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Structural characterization of the HCoV-229E fusion core

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ABSTRACT

HCoV-229E spike (S) protein mediates virion attachment to cells and subsequent fusion of the viral and cellular membranes. This protein is composed of an N-terminal receptor-binding domain (S1) and a C-terminal trans-membrane fusion domain (S2). S2 contains a highly conserved heptad repeat 1 and 2 (HR1 and HR2). In this study, the HRs sequences were designed and connected with a flexible linker. The recombinant fusion core protein was crystallized and its structure was solved at a resolution of 2.45 Å. Then we characterized the binding of HR1s and HR2s via both sequence alignment and structural analysis. The overall structures, especially the residues in some positions of HR2 are highly conserved. Fourteen hydrophobic and three polar residues from each HR1 peptide are packed in layers at the coiled-coil interface. These core amino acids can be grouped into seven heptad repeats. Analysis of hydrophobic and hydrophilic interactions between HR2 helix and HR1 helices, shows that the HR1 and HR2 polypeptides are highly complementary in both shape and chemical properties. Furthermore, the available knowledge concerning HCoV-229E fusion core may make it possible to design small molecule or polypeptide drugs targeting membrane fusion, a crucial step of HCoV-229E infection.

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1. Introduction

The genus Coronavirus (CoVs) belong to the order Nidovirales, family Coronaviridae and subfamily Coronavirinae. Recently, the Coronavirus Study Group of the International Committee for Taxonomy of Viruses has proposed three genera: including the mammalian genera Alpha- and Beta- and the bird-associated genera Gamma- and Detacoronavirus recently [1–3], to replace the traditional CoV groups 1, 2, and 3 [4,5]. CoVs are a diverse group of enveloped, plus-stranded RNA viruses that infect humans and many animal species, in which they can cause respiratory, enteric, hepatic, central nervous system and neurological diseases of varying severity [3]. In 2003, a novel aggressive human coronavirus termed Severe Acute Respiratory Syndrome coronavirus (SARS-CoV), which is most related to genus Betacoronavirus, caused a large epidemic with a case fatality rate of ~10%, creating a global panic and economic damage [6]. In 2012, the newly emerging Middle East respiratory syndrome coronavirus (MERS-CoV), a highly pathogenic virus with an approximately 50% fatality rate, can cause a serious respiratory manifestations, including fever, cough, shortness of breath, and acute respiratory distress syndrome (ARDS) [7,8]. To date, it has been shown that four other human coronaviruses HCoV-229E, HCoV-OC43, HCoV-NL63 and HKU1 [9–11], cause a significant portion of upper and lower respiratory tract infections in humans, including common colds, bronchiolitis and neurological disease [12]. To date, no medication is available to effectively prevent or cure HCoV-229E-related diseases. Thus, it is important to design and develop new vaccines or drugs that are specific for the HCoV-229E-related diseases.

CoVs are enveloped virus, suggesting that the fusion between the cell membrane and the viral envelope is essential for the initiation of viral invasion. Following binding with host receptor, CoVs release viral genetic materials into the cytoplasm through direct membrane fusion [13,14]. Membrane fusion is mediated by spike protein S2 subunit. This subunit is expected to present rearrangement of its characteristic elements called heptad repeats (HRs) to form a stable 6-helix bundle fusion core [14]. Although great advances in the knowledge of CoVs protein and host factors

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have been achieved, no licensed vaccines or specific drugs are available to prevent CoVs infection [15]. In view of prevalence of epidemics due to CoVs, it is essential to develop novel broad-spectrum antivirus drugs. Since fusion with the host membrane is a vital early step in the life cycle of the virus, fusion inhibitors could be designed as antiviral drugs. Such as Human immunodeficiency virus I inhibitor T20, a peptide mimic of HR2 and acting through competitively binding to HR1, is the first fusion inhibitor targeting membrane fusion [16,17].

To date, some studies discovered the highly conserved conformation of fusion core, based on four solved crystal structures namely that from NL-63, MHV, SARS and MERS [18–21]. Little if any experimental evidence has been presented to propose that the HCoV-229E fusion core shares the same features. In order to unravel the structure basis of HCoV-229E membrane fusion, we solved the structure of fusion core of HCoV-229E and characterized the binding of HR1s and HR2s via both sequence alignment and structural analysis. The overall structures shows that the HR1 and HR2 polypeptides are highly complementary in both shape and chemical properties. Although HCoV-229E fusion core presenting the highly conserved sequences compare with MHV, SARS, MERS, and NL-63, the HR2 have some differences in stereo configuration. Furthermore, the available knowledge concerning HCoV-229E fusion core may provide important details for the design of small molecule or polypeptide drugs that target HCoV-229E infection.

