Research Article

Cryo-EM structure of glycoprotein C from Crimean-Congo hemorrhagic fever virus

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ABSTRACT

Crimean-Congo hemorrhagic fever virus (CCHFV) is a causative agent of serious hemorrhagic diseases in humans with high mortality rates. CCHFV glycoprotein Gc plays critical roles in mediating virus-host membrane fusion and has been studied extensively as an immunogen. However, the molecular mechanisms involved in membrane fusion and Gc-specific antibody-antigen interactions remain unresolved largely because structural information of this glycoprotein is missing. We designed a trimeric protein including most of the ectodomain region of Gc from the prototype CCHFV strain, IbAr10200, which enabled the cryo-electron microscopy structure to be solved at a resolution of 2.8 Å. The structure confirms that CCHFV Gc is a class II fusion protein. Unexpectedly, structural comparisons with other solved Gc trimers in the postfusion conformation revealed that CCHFV Gc adopted hybrid architectural features of the fusion loops from hantaviruses and domain III from phenuiviruses, suggesting a complex evolutionary pathway among these bunyaviruses. Antigenic sites on CCHFV Gc that protective neutralizing antibodies target were mapped onto the CCHFV Gc structure, providing valuable information that improved our understanding of potential neutralization mechanisms of various antibodies.

1. Introduction

Crimean-Congo hemorrhagic fever virus (CCHFV, genus Orthornairovirus, family Nairoviridae, order Bunyavirales) is a prevalent tick-borne virus (Abudurexiti et al., 2019; Kuhn et al., 2020), which causes severe hemorrhagic fever in humans with a case fatality rate ranging from 3% to 80% (Pavel et al., 2020). CCHFV has been classified as a biosafety level 4 (BSL-4) pathogen (Weidmann et al., 2016) and has been declared by the World Health Organization (WHO) as a R&D Blueprint priority pathogen (Mehand et al., 2018). Despite being a deadly pathogen, there is currently no licensed vaccine against CCHF disease (Al-Abri et al., 2017; Tipihand Burt, 2020; Tahir Ul Qamar et al., 2021).

CCHFV contains a tripartite genome composed of small (S), medium (M) and large (L) negative-stranded RNA segments (Bente et al., 2013; Zivcec et al., 2016). The M segment encodes the glycoprotein precursor that is processed by cellular proteases into several proteins, including the N-terminal mucin-like domain (MLD), two non-structural proteins (GP38 and NSm) and two structural glycoproteins (Gn and Gc) (Fig. 1A) (Bergeron et al., 2007; Guardado-Calvo and Rey, 2017). Glycoproteins Gn and Gc form a lattice of putative tetrameric projections on the viral membrane surface according to recent cryo-electron tomography studies of a nairovirus (Hazaravirus) (Punch et al., 2018). The CCHFV Gc glycoprotein is predicted to be a class II membrane fusion protein (Garry and Garry, 2004) that mediates fusion of the viral envelope with the cellular membrane, and this fusion event is triggered by low pH in multivesicular bodies (Shtanko et al., 2014).

Within the Bunyavirales order, most fusion proteins adopt a class II fusion fold, with the remarkable exception of arenaviruses whose GP2 is a class I fusion protein (Hulswit et al., 2021). Class II fusion proteins of bunyaviruses in their postfusion conformations are homotrimers, which have been structurally studied mainly by X-ray crystallography and are limited to the families Phenuiviridae [Rift Valley fever virus (RVFV) and severe fever with thrombocytopenia syndrome virus (SFTSV)] (Haldorsen et al., 2016; Guardado-Calvo et al., 2017) and Hantaviridae [Puumala virus (PUUV), Hantaan virus (HTNV), Andes virus (ANDV) and Maporal virus (MAPV)] (Guardado-Calvo et al., 2016; Willensky et al., 2016).
Fig. 1. Expression of CCHFV sGc-trimers. A Domain diagram of the CCHFV M RNA segment encoding MLD, GP38, Gn, NSm and Gc and transmembrane (TM) regions, which are shown in dark gray. The first residue of each mature protein is provided above. The lower schematic outlines the ectodomain organization of Gc based on our cryo-EM structure: domain I (DI) in red, domain II (DII) in yellow with the fusion loop (FL) in orange, domain III (DIII) in blue and the stem region in magenta. The first residue of each domain is provided above. B SEC elution profiles of CCHFV Gc monomer and trimer are shown as dashed and solid lines, respectively. There are two oligomeric states for sGc-trimers, as indicated by peaks 1 and 2. C SDS-PAGE analysis of degradation of purified sGc-trimers (marked by a red arrow) at 4 °C after 60 days. D and E Negative staining analysis of purified sGc-trimers in peaks 1 and 2 from Fig. 1B. Red arrows indicate rosette-shaped particles formed by aggregation of sGc-trimers (D) and cone-shaped sGc-trimers (E). Scale bar: 50 nm. MLD, mucin-like domain; SEC, size-exclusion chromatography; CCHFV, Crimean-Congo hemorrhagic fever virus; sGc-trimer, trimer of the soluble Gc ectodomain.
2. Materials and methods

