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Antiviral drug design based on structural insights into the N-terminal domain and C-terminal domain of the SARS-CoV-2 nucleocapsid protein

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Article

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1. Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is a positive single-stranded RNA virus (+ssRNA) with a genome of ~30 kb [1]. According to sequence alignment and bioinformatics analysis results, SARS-CoV-2 is identified as a member of the genus Betacoronavirus, which also contains SARS-CoV and MERS-CoV [2,3]. The genome of SARS-CoV-2 mainly encodes four structural proteins, sixteen nonstructural proteins, and other accessory proteins. The four structural proteins are spike (S), membrane (M), envelope (E), and nucleocapsid (N) proteins [4,5]. Spike protein mediates the recognition and membrane fusion with host cells by angiotensin-converting enzyme 2 (ACE2), which is the main target of antibody and vaccine development [5–7]. Sixteen nonsstructural proteins, including nsp1–16, mainly mediate viral synthesis and processing, and play important roles in virus proliferation [8,9].

Nucleocapsid (N) protein plays crucial roles in the life cycle of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), including the formation of ribonucleoprotein (RNP) complex with the viral RNA. Here we reported the crystal structures of the N-terminal domain (NTD) and C-terminal domain (CTD) of the N protein and an NTD-RNA complex. Our structures reveal a unique tetramer organization of NTD and identify a distinct RNA binding mode in the NTD-RNA complex, which could contribute to the formation of the RNP complex. We also screened small molecule inhibitors of N-NTD and N-CTD and discovered that ceftriaxone sodium, an antibiotic, can block the binding of RNA to NTD and inhibit the formation of the RNP complex. These results together could facilitate the further research of antiviral drug design targeting N protein.

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the importance of the N protein, its RNA binding and dimerization domains may be crucial targets for drug development.

The structure of the full-length N protein is still unknown due to its intrinsically disordered regions, but the structures of NTD and CTD of the N protein (designated as N-NTD and N-CTD) of viruses from genus Betacoronavirus, such as SARS-CoV-2, SARS-CoV, Middle East respiratory syndrome coronavirus (MERS-CoV), and human coronavirus (HCoV)-OC43, have been reported [23–31]. Several key residues involved in RNA binding and interaction have also been identified. However, the crystal structure of N-NTD bound to RNA and the molecular mechanism of RNP formation remain unclear.

Here, we report the crystal structure of NTD of the N protein of SARS-CoV-2 at a resolution of 1.90 Å, the crystal structure of NTD bound RNA at a resolution of 2.25 Å, and the crystal structure of CTD at a resolution of 3.0 Å. Through the Nuclear Magnetic Resonance (NMR) titration experiment, we identified ceftriaxone sodium, which is the third-generation cephalosporin and a commonly used antibiotic in the clinic, as an inhibitor of RNA binding to NTD. The new crystal structural information from this paper provides a basis for the study of the molecular mechanism of RNP formation in SARS-CoV-2, and could contribute to the development of other antiviral drugs targeting NTD of the N protein and its RNA binding sites.

2. Materials and methods

2.1. Cloning, expression, and purification of N-NTD and N-CTD

The genes coding SARS-CoV-2 N-NTD (residues 48–173) and N-CTD (residues 250–365) were chemically synthesized with codon optimization (GenScript, Nanjing, China). Two genes were cloned into the pET-28a vector (Vazyme Biotech Co., Ltd., Nanjing, China) with a 5’ ATG starting sequence and C-terminal tag of the virus (TEV) protease site followed by a hexa-histidine (6 × His) tag separately. N-NTD and N-CTD constructs were extracted from E. coli DH5α (Tiangen Biotech Co., Ltd., Beijing, China) and then expressed into E. coli BL21(DE3) (Mei5 Biotechnology Co., Ltd., Beijing, China) grown in LB medium. When bacterial cultures were grown to an A600nm of 0.6–0.8 at 37 °C, isopropyl-β-D-1-thiogalactopyranoside (IPTG) with a final concentration of 1 mmol/L was added to induce expression at 16 °C for 18–20 h.