2. Materials and methods

2.1. Cloning, expression and purification

The fusion core construct (FC1), including different truncated heptad-repeat regions (HR1 and HR2) of HcoV-229E spike protein (GenBank™ accession number BAL45641.1), were predicted by LearnCoil-VMF [22] (Fig. 1) and further modified according to comparison with the available data on CoV pathogenesis in their corresponding conserved regions. In the FC1 construct, HR1 and HR2 regions cover the amino acids 786–853 and 1051–1102, respectively. Subsequently, truncated HR1 and HR2 were connecting with a flexible linker (LVPRSGSGGGGLEVLFQGP). The coding sequence, optimized codons for Escherichia coli, was synthesized by BeijingAugCT Biotechnology Co., Ltd., amplified by PCR and inserted between the BamHI and XhoI restriction sites of pET28a vector (Novagen). The protein production information is showed in Table 1.

The construct expressed protein with an C-terminal hexa histidine tag. Corresponding proteins were over expressed in E. coli BL21 (DE3) (TransGen Biotech). Cells were grown aerobically at 37 °C in Luria-Bertani medium supplemented with 100 μg/ml kanamycin. When the cell density (OD600) reached 0.6, the culture was first cooled for 2 h at 4 °C and then induced at 16 °C for 18 h with 0.5 mM isopropyl-β-D-thiogalactoside (IPTG).

Cells were harvested by centrifugation at 8000g for 15 min. The
cell pellet was re-suspended in McAc 0 buffer (25 mM Tris·HCl, 500 mM NaCl, pH 8.0) and lysed by sonication. Unbroken cells and debris were removed by spinning the lysate at 12000 g for 40 min at 4 °C. Supernatant containing soluble target protein was then loaded onto a Ni-NTA column (GE Healthcare) previously equilibrated with McAc 0 buffer. After thorough washing with buffer, protein bound to the column was eluted with McAc 500 mM imidazole. The protein solution was concentrated to 1 ml and applied to a Resource Q anion-exchange chromatography column (GE Healthcare) equilibrated with buffer A containing 25 mM Tris, 150 mM NaCl, pH 8.0. The fractions were pooled, concentrated to 500 μl, and applied to a Superdex® 200 gel filtration column (GE Healthcare) equilibrated with buffer A containing 150 mM NaCl. The purity of the protein was estimated to be greater than 95% by SDS-PAGE analysis. Then the target protein were concentrated to 15–20 mg/ml and stored at −20 °C until further use.

2.2. Crystallization

Crystals of HCoV 229E fusion core were grown at 293 K using the sitting drop vapour diffusion technique. A 1 μl protein solution (10 mg/ml, 25 mM Tris, 150 mM NaCl, pH 8.0) was mixed with 1 μl reservoir solution and equilibrated over 100 μl of reservoir solution. Crystals were obtained in condition containing 1.5 M ammonium sulfate, 12% (v/v) glycerol, 100 mM Tris/HCl, pH 8.5; The crystal was flash-frozen in liquid nitrogen. Crystallization details are list in Table 2. X-ray diffraction data were collected at beamline 5 A of the Photon Factory (Japan). All data were processed with the HKL2000 Suite of programs [23].

2.3. Data processing, structure solution and refinement

The crystal structure of HCoV-229E fusion core was determined by molecular replacement with the program Phaser [24] using the structure of the HCoV-NL63 N57C42 (PDB code 2IEQ) trimer as the search template. The initial model was first refined with the Phenix AutoBuild wizard [25]. The model was then improved further with iterative cycles of manual rebuilding in coot [26] and refinement by Phenix. refine [27]. All structures were judged to have good stereochemistry according to the Ramachandran plot calculated by MolProbity [28]. A summary of the data collection, phasing and structure refinement statistics is listed in Table 3 and Table 4.

2.4. Analytic ultracentrifugation (AUC)

Sedimentation velocity (SV) experiments were performed with a Beckman/Coulter Optima XL-1 analytical ultracentrifuge or AUC equipped with two-channel centerpieces and sapphire windows (Beckman Coulter). The experiment was carried out at 4 °C using purified protein in buffer containing 25 mM Tris, 150 mM NaCl, pH 8.0. Protein sample (0.3 mg/ml) was centrifuged at 4 °C at 42000 rpm and migration of protein was monitored based on absorbance at 280 nm. The SV data were analyzed with the SEDFIT program [29].

