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Structure of SARS-CoV-2 spike protein
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The COVID-19 (coronavirus disease 2019) pandemic, caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has led to loss of human life in millions and devastating socio-economic consequences worldwide. The disease has created urgent needs for intervention strategies to control the crisis and meeting these needs requires a deep understanding of the structure-function relationships of viral proteins and relevant host factors. The trimeric spike (S) protein of the virus decorates the viral surface and is an important target for development of diagnostics, therapeutics and vaccines. Rapid progress in the structural biology of SARS-CoV-2 S protein has been made since the early stage of the pandemic, advancing our knowledge on the viral entry process considerably. In this review, we summarize our latest understanding of the structure of the SARS-CoV-2 S protein and discuss the implications for vaccines and therapeutics.

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Introduction
Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the causative agent of the COVID-19 (coronavirus disease 2019) pandemic \cite{1}, and its infection has led to millions of lives lost and devastating socio-economic consequences throughout the globe. There are urgent needs for innovative vaccine and therapeutic strategies to control this unprecedented crisis, as well as potential future needs if it becomes seasonal with continuous emergence of new variants. A deep understanding of the structure-function relationships of viral proteins and relevant host factors will be required in order to meet these needs. Coronavirus (CoVs) are enveloped positive-stranded RNA viruses that enter a host cell by fusion of its envelope lipid bilayer with the target cell membrane. This first critical step of viral infection is catalyzed by its trimeric spike (S) protein, which decorates the virion surface as a major antigen and induces neutralizing antibody responses. The protein is therefore an important target for development of diagnostics, therapeutics and vaccines. Remarkable progress in the structural biology of SARS-CoV-2 S protein has been made since the initial outbreak of the virus \cite{2}, substantially advancing our molecular understanding of the viral entry process. Here we summarize our current knowledge on the structure of the SARS-CoV-2 S protein and discuss the implications for vaccines and therapeutics.

Overall structure of SARS-CoV-2 S protein
The SARS-CoV-2 spike glycoprotein is a type I membrane protein (Figure 1a), which forms a trimer, anchored to the viral membrane by its transmembrane segment, while decorating the virion surface with it large ectodomain (Figure 1b). It binds to the receptor angiotensin-converting enzyme 2 (ACE2) on a host cell and undergo large structural rearrangements to promote membrane fusion \cite{1,3}. The protein is heavily glycosylated with each protomer containing 22 N-linked glycosylation sites \cite{4,5}. The full-length S protein of the Wuhan-Hu-1 strain from the initial outbreak has 1273 amino acid residues, including a N-terminus signal peptide, a receptor-binding fragment S1 and a fusion fragment S2. S1 can be further divided into N-terminal domain (NTD), receptor-binding domain (RBD) and C-terminal domains (CTD1 and CTD2), while S2 includes fusion peptide (FP), fusion-peptide proximal region (FPPR), heptad repeat 1 (HR1), central helix (CH), connector domain (CD), heptad repeat 2 (HR2), transmembrane segment (TM) and the cytoplasmic tail (CT), depicted in Figure 1a.

Structures of S protein fragments derived from the Wuhan-Hu-1 strain, including the S ectodomain stabilized in its prefusion conformation \cite{6,7}, RBD-ACE2 complexes \cite{8,9,10,11}, and segments of S2 in the postfusion state \cite{12}, were determined within the first several months of the pandemic. Soon after, structures of detergent-solubilized, full-length S proteins in both prefusion and postfusion conformations \cite{13,14}, as well as those of the intact S trimer on the virion surface, studied by cryo-electron tomography \cite{15,16,17,18}, were also reported (Figure 1b and c). Overall, the SARS-CoV-2 S
Distinct conformational states of the SARS-CoV-2 spike protein. (a) Schematic representation of the SARS-CoV-2 spike protein organization. Segments of S1 and S2 include: NTD, N-terminal domain; RBD, receptor-binding domain; CTD1, C-terminal domain 1; CTD2, C-terminal domain 2; S1/S2, S1/S2 cleavage site; S2', S2' cleavage site; FP, fusion peptide; FPPR, fusion peptide proximal region; HR1, heptad repeat 1; CH, central helix region; CD, connector domain; HR2, heptad repeat 2; TM, transmembrane anchor; CT, cytoplasmic tail; and tree-like symbols for glycans. (b) Left: viral SARS-CoV-2 S trimer in the prefusion conformation (EMD-30430; Ref. [15]), fitted with the structures of purified proteins (PDB ID: 7KRR and 6XR8; Refs. [13**,50**]). Right: cryo-EM structure of the full-length S trimer in the RBD-down conformation (PDB ID: 6XR8). (c) Left: viral SARS-CoV-2 S2 trimer in the postfusion conformation (EMD-30428; Ref. [15]), fitted with the structure of the purified protein (PDB ID: 6XRA; Ref. [13**]). Right: cryo-EM structure of the full-length S2 trimer in the postfusion conformation (PDB ID: 6XRA). (d) Additional structures of coronavirus S proteins, including the full-length SARS-CoV-2 S2 trimer carrying G614 in the one RBD-up conformation (PDB ID: 7KRR), the stabilized soluble SARS-CoV-2 S trimer in the RBD-down conformation (PDB ID: 6VXX; Ref. [7]), the stabilized soluble SARS-CoV-2 S trimer in the one RBD-up conformation (PDB ID: 6VSB; Ref. [6**]). (e) MHV (mouse hepatitis virus) S2 in the postfusion state (PDB ID: 6B3O; Ref. [25]), and SARS-CoV S2 in the postfusion state (PDB ID: 6M3W; Ref. [24]).