Different bunyaviral families. In addition, we analyzed the distribution of evolutionary studies revealed some interesting features shared by structure by cryo-electron microscopy (cryo-EM). Structure-based of a C-terminal GCN4 trimerization motif and determined its atomic

...insertion of a GCN4 trimerization motif (MKQIEDKIEEILSKHYIE-...into the pMT/BiP/V5-His vector (Invitrogen). sGc-trimers were designed IbAr10200 strain (GenBank accession no. AAM48106) was subcloned...genic sites (Fels et al., 2021).

In this study, we constructed a CCHFV Gc-trimer by taking advantage of a C-terminal GCN4 trimerization motif and determined its atomic structure by cryo-electron microscopy (cryo-EM). Structure-based evolutionary studies revealed some interesting features shared by different bunyaviral families. In addition, we analyzed the distribution of critical epitope residues on the Gc structure and proposed potential neutralization mechanisms.

2. Materials and methods

2.1. Protein expression and purification

The DNA sequence encoding the soluble ectodomain of Gc (sGc, corresponding to the M segment residues 1049–1569) of CCHFV IbAr10200 strain (GenBank accession no. AAM48106) was subcloned into the pMT/BiP/VS-His vector (Invitrogen). sGc-trimers were designed by insertion of a GCN4 trimerization motif (MKQIEDKIEEILSKHYIE-NEIARIKKGILGE) between sGc and the C-terminal V5-His tag. A flexible linker (GSGG) was introduced between sGc and the trimerization motif.

A mixture of expression vector and selection vector (19:1) were co-transfected into 1 mL of S2 cells (2 × 10^6 cells/mL) with 8 μL Cellycent® II (10362100; Thermo Fisher Scientific Inc., Carlsbad, CA, USA). The transfection mixture was removed and replaced with 2 mL of ESF921 medium (96-001-01; Expression Systems, Davis, CA, USA) with 5% fetal bovine serum (FBS; 10099141; Gibco, Australia) after 6 h. Then, 300 μL of hygromycin-B (V900372; Sigma-Aldrich, USA) was added to S2 medium (96-001-01; Expression Systems, Davis, CA, USA) equilibrated with 25 mmol/L Tris-HCl, 150 mmol/L NaCl, pH 8.0.

2.2. Cryo-EM sample preparation, data collection and processing

The purified protein was first negatively stained and examined under transmission electron microscope (TEM) to evaluate sample quality. Four microliters of a specimen at a concentration of ~ 0.01 mg/mL was applied onto glow-discharged copper grids supported by a continuous layer of carbon film for 30 s at room temperature. The grids were then submerged into 2% (w/v) uranyl acetate solution for 60 s. The air-dried grids were loaded onto a FEI Tecnai G2 20 TWIN microscope (FEI, United States) (operating at 200 kV) equipped with an Olympus SIS Cantega 2K × 2K CCD (Olympus, Japan).

For cryo-EM specimen preparation, 3.5 μL of the protein sample at a concentration of ~ 0.3 mg/mL was applied onto glow-discharged holey carbon copper grids (Quantifoil R 1.2/1.3, 300-mesh). After excess sample was removed by blotting with filter paper for 5.5 s, the grid was plunge-frozen in liquid ethane using a Leica EM GP2 (Leica Microsystems, Germany) at 25 °C and 85% humidity.