2.5. Circular dichroism spectroscopy

Circular dichroism (CD) spectra were performed on a Biologic W. Zhang et al. / Biochemical and Biophysical Research Communications 497 (2018) 705–712

| Table 1 | Protein production information. |
|---------|--------------------------------|
| Source organism | HcoV-229E |
| DNA source | synthesized by BeijingAugCT Biotechnology Co.,Ltd. |
| Forward primer | CGGGATCCGAAACACAGAAAATTCT |
| Reverse primer | CCCCTGAGGAGCCATTTCGAGTC |
| Cloning vector | pET28a |
| Expression vector | pET28a |
| Expression host | Escherichia coli, |
| Complete amino acid sequence of the construct produced | ENQKILAASFNKAMTNIVDAFTGVNDAITQTSQALQTVA |

| Table 2 | Crystallization conditions. |
|---------|-----------------------------|
| Method | sitting drop vapour diffusion |
| Plate type | 48 well |
| Temperature (K) | 293 |
| Protein concentration | 10 mg/ml |
| Buffer composition of protein solution | 25 mM Tris, 150 mM NaCl, pH 8.0 |
| Composition of reservoir solution | 1.5 M Ammonium sulfate, 12% (v/v) Glycerol, 100 mM Tris/HCl, pH 8.5 |
| Volume and ratio of drop | 1:1 (2 μl) |
| Volume of reservoir | 100 μl |

| Table 3 | Values for the outer shell are given in parentheses. |
|---------|------------------------------------------------------|
| Diffraction source | Photon Factory (Japan) |
| Wavelength (Å) | 1.0000 |
| Crystal-detector distance (mm) | 350 |
| Rotation range per image (°) | 0.5 |
| Total rotation range (°) | 180 |
| Space group | P1 |
| a, b, c (Å) | 30.1, 77.3, 77.4 |
| α, β, γ (°) | 76.9, 85.8, 90.2 |
| Mosaicity (°) | 0.3 |
| Resolution range (Å) | 50.00–2.44 (2.49–2.44) |
| Total No. of reflections | 46795 |
| No. of unique reflections | 24629 |
| Completeness (%) | 97.9 (94.4) |
| Redundancy | 19.1 (1.9) |
| Rmerge | 3.9 (23.5) |
| Overall R factor from Wilson plot (Å²) | 48.8 |
M450 spectropolarimeter (Biologic Science Inc.) equipped with a thermoelectric temperature controller. Spectra of HCoV-229E fusion core (0.1 mg/ml) were measured by taking data points every 1.0 nm from 195 nm to 260 nm at 20°C in 25 mM Tris, 150 mM NaCl, pH 8.0.

3. Results and discussion

3.1. Generation and characterization of HCoV-229E fusion core

From the NCBI database, we obtained the primary sequence of HCoV-229E spike protein (Gen Bank: BAL45641.1). In view of the prediction of the LearnCoil-VMF program and the multiple alignments of the putative heptad repeat regions of HCoV-229E with those of several other coronaviruses, the final construct of the HCoV-229E fusion core was designed. The fusion core consists of HR1, a truncated type of the LearnCoil-VMF predicted HR1 (residues 786–853), connected with HR2, an extended version of the...
Fig. 3. a) Electrostatic potential surface of three central HR1 helices, as calculated using PyMOL, shows the hydrophobic grooves formed between each of the two adjacent HR1 helices. One of three HR2 segments in the grooves are shown in lines representation. The residues that insert into hydrophobic grooves are shown in stick representation. b) The hydrophilic interactions between HR1 and HR2 helices. The residues involved in forming hydrogen bonds are shown in stick representation and are labelled, as well as hydrogen bonds in yellow dashed lines. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
LearnCoil-VMF predicted HR2 (1051–1102) (Fig. 1a) and a flexible 22-residue linker (LVPRSGCGSGGGEVLFQGP). This flexible linker has been shown to work successfully with the fusion core of SARS-CoV and MERS-CoV that allows for easy expression and purification of the fusion core complex [19,30]. The recombinant protein, designated fusion core (FC1), was expressed in E. coli and purified by affinity chromatography and size exclusion chromatography. FC1 was eluted at approximately 15.8 ml on a calibrated Superdex 200 10/300 column, corresponding to a molecular mass of ~45 kDa. AUC was also used to investigate the molecular mass Fig. 1b which gave a value of 49.7 kDa. As the single HR1-linker-HR2 chain is ~15.4 kDa, these data indicate a trimer has formed in solution, as expected. We further tested the protein by CD spectroscopy. Double minima at 208 and 222 nm were observed, demonstrating that the FC1 has a high alpha-helical content (Fig. 1c). The fusion core designed here shared the feature of the helix wheels (Fig. 1d), a typical characteristic of class I fusion protein [19].

3.2. Overall structure of the HCoV-229E fusion core

In the three-dimensional structure of HCoV-229E fusion core, clear electron densities could be traced for residues Q788-G838 in HR1 and T1060-V1097 in HR2. Residues L791-V836 of HR1 fold into a 13-turn \( \alpha \)-helix stretching the entire length of the coiled coil, whereas residues Q788-I790 at the N terminus and residues Q837-G838 at the C terminus form two extended loops, respectively. The HR1 helices form an interior, parallel trimeric coiled-coil. Then three HR2 helices pack in an oblique, left-handed and antiparallel direction into hydrophobic grooves on the surface of this trimeric coiled-coil (Fig. 2a). This leads to a 6-helix bundle of approximately 13 Å in semidiameter and approximately ~80 Å in height (Fig. 2b).