structure shows many similarities to those of other coronavirus spike proteins [19–23]. In the prefusion structure, the S1 fragment, adopting a ‘V’ shaped architecture with the NTD at one arm and the RBD, CTD1 and CTD2 at the other (also see Figure 2a), which wrap around the central helical bundle formed by the prefusion S2 fragment, projecting the N-terminal end of HR1 toward the viral membrane. Three RBDs form the apex of the S trimer, sampling two distinct conformations — ‘up’ representing a receptor-accessible state and ‘down’ representing a receptor-inaccessible state (Figure 1b). The three NTDs are located at the periphery of the trimer, each making contacts with the RBD from the adjacent protomer. The CTD1 and CTD2 pack underneath the RBD against S2 and between the two neighboring NTDs, indicating they could modulate these domains and play
Structures of NTD and its antibody complexes.
(a) Cryo-EM structure of S1 fragment from the full-length SARS-CoV-2 S trimer (PDB ID: 6XR8), with the NTD highlighted in blue and the rest of S1 in gray. (b) Close-up view of the NTD in the SARS-CoV-2 S protein. (c) The NTD (in blue) from its complex with 4A8 is superposed with the domain from the full-length S trimer in gray, showing shifts of the five surface loops (N1–N5). (e) and (f) Close-up view of the binding interface for the NTD-4A8 and NTD-DH1052 complexes with contacting residues in the NTD highlighted in sticks. (d) Superposition of the structures of the NTD in complex with antibody 4A8 Fab (PDB ID: 7C2L; Ref. [28*]) and DH1052 Fab (PDB ID: 7LAB; Ref. [32]), as indicated. Heavy and light chains of 4A8 are colored in red and pink, respectively, and those of DH1052 are in green and cyan, respectively.
important roles in the structural rearrangements required for membrane fusion.

In the postfusion conformation, S1 dissociates as a monomer, while S2 adopts a rigid, baseball-bat-like shape (~220 Å long), and the HR1 flips over to form a continuous long helix together with the CH, which is further surrounded by short helices and β-sheets at the distal end of the membrane (Figure 1c and e). The connector domain (CD), together with a segment (residues 718–729) in the S1/S2–S2’ fragment, form a three-stranded β sheet, and residues 1127–1135 join the connector β sheet to expand it into four strands. Another segment (residues 737–769) in the S1/S2–S2’ fragment makes up three helical regions locked by two disulfide bonds that pack against the groove of the CH part of the coiled coil to form a short, six-helix bundle structure (6Hβ-1). The N-terminal region of HR2 adopts a one-turn helical conformation and also packs against the groove of the HR1 coiled coil; the C-terminal region of HR2 forms a longer helix that makes up the second six-helix bundle structure with the rest of the HR1 coiled coil (6Hβ-2) [13**,24,25].

**N-terminal domain**

At the periphery of the spike (Figure 1b) [13**], the NTD projects away from the threefold axis, and can be divided into the top, core and bottom regions (Figure 2b). The core structure has a gallerin-like antiparallel β-sandwich fold, formed by one six-stranded β-sheet and the other with seven strands. The top region has two antiparallel β strands connected by a short loop, while the bottom region is primarily made up of two short β sheets and a helix. The overall structure of the NTD is surrounded by eight N-linked glycans and similar to that of the S proteins from Middle East respiratory syndrome coronavirus (MERS-CoV) [26] and bovine coronavirus [27].