Cells were harvested by centrifugation (4000 r/min, 15 min, 4 °C) (Beckman Coulter, Brea, USA) and then resuspended in lysis buffer (50 mmol/L Tris-HCl pH 8.0, 500 mmol/L NaCl, 5% glycerol). Cells were lysed with a high-pressure homogenizer operating (ATS Engineering Limited, Suzhou, China) at 700–900 bar. Lysates were cleared by centrifugation (18,000 r/min, 30 min, 4 °C) (Beckman Coulter). The supernatant was loaded on Ni-NTA beads (Qianchun Bio, Yancheng, China) and then washed with buffer containing 40 and 60 mmol/L imidazole. Finally, the protein was eluted with lysis buffer containing 250 mmol/L imidazole. The protein was incubated with TEV protease at 4 °C for overnight and passed through Ni-NTA beads (Qianchun Bio) again to remove the 6 × His tag (protein: TEV protease = 100:1). Protein without 6 His tag was concentrated by Centrifugal Filter Unit (Merk Millipore Ltd., Yancheng, China) and then washed with TEV protease at 4 °C and 18,000 r/min, 30 min.

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The supernatant was loaded on Ni-NTA beads (Qianchun Bio, Yancheng, China) and then washed with buffer containing 40 and 60 mmol/L imidazole. Finally, the protein was eluted with lysis buffer containing 250 mmol/L imidazole. The protein was incubated with TEV protease at 4 °C for overnight and passed through Ni-NTA beads (Qianchun Bio) again to remove the 6 × His tag (protein: TEV protease = 100:1). Protein without 6 His tag was concentrated by Centrifugal Filter Unit (Merk Millipore Ltd., Yancheng, China) and then washed with TEV protease at 4 °C and 18,000 r/min, 30 min.

Purification of protein samples for NMR is similar to the above protocol. E. coli BL21(DE3) (Mei5 Biotechnology Co., Ltd.) should be grown in M9 medium containing U-[13C, 15N] (Cambridge Isotope Laboratories, Inc., Turksbury, USA). The SEC buffer for purified protein should contain 20 mmol/L Na2HPO4 pH 5.5, 50 mmol/L NaCl, 0.01% NaN3.

2.2. Preparation of RNA

To examine the binding of RNA and N-NTD, dsRNA was prepared by two RNA oligonucleotides with sequence of 5’-CACUGAC-3’ and 5’-GUCAUGU-3’. To anneal the RNA duplex, two RNA oligonucleotides were mixed at an equal molar ratio in annealing buffer (10 mmol/L Tris-HCl pH 8.0, 25 mmol/L NaCl, 2.5 mmol/L ethylenediamine tetraacetic acid (EDTA)), denatured by heating to 94 °C for 5 min and then slowly cooled to room temperature.

2.3. Crystallization and data collection

SARS-CoV-2 N-NTD, N-CTD, and N-NTD and dsRNA complex were crystallized by the sitting drop method. Crystals of N-NTD were grown with crystal buffer (10 mmol/L NaCl, 100 mmol/L 2-morpholinoethanesulfonic acid (MES) pH 6.0, 20% polyethylene glycol (PEG) 6000 at 20 °C. Crystals of N-CTD were grown with crystal buffer (27 mmol/L NaNO3, 27 mmol/L Na2HPO4, 27 mmol/L (NH4)2SO4, 100 mmol/L Tris, 1 mol/L bicine pH 8.5, 20% polyethylene glycol monomethyl ethers (PEGMME) 500, 10% PEG 20,000, 100 μmol/L 1.8-anilinonaphthalene sulfonate) at 20 °C. Crystals of N-NTD and dsRNA complex were grown with crystal buffer (100 mmol/L MIB buffer pH 6.0, 25% PEG 1500) at 20 °C. The MIB buffer is prepared by mixing sodium malonate, imidazole, and boric acid according to 2:3:3. Crystals are frozen directly in liquid nitrogen.