Similar to the core structure of S2 from HCoV-NL63 [18], fourteen hydrophobic and three polar residues from each HR1 peptide are packed in layers at the coiled-coil interface. These amino acids had been grouped into seven heptad repeats (Fig. 2c). In the last two of these repeats, the side-chains of the “a” and “d” residues in one \( \alpha \)-helix point directly into the hole formed between the side-chains of four residues in an adjacent HR1 helix that exhibits canonical knobs-into-holes packing (Fig. 2d). In contrast, the first six heptad repeats of HR1 lack any regular 3\(-\varepsilon\) periodicity. Instead, cross-sectional layers containing an “x-like” symmetric pattern (Fig. 2e) [31] alternate with layers containing a “y-like” packing; see Fig. 2f [32]. Side chains at the \( x \) position project toward the center of the hydrophobic core (Fig. 2e), and similarly alternating side chains at the \( y \) positions pack in a hexagonal arrangement (Fig. 2f) [18]. All these core side chains (excluding alanine) adopt their well-populated rotamer conformations in \( R \)-helices.

3.3. Interactions between the HCoV-229E fusion core helices

We further characterized the atomic details mediating the formation of the hairpin trimer. Consistent with the HR features, the hydrophobic residues at positions “a” and “d” in HR1 are aligned on one side of the helix, forming an interface of strong hydrophobicity.

![Fig. 4.](image-url) In the upper panel, representatives of each genera were chosen for comparison. Superposition of the 2-helix bundle crystal structures of HCoV-229E (yellow) with NL-63 (green), MHV (blue), SARS (violet) and MERS (dark blue) showed ~0.61 Å, ~0.94 Å, ~0.70 Å and ~9.36 Å root mean square deviation for approximate 80 Ca. In the lower panel, it shows that all of the residues at d position of HR2 are highly conserved. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
Three HR1 helices thereby pack against each other and stack the hydrophobic helical laterals in the center of the coiled-coil core. These three hydrophobic grooves become the binding sites of three HR2 helices. Although a notable difference in length is observed between the HR1 and HR2 helices (~13 turns versus ~8 turns), both the N- and C-terminal tails of the HR2 region pack in an orderly manner against the hydrophobic grooves of a central three-helical coiled-coil, mainly through hydrophobic interactions involving L1062, L1064, L1068, L1071 and L1078 at the N-terminal position and L1085, L1089 and L1092 at the C-terminal position of the HR2 helix (Fig. 3a).

The hydrophilic interactions between one HR2 helix and its neighbouring HR1 helices consist of 12 hydrogen bonds, mostly distributed in the regions around the N- and C-terminal ends of the HR2 helices (Fig. 3b). Around the N-terminal position of the HR2 helix, its L1091, S1093 and V1097 form hydrogen bonds with N796 of its fused HR1 helix and Q818, K829 of the adjacent HR1 helix. The concentrated arrangement of hydrogen bonds constitute two anchoring points at both ends of one short HR2 helix; further stabilizing its binding with the central hydrophobic grooves.

### 3.4. Sequence alignment and structure comparison

Since fusion core proteins presented comparatively high sequence similarity within each CoVs genera, representatives of each genera were chosen for comparison. In genera Betacoronavirus, structures from MHV, SAES and MERS are available; in genera Alphacoronavirus, only NL-63 and HCoV-229E; while in genera Gamma- and Delta-coronavirus, no structures are presently available. Superposition of the 2-helix bundle crystal structures of HCoV-229E with NL-63, MHV, SARS and MERS showed ~0.60 Å, ~0.70 Å and ~0.94 Å root mean square deviation (rmsd) for ~80 Cα (Fig. 4), respectively. The common regions include the following peptide fragments: residues 1060–1082 in HCoV-229E, residues 1244–1266 in NL-63, residues 1227–1249 in MHV, residues 1156–1178 in SARS and residues 1258–1280 in MERS. The residues in the α position (1061, T1068, S1075, V1082) and d position (L1064, L1071, L1078), showing relatively conservation, pack against hydrophobic grooves formed on the surface of the HR1 core. All of the helical elements and a majority of the extended HR2 loops could be well aligned. However, some terminal residues exhibit conformational variance, and the helices in HR2 differ in length. As described above, all of the residues at a/d position of HR2 are highly conserved (either Leu and Ile). Their presence is essential to form an interface of strong hydrophobicity. In addition, e/g positions of HR1 are also conserved. Thus, the overall structures and the hydrophobic character of the interaction between HR1 and HR2 is conserved in both fusion cores, suggesting the possibility for broad-spectrum inhibitor design targeting fusion core proteins of all CoVs.

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