The exact function of the NTD in SARS-CoV-2 S remains unknown, although NTDs of other coronaviruses have been shown to recognize sugars upon initial attachment or specific protein receptors, or play a role in the prefusion-to-postfusion transition [27]. Nonetheless, NTD-targeted neutralizing antibodies (nAbs), with a potency at the nM level, have been isolated from SARS-CoV-2 infected patients [28**], suggesting a functionally critical role of this domain. High-resolution structures of the S protein in complex with NTD-directed neutralizing antibodies (4A8, FC05, CM25, 4-18, S2M28, and DH1205) have been determined [28**,29*,30,31*,32], showing that these antibodies primarily bind to two glycan-free surfaces of the domain, designated NTD-1 and NTD-2 regions, respectively (Figure 2b; Ref. [33]). Most antibodies target the NTD-1 region, which is thus named the NTD-1-antigenic supersite. It is located at the edge of the NTD top-core region, including five surface loops: N1 (residues 14–26), N2 (residues 67–79), N3 (residues 141–156), N4 (residues 177–186), N5 (residues 246–260) (Figure 2c), and a β-hairpin structure near N3, surrounded by four N-linked glycans (Asn17, Asn74, Asn122 and Asn149). These loops reconfigure upon binding to various antibodies (Figure 2c).

In the S-4A8 complex structure (Figure 2d) [28**], the third complementarity determining region (CDR3) of the 4A8 heavy chain inserts to a cleft formed by the N3 β-hairpin/loop and N5 loop, while the CDR1 and CDR2 interact with the tips of the two loops. Moreover, the glycan at Asn149 is very close to the interface and may also contribute to antibody binding (Figure 2c). Other antibodies, such as S2M28, 4-18, DH1050, CM25, FC05, 12C9 [33], also use their CDR1-3 to contact the N3 and N5 loops, but some interact with the nearby N1 loop or the glycan at Asn17 as well. Despite the differences in approaching angles among these antibodies, their interface with the NTD-1 is highly conserved. Up till now, NTD-2 is recognized by non-neutralizing antibodies, such as by DH1052 and 81D6 [33]. The CDR loops of both heavy and light chains in DH1052 interact with the surface formed by residues spanning 27–32, 59–62 and 211–218 in the NTD (Figure 2f), with possible involvement of the glycan at Asn603 of the CTD-2. Not surprisingly, the newly emerged SARS-CoV-2 variants of concerns, including Alpha (lineage B.1.1.7), Beta (B.1.351), Gamma (B.1.1.28) and Delta (B.1.617.2), all have mutations and/or deletions within the NTD-1-supersite, rendering resistance to neutralization by NTD-directed antibodies [34,35].

**Receptor binding domain**

The RBD contains two subdomains — a five-stranded antiparallel β sheet connected by short helices and loops, and an extended loop, named receptor binding motif (RBM) [8**,9*,36]. In the host cell, ACE2 is an important component of renin-angiotensin system (RAS) and catalyzed the hydrolysis of angiotensin II to angiotensin 1–7 [9*]. The full-length human ACE2 is also a chaperone of the amino acid transporter B’AT1 and forms a homodimer mediated by its neck domain in the presence of B’AT1 (Figure 3a) [10]. Cryo-EM structures of the soluble uncleaved S protein in complex with monomeric ACE2 show that the S trimer can bind one, two or three ACE2s in the RB-D loop conformations (Figure 3b) [37,38]. The crystal structure of the SARS-CoV-2 RBD-ACE2 complex reveals a similar structure to the SARS-CoV RBD-ACE2 complex [8**,9*,10]. A gently concave outer surface of the extended RBM interacts with the N-terminal helix of the claw-like peptidase domain (PD) of ACE2 (Figure 3c) [8**,9*,10]. Hydrogen bonds and salt bridges between a series of polar residues, such as K417, E484, N487 and N501 of the RBM and D30, K31, H34, Y41 and K353 of the ACE2, dominate the RBD-ACE2 interaction (Figure 3c) [8**,9*,10]. Additional hydrophobic interactions between F486 of the RBM and L79, M82 and Y83 of the ACE2 also contribute to the receptor
Structures of ACE2, ACE2-S complexes and RBD-antibody complexes.
(a) Cryo-EM structure of the full-length ACE2 in complex with BQT1 (PDB ID: 6M17; Ref. [10], with the peptidase domain (PD) of ACE2 in pink, its neck domain in magenta, the transmembrane helix in yellow, and BQT1 in gray. (b) The side view (left) and the top view (right) of cryo-EM structure of the soluble S trimer complexed with three ACE2s (PDB ID: 7KJ4; Ref. [37]). (c) Left, the crystal structure of SARS-CoV-2 RBD in complex with ACE2 (PDB ID: 6M0J; Ref. [8**]). With ACE2 in pink and the RBD in cyan. Middle and right, close-up views of the binding interface with contacting residues from the N-terminal helix of the ACE2 and the RBM of the RBD shown in sticks. (d) Left, cryo-EM structure of the RBD in complex with antibody REGN10933 (PDB ID: 6XDG; Ref. [43**]). Middle, cryo-EM structure of the RBD in complex with antibody REGN10987 (PDB ID: 6XDG; Ref. [43**]). Right, crystal structure of the RBD in complex with antibody CR3022 (PDB ID: 6YLA; Ref. [45]). The RBD is shown in cyan and heavy and light chains of the antibodies in various colors. The RBM is highlighted in dark blue.
engagement (Figure 3c) [8**,9*,10]. Mutations of the key residues, such as N501Y, K417N and E484K, have been identified in the fast-spreading variants of concern, leading to enhanced affinity for ACE2 and immune evasion [39,40].