2.4. Structure determination and refinement

X-ray diffraction data were collected at beamline using a wavelength of 0.979361 Å (NDT), 1.18057 Å (NTD-RNA), or 0.979150 Å (CTD). The structure was determined by molecular replacement (MR) with Phaser (STFC Daresbury Lab, Warrington, UK), using the X-ray crystal structure of SARS-CoV N-NTD (PDB ID: 2OFZ) or N-CTD (PDB ID: 2CJR) as the search model [26,28,32]. All initial models were further refined with Refmac5 (STFC Daresbury Lab, Warrington, UK) as part of CCP4 program (STFC Daresbury Lab) and interactively adjusted by COOT (STFC Daresbury Lab) [33]. The X-ray diffraction and structure refinement statistics are summarized in Table S1 (online).

2.5. Surface plasmon resonance (SPR) assay

The binding affinity of ceftriaxone sodium (MedChemExpress, Monmouth Junction, USA) and RNA to SARS-CoV-2 N-NTD were verified with an SPR assay by a Biacore S200 instrument (GE Healthcare). SARS-CoV-2 N-NTD was immobilized on a CM5 sensor chip (GE Healthcare). The channel of the blank group flowed through the chip with PBS buffer, and the channels of the experimental group were diluted to different concentrations of ceftriaxone sodium (MedChemExpress) or RNA with PBS buffer and flowed through the chip. Equilibrium dissociation constant (Kd) values were calculated and analyzed by Biacore S200 analysis software (GE Healthcare). In the competition experiment between ceftriaxone sodium (MedChemExpress) and RNA, 100 μmol/L ceftriaxone sodium (MedChemExpress) was added to the PBS buffer of blank groups and experimental groups.
2.6. NMR spectroscopy

The NMR experiments were carried out at 25 °C on a Bruker Avance 600 MHz spectrometer (Bruker, Billerica, USA) equipped with a 5 mm triple-resonance TCI cryogenic probe. The U-[15C, 15N] NMR samples (Cambridge Isotope Laboratories, Inc.) were prepared in the SEC buffer, 5% D2O/95%–95% H2O. A series of double- and triple-resonance spectra were recorded to sequence-specific resonance assignments, including 2D 1H-15N HSQC, 3D HNCACB, and 3D CBCA(CO)NH. For NMR titrations with candidate small molecule compounds (TargetMol Chemicals Inc., Boston, USA), 2D 1H-15N HSQC spectra were collected on the 15N-labelled N-NTD/N-CTD in the absence and in the presence of small molecules at different ratios. Small molecular compounds are usually dissolved into DMSO, which may induce shifts in the NMR peaks. To eliminate the effect of DMSO, we dissolved small molecular compounds into the SEC buffer of N-NTD/N-CTD and repeated NMR titration experiments. All NMR spectra were processed and analyzed with Sparky (University of California, San Francisco, USA) and NMRPipe (National Institute of Standards and Technology (NIST), Rockville, USA). The chemical shift perturbations (CSPs) were calculated using the equation:

\[ \text{CSP} = \sqrt{\langle AH \rangle^2 + (0.2\text{AN})^2}, \]

where the significantly perturbed residues are defined as those with the CSP values > average ± STD.