Cryo-EM single particle data collection was performed using a CRYO ARM 300 electron microscope (JEOL, Japan) operated at 300 kV with an in-column energy filter (slit width 30 eV). A total of 3330 images were recorded at a nominal magnification of 80,000 ×, corresponding to 0.61 Å per pixel by a K3 direct electron detector (Gatan, Japan). Each image stack was dose-fractionated to 40 frames with an accumulated dose of ~ 48 e/Å^2 and a total exposure time of 2 s.

After manual selection, 2688 image stacks were subject to beam-induced motion correction using the MotionCor2 implementation in RELION (version 3.1) (Zivanov et al., 2019). Dose-weighted micrographs were then imported into cryoSPARC (version 3.2) for subsequent data processing procedures (Punjani et al., 2017). The contrast-transfer function parameters were fitted by the patch-based CTF estimation procedure. A total of 2514 micrographs were filtered with a CTF fit resolution better than 4.5 Å and a detected focus range of 0.5–2.5 μm.

Totally, 323,944 particles were auto-picked in 250 micrographs by the blob picker tool within cryoSPARC and were subjected to reference-free 2D classification. Five representative classes were selected as references for template-based particle picking for the whole data set. After 2D classification of the resultant 2,060,865 particles, 956,121 particles were selected.
selected from 13 good classes for ab initio generation of four volumes, which were used as initial models for heterogeneous refinement. When C3 symmetry was applied, 187,292 particles selected from the best 3D class yielded a map at an average resolution of 2.8 Å [Fourier shell correlation (FSC) = 0.143] using non-uniform refinement and CTF refinement.

2.3. Atomic model building and refinement

A starting atomic model was built using the phenix.map_to_model tool (Afonine et al., 2018). The only fragment with the correct register was N1377–T1425, including the longest α helix (α2). The crystal structure of RVFV Gc in the postfusion conformation (PDB ID: 6EGT) was used as a guide to trace the backbone of the CCHFV sGc-trimer. The coordinates were modified and the sequence was assigned using Coot (Emsley et al., 2010). The model was improved through multiple cycles of real-space refinement using phenix.real_space_refinement with geometric and secondary structure restraints, and subsequent manual adjustments in Coot. The structure was validated using the phenix.validation_cryoem tool. The model refinement statistics generated in Phenix are summarized in Table 1.

UCSF ChimeraX (Goddard et al., 2018) or PyMOL (The PyMOL Molecular Graphics System, version 1.8, Schrödinger, LLC) was used to visualize the 3D volume map and the atomic model, and local resolution variations were estimated using cryoSPARC. Sequences were aligned using ClustalW, and the secondary structure was analyzed by DSSP (Joosten et al., 2011) and illustrated with ESPript 3.0 (Robert and Gouet, 2014).