The RBD is a dominant target of nAbs elicited by either natural infection or vaccination, confirming its pivotal role during infection [41,42]. The RBD-directed nAbs can recognize multiple distinct epitopes, showing great potencies at the pM-nM level in vitro neutralization assays (Figure 3d) [42]. The nAbs that target the ACE2-binding-site, such as REGN10933, C144 and S2H14, directly compete for ACE2 association [41,42,43**,44]. Those recognizing the non-ACE2-binding-site, such as REGN10987 and C135, probably prevent ACE2 binding either by clashing with ACE2 or by blocking the transition of the RBD from the ‘down’ to the ‘up’ conformation [42,43**,44]. Other nAbs against the so-called ‘cryptic supersite’, such as CR3022 and S304, can destabilize the S trimer and induce S1 dissociation [41,42,45]. Although the great potency of this class of antibodies makes them promising therapeutic agents, emergence of resistant variants could limit their clinical applications for treating the COVID-19.

A recombinant human ACE2, named APN01, is under evaluation as a treatment for COVID-19 in a phase 2 clinical trial, based on the favorable results from a previous phase 1 trial [46], and evidence that the protein blocks SARS-CoV-2 infection effectively in vitro [47]. Other ACE2-based fusion inhibitors have been developed with optimized binding and potency comparable to those of the nAbs [37,48,49]. The ACE2 constructs with multivalency, such as the dimeric protein sACE2Δv2.4-IgG1 carrying the mutation T27Y/L79T/N330Y and the trimeric protein ACE2-foldon T27W, can inhibit the viral infection with a potency 1000-fold and 1700-fold greater than that of the monomeric soluble ACE2 with the wildtype sequence [37,48]. Substitution of T27 with an aromatic residue appears to further stabilize the binding interface through non-polar interactions with residues Y489, F456 and Y473 of the RBD (Figure 3c). In addition, a series of miniproteins, created using computer-generated scaffolds to mimic the N-terminal helix of ACE2, can bind the RBD and inhibit viral infection at a concentration below the nM level [49]. These ACE2-derived inhibitors may show even greater potency to those SARS-CoV-2 variants that have gained increased receptor binding than the Wuhan-Hu-1 virus. Nonetheless, pharmacokinetics, in vivo efficacy and safety profile of these new designs still require further validation.

C-terminal domains

The C-terminal domains (CTDs) are formed primarily by β-structures of segments from S1, as well as the N-terminal segment of S2 adjacent to the furin cleavage site (Figure 4). CTD1 contains two antiparallel β-sheets,
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Figure 5

Structures and proposed conformational changes of SARS-CoV S2.
(a) Close-up view of S2 in the prefusion (left) and postfusion (right) conformations from PDB ID: 6XR8 and 6XRA, with the fusion peptide (FP) highlighted in purple, the FPPR in red, central helix (CH) in gold, connector domain (CD) in green, HR1 in orange and HR2 in green. (b) Proposed structural transition of the HR1 from the prefusion to postfusion conformation. (c) Proposed conformational change of the HR2. (d) Six-helix bundle structures in the postfusion S2 with HR1 in orange and HR2 in green.

with two strands and four strands, respectively. CTD2 also has two β-sheets: a four-stranded one and another four-stranded one that includes a strand from the S2 subunit [6**,7**,13**,50**]. In the RBD-down conformation of the S trimer, a structural element in the CTD2, named the ‘630 loop’, becomes well-ordered in the G614 variant while disordered in the Wuhan-Hu-1 strain [13**,50**]. When structured, the 630 loop inserts into a gap between the NTD and CTD1 of the same protomer, stabilizing the CTD2 structure. It is also located in the vicinity of the S1/S2 boundary as well as the FPPR of a neighboring protomer [50**]. The FPPR and the 630 loop help retain the RBDS in the down conformation but move out of their positions when the adjacent RBD flips up. Thus, the CTDs, together with the FPPR and the 630 loop are key components of the S fusion machinery that modulate the fusogenic structural rearrangements of S protein.