2.7. Molecular docking

The structure of the N-NTD in complex with ceftriaxone sodium (MedChemExpress) was calculated using HADDOCK (Utrecht University Bonvin Lab, Utrecht, the Netherlands) in combination with crystallography and NMR system (CNS) [34,35]. The actual docking was performed using the NMR structure of N-NTD (PDB ID: 6Y13) and the 3D structure of ceftriaxone sodium (PubMed CID: 5479530) (MedChemExpress) according to a standard protocol, which is described as follows. The topology and parameters files of ceftriaxone sodium (MedChemExpress) were generated with ACYPYPE (Vrije Universiteit Brussel Wim F Franken Lab, Brussels, Belgium) [36]. The N-NTD residues with CSP > 0.8 ppm and at least 20% solvent accessibility were chosen as active residues (T49, A50, W52, T54, G60, A90, T91, R92, L104, S105, R107, Y109, L139, A152, and Y172), while adjacent solvent-exposed residues were additionally selected as passive residues (N48, L55, H59, K61, R89, R93, D103, P106, A138, N140, P151, N153, F171, and A173). All atoms of ceftriaxone sodium (MedChemExpress) were defined as active for the experimentally driven docking protocol. In addition, three regions in the N-NTD were defined as fully-flexible segments for the advanced stages of the docking calculation (the N-terminal N48–A50, the central loop I94–M101, and the C-terminal G170–A173). The final set of 200 water-refined structures was clustered using a Fraction of Common Contacts approach with a default cut-off of 0.75. Finally, the structure of NTD-ceftriaxone sodium with the lowest energy score was selected for detailed analysis and display by PyMOL (The PyMOL Molecular Graphics System, version 2.1.1; Schrodinger, New York, USA).

3. Results

3.1. Crystal structure of N-NTD

We designed a SARS-CoV-2 N-NTD construct containing 48–173 residues of the N protein, expressed and purified as the method described (Fig. S1 online). The crystal structure of SARS-CoV-2 N-NTD (PDB ID: 7XX1) was determined to a resolution of 1.90 Å by molecular replacement using the structure of SARS-CoV N-NTD (PDB ID: 2OFZ) as the search model [26]. The final structure was refined to R-factor and R-free values of 0.210 and 0.258, respectively. The additional data are given in Table S1 (online). Each asymmetric unit (ASU) contains four N-NTD molecules with overall root mean square deviation (RMSD) values of 0.43–1.06 Å between each monomer, each of which binds one zinc ion and one MES molecule (Fig. 1a, c). In the tetramer, MES molecules, which are buffer components, are found to form hydrogen bonds with two adjacent N-NTD monomers, making the entire tetramer assume a cyclic configuration (Fig. 1b). Compared with the structures of other N-NTD tetramers, our circular tetramer structure is more stable under the action of MES molecules, which may contribute to the oligomerization of the N protein. All four monomers in one asymmetric unit of N-NTD show a right-handed shape and contain a structural core and a flexible loop similar to other coronaviruses’ N-NTDs [26,30,31]. The structural core comprises of a four-stranded antiparallel β-sheet and a short 310 helix, while the flexible loop is a protruding β-hairpin (Fig. 1c). As depicted on the electrostatic potential surface, the junction between the protruding β-hairpin and the core structure forms a positively charged pocket, serving as a potential RNA binding site (Fig. 1d).

At present, there are several reports on the structures of SARS-CoV-2 N-NTD [23,24]. To further explore the structural information, we compared our structure with the reported structures (PDB ID: 6M3M, 7CDZ) (Fig. 1e) [23,24]. The RMSD values between our N-NTD structure and the other two N-NTD structures are 0.630–1.149 Å (PDB ID: 6M3M) and 0.635–1.190 Å (PDB ID: 7CDZ), respectively. In addition to the differences in the flexible N- and C-termini, there are still two differences in the β-hairpin region. First, compared with the other two N-NTD structures, our N-NTD structure shows an extended β-hairpin, similar to MERS-CoV N-NTD (PDB ID: 4UD1) [31]. Second, the β-hairpin shows a movement towards the RNA binding site as described later. These differences may be caused by the flexibility of β-hairpin and the influence of MES molecules. The MES molecule binds to R107 at the bottom of the β-hairpin region via hydrogen bond interactions (Fig. 1f).