3. Results

3.1. Expression of CCHFV Gc

sGc, (~ 60 kDa), which was constructed by truncating the C-terminal transmembrane domain and cytoplasmic domain (residues 1595–1684 in the polyprotein) (Fig. 1A), was expressed using a Drosophila expression
Fig. 3. The postfusion structure of the CCHFV Gc ectodomain. A The cryo-EM map of CCHFV sGc-trimers. The upper map is a side view of sGc-trimers with the region corresponding to one Gc protomer shown in green and the density corresponding to the other two subunits shown in light gray. The bottom is the top view of the map along the C3 axis. Contour threshold used in ChimeraX was set to 0.44. B Ribbon diagram of one Gc ectodomain oriented roughly as shown in panel (A) and color-coded according to Fig. 1A. The β strands “A0–J0” of domain I are labeled by letters with a subscript. The β strands “a–m” of domain II are labeled with lowercase letters, whereas the three helices are labeled α1–3. The β strands “A–H” of domain III are labeled with capital letters. The C-terminal “stem” (magenta) extends along the interface between domains II from neighboring protomers towards the fusion loop (orange). Magnified view (upper left) of the cryo-EM density of the stem region (residues Gln 1546 to Ile1559) is shown as a transparent surface with the underlying atomic model in stick representation. Magnified view (bottom) highlights the connecting region between domains I and III with residues shown in stick representation under the cryo-EM density. Residues are labeled and colored to indicate the domain boundary, and the carbon, oxygen and nitrogen atoms in magnified amino acid residues are colored with gray, red and blue respectively. CCHFV, Crimean-Congo hemorrhagic fever virus; sGc-trimer, trimer of the soluble Gc ectodomain; DI, domain I; DIII, domain III.
system and was purified by nickel affinity chromatography. SEC analysis revealed that purified sGc was monomeric in solution (Fig. 1B). Efforts to obtain sGc crystals were hampered by the poor stability of sGc. A C-terminal trimerization motif (GCN4) with a flexible glycine-serine linker (GSGG) (Hernandez Alvarez et al., 2008; Chan et al., 2012) was appended to sGc to obtain a stable Gc candidate for structural analysis. This sGc product was purified as stable trimers (hereafter referred to as sGc-trimers) with no significant degradation over 60 days at 4 °C (Fig. 1C). SEC analysis of sGc-trimers revealed that there were two oligomeric states in solution (Fig. 1B). One (peak 1) was the aggregation of trimers, which showed a rosette pattern under TEM (Fig. 1D). Negatively stained sGc-trimers (peak 2) were homogeneous cone-shaped particles (Fig. 1E), which were used for subsequent cryo-EM studies.

3.2. Single-particle reconstruction of sGc-trimers

Cryo-EM images (Fig. 2A) of sGc-trimers were collected using a K3 camera mounted on a JEOL CRYO ARM 300 electron microscope. Using a template-based picking strategy, about two million particles for reference-free 2D classification in cryoSPARC were selected (Punjani et al., 2017). Dominant 2D classes showed essentially C3 symmetry (Fig. 2B, classes a and b), indicating that most of the particles oriented with their three-fold axis coinciding with the direction of the electron beams. Using heterogeneous refinement implemented in cryoSPARC, we were able to select ~190,000 particles with an even orientation distribution for final reconstruction of the structure at 2.8 Å resolution when C3 symmetry was applied (Fig. 2C and D).

3.3. CCHFV Gc is a class II membrane fusion protein

The cryo-EM structure of sGc-trimers at atomic resolution enabled tracing of the polypeptide chain from Lys 1078 to Ile1559 (Fig. 3A and B), except for some loop regions: Glu1094–Ile1100, Gly1123–Ser1130, Pro 1195–Gly 1202, Gly1286–Phe1289, Gly1324–His1343 and Phe1501–Thr1503. Our final model contained 482 amino acids per monomer. Interestingly, the C-terminal GCN4 trimerization motif was also missing in the density map, suggesting that the trimerization motif facilitated Gc monomer packing but was not required in forming the final stable Gc trimer structure. One or two N-linked glycosylation sites have been identified in the Gc crystal structures of RVFV, SFTSV, HTNV and PUUV (Guardado-Calvo et al., 2016, 2017; Halliderson et al., 2016; Willensky et al., 2016), and two N-linked glycosylation sites (Asn1194 and Asn1345) were predicted in our model by NetNGlyc (Sun et al., 2007); however, no clear glycans density was observed near the two asparagine residues in our cryo-EM map. The close proximity of Asn1194 and Asn1345 to two disordered loops (Pro1195–Gly1120 and Gly1324–His1343) might contribute to the structural flexibility of the glycans. Statistics on cryo-EM data collection, 3D reconstruction and model refinement are summarized in Table 1.

Like other class II fusion proteins, the ectodomain of CCHFV Gc can be divided into three domains, I, II and III (Fig. 3B). Structure-based sequence alignment showed that the secondary elements of CCHFV Gc largely coincide with those of RVFV and HTNV (Fig. 4A and B), even though the total sequence identity was only 15.7% and 15.3%, respectively. sGc-trimers have a central trimeric core formed by parallel interaction of adjacent protomers involving roughly the N-terminal three-quarters of sGc (domains I and II). The remaining C-terminal portion of sGc packs laterally against the core trimer, which involves domain III and part of the stem region. CCHFV Gc trimerizes through extensive interactions between the adjacent protomers and the total surface area buried in trimer interfaces is ~16,000 Å² (compared with the ~50,000 Å² of total solvent accessible surface area of the trimer).