S2 structure
In the prefusion conformation [13**], three S2 subunits tightly pack around a central three-stranded coiled-coil of ~140 Å long, formed by CH (Figure 1b). Portion of the HR1 together with another segment of S2 (residues 758–784) adopt α-helical conformation and assemble into a nine helix-bundle with the central coiled-coil, forming the most rigid part of the entire S trimer. The CD region links CH and the C-terminal HR2 through a linker region (Figure 5a). The FP forms a short helix and tucks in a pocket formed by two neighboring S protomers. The structured FPPR clashes with the CTD1 if the RBD moves up and thus appears to help clamp the prefusion S trimer in the closed, RBD-down conformation. It has also been suggested to function as a pH-dependent switch domain that modulates the RBD position [38]. The remaining HR2, TM and CT segments are disordered in the most S trimer structures, but show low-resolution density in the cryo-ET reconstructions that can be tilted away from the threefold axis of the trimer with an angle from 17° to 60° [16**].

In the postfusion conformation [13**,24], the HR1 and CH form a continuous α-helix and three copies of them assemble into a long central three-stranded coiled-coil of ~180 Å (Figure 5a). Two proline substitutions at the
boundary between the HR1 and CH to prevent formation of the postfusion helix have been introduced to stabilize the prefusion conformation and such a design has been used for structural studies and the first-generation vaccines [6,7,51]. Part of the HR2 folds into α-helix and packs against the groove between two HR1-CH helices to form a six-helix bundle structure, reminiscent of the postfusion organization of other viral fusion proteins [52,53]. The CD remains unchanged from the prefusion conformation, as a three-stranded β-sheet covering the C-terminal end of HR1-CH helices. Comparison of the prefusion and postfusion conformations of S suggests that HR1 undergoes large rearrangements to form a coiled-coil, translocating its N-terminal end by a large distance to project the FP towards the target cell membrane (Figure 5b). In addition, the HR2 and the TM at its C-terminal end must fold back to pack along the groove of the HR1-CH coiled-coil to form the postfusion six-helical bundle (Figure 5c). These reordering events effectively bring the viral and target cell membranes close together, ultimately leading to membrane fusion (Figure 5d). Interestingly, five N-linked glycans decorate the postfusion S2 surface along the long axis with a regular spacing and may protect the S2 from the host immune responses.

Implications for vaccines and therapeutics
The SARS-CoV-2 S protein is the key component of almost all the first-generation COVID-19 vaccines [51]. Based on structural studies, concerns have been raised that the inactivated-virus vaccines or those used the wildtype sequence of the Wuhan-Hu-1 strain may have too many postfusion spikes and induce mainly non-neutralizing antibodies [13,18]. Indeed, these vaccines have induced lower levels of neutralizing antibody responses than other S constructs containing stabilization modifications to prevent conformational changes [54]. Additional studies have identified the G614 S trimer as a possible superior immunogen candidate [50,55], as it is naturally constrained in a prefusion state presenting both the RBD-down and RBD-up conformations with great stability. Moreover, the global spread of SARS-CoV-2 and the consequently vast number of replication events make emergence of new variants inevitable, and substantially increases the genetic diversity of the virus, which will bring much greater challenges for vaccine development than it was at the beginning of the pandemic. Indeed, genetic diversity is also the major hurdle for development or optimization of vaccines against several other human pathogens, such HIV-1, hepatitis C virus and influenza virus [56–58]. If SARS-CoV-2 becomes seasonal, structure-based innovative strategies will likely be needed for developing next-generation vaccines designed to elicit broadly neutralizing antibody responses. Likewise, high-resolution structural information has been instrumental for creating peptide-based and ACE2-based fusion inhibitors [12,37,49,59], it will undoubtedly continue to serve as a foundation for rational design of antiviral therapeutics to fight against the pandemic.

Conclusion
Tremendous progress in the structural biology of SARS-CoV-2 spike protein has been made since the initial outbreak of the virus. The structural knowledge not only fills the major gap in our understanding of the viral entry process, but also provides a solid foundation for development and optimization of vaccines and therapeutics against the current and future pandemics of coronaviruses.

Conflict of interest statement
Nothing declared.

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