3.2. Crystal structure of N-NTD and RNA complex

The electrostatic potential surface of N-NTD reveals a potential RNA binding site [26,30,37,38]. To obtain the structure of the N-NTD and RNA complex, we selected an RNA sequence (5'-GACUGAC-3' and 5'-GUCAGUG-3') as a duplex of RNA (dsRNA), which binds to N-NTD [38]. The Kd value of N-NTD and dsRNA detected by SPR analysis was 8 mmol/L (Fig. S2a online). The crystal structure of the SARS-CoV-2 NTD-RNA complex (PDB ID: 7XWZ) was solved at a resolution of 2.25 Å. Each ASU contains two N-NTD molecules and two dsRNA fragments (Fig. 2a). Structural superimposition of N-NTD in the apo and RNA-bound forms reveals significant conformational changes within the β-hairpin region (Fig. 2b), highlighting the intrinsic flexibility of these RNA-interacting regions, which would allow each N-NTD monomer binds to dsRNA in different ways. Of our two N-NTD monomers, N-NTD binds predominantly to one of the strands in the dsRNA. In monomer 1, the hydrogen bond interaction between T109, Y111, and RNA involves O2′ atoms and O3′ atoms in the RNA phosphate backbone, the hydrogen bond between R88 and RNA includes only an O1′ atom, while the hydrogen bond between T49 and RNA includes only an O2′ atom (Fig. 2c, d). However, hydrogen bond interactions between R92, R107, and RNA involve O1′ atoms and O2′ atoms in RNA phosphate backbones in monomer 2 (Fig. 2c, e). These binding modes are also different from that of the docking model of the NTD-RNA complex (PDB ID: 7ACS) by NMR (Fig. 2b) [38]. In the NMR-based model, three arginine resi-
dues (R92, R107, and R149) directly bind to RNA. Although the amino acid residues affected by the three binding modes are distinct, the binding site is obviously at the same position, the positively charged junction between the protruding β-hairpin and the core structure. Distinct binding modes may indicate that N-NTD binds to RNA as much as possible in various ways to promote the formation of RNP.

3.3. Structure of N-NTD and ceftriaxone sodium complex

N-NTD and N-CTD contribute to the formation of RNP, so small molecule drugs that bind to them and block this process can be used as inhibitors targeting the N protein. We screened a large number of small molecule compounds from literature reports and compound libraries to identify potential drugs. Finally, we

Fig. 1. Crystal structure of SARS-CoV-2 N-NTD. (a) The overall structure of SARS-CoV-2 N-NTD tetramer (PDB ID: 7XX1). Each monomer is represented by a different color. (b) Hydrogen bonds between an MES molecule and different N-NTD monomers. Each monomer is represented by a different color. Hydrogen bonds are represented by black dotted lines, and amino acids involved in hydrogen bond interactions are marked. (c) The superimposition of four monomers in an asymmetric unit shown as a cartoon representation in two orientations. (d) The electrostatic potential surface of SARS-CoV-2 N-NTD in two orientations. Blue indicates positive charge potential, while red indicates negative charge potential. (e) Superposition (based on Cα) of our SARS-CoV-2 N-NTD structure (PDB ID: 7XX1, wheat) and other published SARS-CoV-2 N-NTD structures (PDB ID: 6M3M, gray; PDB ID: 7CDZ, pink). (f) Hydrogen bonds between an MES molecule and an N-NTD monomer. Hydrogen bonds are represented by black dotted lines, and amino acids involved in hydrogen bond interactions are marked.
identified several candidate compounds. Because of the difficulties in growing crystals of protein and small molecular compounds, we chose to characterize the binding of these small molecule compounds to N-NTD by NMR titration experiments. Residues of N-NTD are specifically perturbed by ceftriaxone sodium dissolved in DMSO or SEC buffer on the 2D 1H-15N HSQC spectra (Fig. S3). Ceftriaxone sodium is a third-generation cephalosporin and was approved by FDA in 1984. It is mainly used for infections caused by pathogens including Neisseria meningitidis, N. gonorrhoeae, and N. lactamica. Meanwhile, the SPR assay also detected the KD value of 8.3 μmol/L for N-NTD and ceftriaxone sodium (Fig. S2b online). Ceftriaxone sodium is a third-generation cephalosporin and was approved by FDA in 1984. It is mainly used for infections caused by pathogens by inhibiting the synthesis of the cell wall.