CCHFV Gc has a long stem region connecting domain III to the transmembrane domain, which spans 49 residues. Electron density for most of this region is missing because of flexibility; however, the first 14 residues of the stem (1546–1559) are visible in the cryo-EM map. The last residue modeled in our structure is Ile1559, which is located ~20 Å from the distal tip of domain II where the fusion loop is situated (Fig. 3B). The remaining 35 residues could readily extend towards the fusion loop to form a “hairpin” conformation with the Gc transmembrane domain and fusion loop at the same end, strongly suggesting that our structure represents CCHFV Gc in its postfusion conformation.

3.4. Distribution of disulfide bonds in Gc

There were 26 cysteine residues in the soluble Gc ectodomain and 24 of these residues are present in the cryo-EM structural model (except Cys1198 and Cys1566). There are 11 disulfide bonds in our model with one disulfide bond in domain I, six in domain II, and the remaining four in domain III. All 11 disulfide bonds are fully conserved among different CCHFV strains (Supplementary Fig. S1).

Domain I of CCHFV Gc is composed of ten β-strands labeled as0–as9 that are arranged in two anti-parallel β-sheets to form a central β-sandwich (Fig. 3B), which is structurally similar to RVFV and HTNV (Fig. 4A and B). The Cys0–Cys9 β-sheets face the trimeric core, whereas the D0E0F0G0 sheet is located on the surface of the Gc trimer. CCHFV Gc has one disulfide bond in domain I between strands F0 and G0 (Cys1236–Cys1246), which is also conserved in domain I of HTNV Gc (Guardado-Calvo et al., 2016).

3.5. Putative fusion loops are structurally similar to those of hantaviruses

Elongated domain II consists of two insertions in domain I (one is between strands E0 and F0 and the other is between strands I0 and J0). Domain II is composed of thirteen β-strands (labeled a through m) and three short helices (α1, α2 and α3). Like other class II fusion proteins, domain II of CCHFV Gc can be divided into two subdomains. One subdomain is proximal to domain I, folding into an open β-barrel composed of six antiparallel β-strands (“αefglm”). In comparison to RVFV and HTNV Gc structures, CCHFV Gc has a longer hairpin structure formed by “lm” β-strands that extend to an adjacent protomer (Fig. 3B). The other subdomain resides at the distal tip of the Gc ectodomain, forming a β-sandwich between the “bcd” β-sheet and the “kj” β-hairpin that projects the “bc” (Thr1164–Cys1169), “cd” (His1187–Gly1204) and “jk” (Cys1360–Pro1366) loops at its outmost end. In our postfusion Gc model, several residues of the “cd” loop are missing. Fairwise sequence alignment shows that the fusion loop of RVFV (Cys818–Pro830) corresponds to a region of CCHFV Gc spanning part of the “cd” loop and complete “d” β-strand (Fig. 4C), suggesting the fusion loop in the two bunyaviral families are not conserved.

Five disulfide bonds in domain II are distributed near the fusion loop region and are essential for structural integrity. The top and middle moieties of the CCHFV Gc “bc” loop are covalently connected with the “d” β-strand by two disulfide bonds (Cys1169–Cys1205 and...
Cys1208 is also similar to that of CCHFV (Fig. 5). It has been reported that instead stabilize the AB and GH loops (Fig. 4C).

β has eight of a single –tip region of domain II (Cys1169 to Cys1360) from RVFV or HTNV Gc with seven β-strands of the adjacent protomer domain II. In contrast to domain III from CCHFV Gc and HTNV Gc with seven β-strands, domain III from CCHFV Gc has eight β-strands with the extra E β-strand, resulting in a more closed β-barrel structure. The four disulfide bonds in domain III function to stabilize the AB and GH loops (Fig. 4C).

