of SD1 and SD2 and comparisons. (a) Comparison of SD1 from four genera of coronaviruses. HKU2 SD1 is colored salmon; SD1 from other coronaviruses are colored gray. Disulfide bonds are shown as red stick. Red arrows indicate the disulfide bonds conserved in all genera of coronaviruses. PDB codes: HCoV-NL63, 5SZS; MHV, 3JCL; IBV, 6CV0; PdCoV, 6B7N. (b) Comparison of SD2 from four genera of coronaviruses. HKU2 SD2 is colored salmon; SD2 from other coronaviruses are colored gray. Disulfide bonds are shown as red stick. Red arrows indicate the disulfide bonds conserved in all genera of coronaviruses. HKU2 SD2 is colored salmon; SD2 from other coronaviruses are colored gray. Disulfide bonds are shown as red stick. Red arrows indicate the disulfide bonds conserved in all genera of coronaviruses. Blue arrows indicate the disulfide bonds only found in HKU2 (and SADS-CoV) and β CoVs. The additional helices of SD2 from HKU2 (and SADS-CoV) and β CoVs are colored yellow and partitioned by gray dashed lines. PDB codes: HCoV-NL63, 5SZS; MHV, 3JCL; IBV, 6CV0; PdCoV, 6B7N.

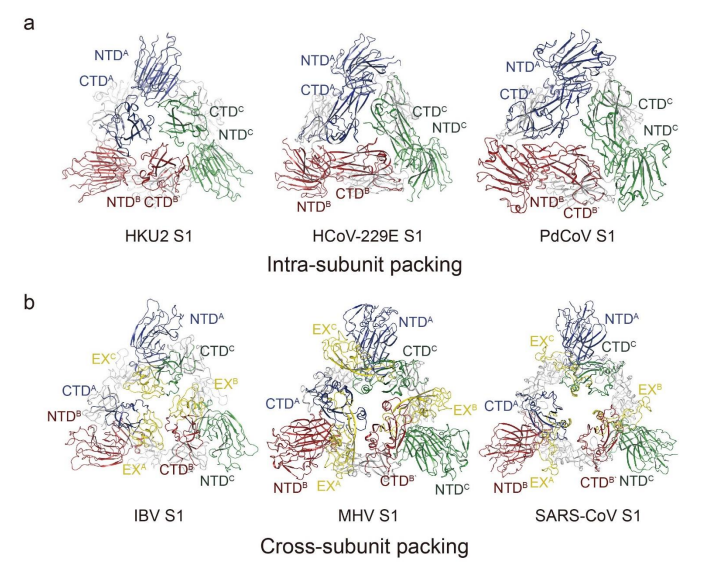


Fig. 5 Quaternary packing of NTD and CTD. (a) α -coronavirus S1 and δ -coronavirus S1 use intra-subunit packing pattern. NTD and CTD from the first monomer are colored blue, the second are colored red, and the third are colored green. PDB codes: HCoV-229E, 6U7H; PdCoV, 6B7N. (b) β -coronavirus S1 and γ -coronavirus S1 use cross-subunit packing pattern. NTD and CTD from the first monomer are colored blue, the second are colored red, and the third are colored green. The extra loop of IBV and the extra domains of β CoVs are colored yellow and labeled as EX. PDB codes: MHV, 3JCL; IBV, 6CV0; SARS, 5XLR.

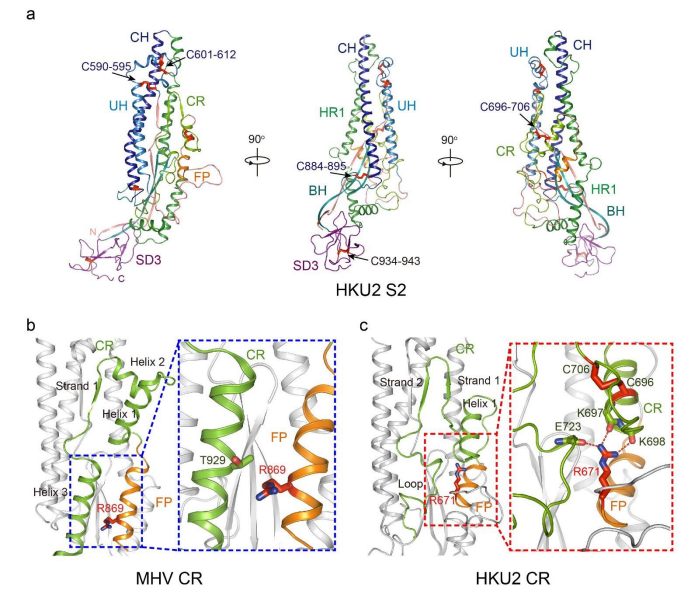


Fig. 6 Structure of HKU2 S2. (a) Side views of HKU2 S2 shown in three directions. Seven segments of S2 are shown as different colors. UH, upstream helix; FP, fusion peptide; CR, connecting region; HR1, heptad repeat 1; CH, central helix; BH, β -hairpin; SD3, subdomain 3. Disulfide bonds are shown as red stick. Disulfide bonds conserved in all coronaviruses are labeled blue; the other disulfide bond is labeled black. (b) Conserved CR represented by MHV CR. CR and FP are colored the same as in a. Helices and strands in CR are labeled. R869 (S2' cleavage site) is shown as stick and colored red. The blue dashed box shows R869 does not interact tightly with MHV CR. PDB code: MHV, 3JCL. (c) Unique feature of HKU2 CR. CR and FP are colored the same as in a. Helices and strands in CR are labeled. R671 (S2' cleavage site) is shown as stick and colored red. The blue colored red. The red dashed box shows detailed interactions between R671 and CR.