Compared with other Gc structures in their postfusion states from the Phenuiviridae and Hantaviridae families, all three domains of CCHFV Gc can be roughly aligned to those of RVFV Gc, whereas superimposable regions of HTNV Gc are limited to domains I and II (Fig. 4A and B). The polypeptide sequence linking domains I and III is well defined in the cryo-EM map (Fig. 3B), validating that this domain arrangement is distinct from “the swapping of domain III” observed in hantaviruses (Guardado-Calvo et al., 2016). There are 23 conserved residues in domain III between CCHFV and RVFV, especially for the A β-strand (5 of the 10 residues being conserved) (Fig. 4C), whereas only 11 residues are conserved between CCHFV and HTNV (Supplementary Fig. S2). In addition, there is one conserved disulfide bond between RVFV and CCHFV in domain III (Cys1455–Cys1458), whereas no conserved disulfide bond is present between HTNV and CCHFV in this region. All these structural observations strongly indicate that nairoviruses and phenuiviruses share a possible common evolutionary origin for domain III.

3.6. Organization of domain III resembles its phenuivirus counterpart

Domain III has an immunoglobulin superfamily C2 subtype fold formed with an eight-stranded (ABCDEFGH) β-barrel-like module (Halaby et al., 1999) (Fig. 3A). The AB, CD and FG loops are close to the β-sheets of the adjacent protomer domain II. In contrast to domain III from RVFV or HTNV Gc with seven β-strands, domain III from CCHFV Gc has eight β-strands with the extra E β-strand, resulting in a more closed β-barrel structure. The four disulfide bonds in domain III function to stabilize the AB and GH loops (Fig. 4C).

fig. 5. Structure and sequence alignment of fusion loops and conserved disulfide bonds between CCHFV Gc and HTNV Gc. A Superposition of Gc fusion loops of CCHFV (yellow) onto HTNV (gray; PDB ID: 5LJZ) reveals highly similar conformations. Residues from Pro 1195 to Gly 1202 in the cd loop of CCHFV Gc are missing in the density map and are depicted as a dashed line. Three conserved disulfide bonds of CCHFV are shown as sticks and in magenta (Cys1169–Cys1205), blue (Cys1193–Cys1360) and green (Cys1208–Cys1368), respectively. Three conserved disulfide bonds of HTNV are shown as gray sticks. B Sequence alignment of bc, cd and kj loops between CCHFV and three hantaviruses. The first and last residues of each loop are numbered on the top of the sequences. Residues highlighted in red are strictly conserved. The cysteine residues of conserved disulfide bonds are labeled by arrows that match the coloring used in (A). Conserved or similar amino acids are highlighted in light blue border, whereas fully conserved amino acids are highlighted in red. It is worthy to note that a cysteine pair (Cys1165-Cys1198, labeled as Cys1198) is highly conserved. Although the cysteine pair forms a disulfide bond in hantaviruses (Guardado-Calvo et al., 2016), further experimental evidence is needed to confirm whether this is also the case in CCHFV Gc, since the density for Cys1198 is missing in the cryo-EM model. CCHFV, Crimean-Congo hemorrhagic fever virus; HTNV, Hantaan virus; PUUV, Puumala virus; ADNV, Andes virus.

Cys1173–Cys1207, and the “bc” loop is also stabilized by an internal disulfide bond (Cys1175–Cys1180). There is a disulfide bond bridging “cd” and “jk” loops (Cys1193–Cys1360), and a disulfide bond stapling together β-strands “k” and “d” (Cys1208–Cys1368) (Fig. 5).

HTNV Gc and CCHFV Gc share three conserved disulfide bonds in the tip region of domain II (Cys1169–Cys1205, Cys1193–Cys1360 and Cys1208–Cys1368), and the overall structure of the fusion loops of HTNV is also similar to that of CCHFV (Fig. 5). It has been reported that instead of a single “fusion loop”, all three loops at the tip of domain II from HTNV Gc insert into the target membrane (Guardado-Calvo et al., 2016). Structural conservation of the fusion loops between the two families suggests that nairoviruses could also use a tri-partite membrane targeting strategy to facilitate viral membrane fusion (Guardado-Calvo et al., 2016). The functional roles of individual loops need to be experimentally determined.

3.7. Epitope sites on CCHFV Gc

Recent research has identified six antigenic sites on CCHFV Gc that protective neutralizing antibodies target: site 1 (Trp 1185, Trp 1199), site 2 (Ser 1309, Val 1314, Asp 1352, Gly 1353), site 3 (Leu 1307, Ile 1229), site 4 (Lys 1297, Asn 1386, Lys 1393), site 5 (Val 1124, Ser 1128, Val 1093) and site 6 (Glu 1500, Asp 1504, Leu 1450, Ile 1522, Leu 1518, Phe 1520) (Fels et al., 2021). We mapped these residues onto the CCHFV Gc trimer and protomer structures (Fig. 6).
Site 1 is located near the fusion loop "cd", and the antibodies targeting this site can effectively block the insertion of the fusion loop into the target membrane. Sites 2 and 3 are embedded in the interior of the trimer, preventing the conformational transition of Gc by blocking the formation of the postfusion homotrimer. Site 4 is situated between the α2 helix and "γ" β-strand. The binding of an antibody to this position probably prohibits the complete folding back of domain III. Site 5 is located at the bottom of domain III and near the linker region between domains I and III, and antibody binding may hinder the movement of domain III. Critical residues in site 6 are located in domain III and binding of antibodies to this site may also affect the free movement of domain III, which is required for the prefusion-to-postfusion transition.

4. Discussion

In this report, we presented a cryo-EM reconstruction of the CCHFV Gc ectodomain at atomic resolution, which enabled us to build a model with clear structural details comparable to previously reported crystal structures of postfusion Gc trimers from other bunyaviral families. During 2D classification of selected sGC-trimer particles, it was observed that most particles in vitreous ice had their three-fold axes oriented perpendicular to the vitreous ice layer, which might be caused by exposure of hydrophobic fusion loops to the air-water interface. Even particles lying flat in the vitreous ice layer showed a head-to-tail arrangement (Fig. 2B, class d), in which the fusion loops of one sGC-trimer might insert into the base region of domain I (near missing residues Glu1094–Ile1100, Gly1123–Ser1130) in the adjacent homotrimer to harbor their hydrophobic fusion loops. Testing with different surfactants to tackle the problem of preferential orientation (Drulyte et al., 2018) may facilitate expansion of cryo-EM studies to other bunyaviral Gc.

In recent years, the range of CCHFV circulation has increased (Bente et al., 2013; Arteaga et al., 2021). This increase has been accompanied by reports of multiple CCHFV strains co-circulating within a geographic region (Goedhals et al., 2014). Our results suggest that CCHFV may have "borrowed" different structural elements from various bunyaviral families, showing a complex evolutionary strategy within this large virus group. Studying the possible transfer of genetic material among bunyaviral families represents an interesting future study.

5. Conclusions

This report describes the atomic-resolution structure of CCHFV Gc in the postfusion conformation using cryo-EM and sheds light on how CCHFV Gc combines structural features from two other bunyaviral families. Our cryo-EM structure of CCHFV Gc provides a structure-based blueprint for developing new neutralizing antibodies that target the glycoprotein surface of this group of important emerging pathogens.

Data availability

The Cryo-EM map has been deposited in the Electron Microscopy Data Bank (EMDB) under accession code EMD-31579, and the corresponding atomic model has been deposited in the Protein Data Bank (PDB) under accession code 7FGF.

Ethics statement

This article does not contain any studies with human or animal subjects performed by any of the authors.

Author contributions

Na Li: conceptualization, formal analysis, investigation, writing–original draft, writing–review and editing. Guibo Rao: data curation, methodology, writing–original draft. Zhiqiang Li: methodology, visualization. Jiayi Yin: data curation. Tingting Chong: methodology. Kexing...
Tian: formal Analysis. Yan Fu: project administration. Sheng Cao: conceptualization, funding acquisition, resources, supervision, writing-review and editing.

Conflict of interest
The authors declare that they have no conflict of interest.

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Appendix A. Supplementary data
Supplementary data to this article can be found online at https://doi.org/10.1016/j.virome.2022.01.015.

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