We added ceftriaxone sodium into the N-NTD protein sample at ratios from 0:1 to 25:1. Superimposition of the 2D 1H-15N HSQC spectra in the presence of ceftriaxone sodium at different ratios revealed remarkable resonance perturbations of many residues (Fig. 3a). Surprisingly, the same small compound can also induce resonance shifts in N-CTD, although relatively few residues were found to move (Fig. 3b). These results indicate that ceftriaxone sodium specifically interacted with both N-NTD and N-CTD. To locate the binding sites of ceftriaxone sodium on the surface of N-NTD, we analyzed chemical shift perturbations (CSPs) values on a per-residue basis. According to CSPs at the max ratio (N-NTD: ceftriaxone sodium = 1:25), a total of 17 residues (T49, A50, S51, W52, F53, T54, G60, A90, T91, R92, L104, S105, R107, Y109, L139, A152, and Y172) showed significant resonance shifts, which, by mapping onto the crystal structure of N-NTD, constituted two binding sites on the surface of N-NTD (Fig. 3a, c). Subsequently, we used the HADDOCK program for the NMR-restraint driven docking simulations of ceftriaxone sodium. The 3D structure with PubChem CID of 5479530 was chosen as a starting conformation of the ceftriaxone sodium. The analysis of CSPs provided a set of “active” solvent-accessible residues on N-NTD that were expanded for surrounding “passive” residues. The selection criteria for active residues were that their CSP values were higher than 0.08 ppm and at least 20% solvent accessibility. The standard docking protocol yielded a set of 200 water-refined structures, in which the structure with the lowest energy score was selected as a representative conformation for the N-NTD in complex with ceftriaxone sodium (NTD-ceftriaxone sodium) (Fig. 3d). As shown in the resulting structure model of NTD-ceftriaxone sodium complex, two main binding sites are located within the junction region, the same location as the RNA binding site (Fig. 3e). Interestingly, significantly perturbed residues include those (R92, R107, and Y109) involved in hydrogen bond interactions under two NTD-RNA binding modes (Fig. 2d, e, Fig. 3d). In addition, we also used SPR to study ceftriaxone sodium and RNA competition for NTD binding. The experimental results revealed that in the presence of 10- to 20-fold excess of ceftriaxone sodium, no RNA binding signal to N-NTD was detected, indicating that ceftriaxone sodium can compete with RNA for N-NTD binding (Fig. S2c online). Therefore, with the N protein as the drug target, our results suggest that ceftriaxone sodium is an inhibitor by blocking the interaction between NTD and RNA.

3.4. Crystal structure of N-CTD

We also designed a SARS-CoV-2 N-CTD construct containing 250–365 residues of the N protein, expressed and purified as the method described (Fig. S3 online). The crystal structure of SARS-CoV-2 N-CTD (PDB ID: 7XWU) was determined by molecular replacement at a resolution of 3.0 Å. The final structure was refined to R-factor and R-free values of 0.219 and 0.268, respectively. The additional data are given in Table S1 (online). Eight N-CTD molecules are paired to form four homo-dimers in one ASU, each dimer also binding one phosphate ion (Fig. 4a). Each monomer is composed of one 310 helix, three α-helices, and two β-strands (Fig. 4b, c). N-CTD molecules form stable dimers, consistent with the result of analytical gel filtration (Fig. S4c online). Extensive hydrogen bonds and hydrophobic interactions stabilize the dimerization of N-CTD. In the dimer, two β-strands from each monomer form an antiparallel β-sheet with eight hydrogen bonds interactions, including S318-Y333, S327-K338, T329-L339, L331-I337, Y333-G335, T334-T334, G335-Y333, G335-Y333, I337-L331, and L339-T329 (residues written in front are from monomer 1) (Fig. 4d). There are also many hydrogen bonds in the interaction region of α-helices from two monomers, such as R277-G316, G278-R319, N285-I320, G316-R277, R319-G278, R319-E280, R319-Q283, and
Fig. 3. NMR characterization of N-NTD/N-CTD and ceftriaxone sodium. (a) Superimposition of HSQC spectra of N-NTD in the absence (blue) and the presence of ceftriaxone sodium at a molar ratio (N-NTD: ceftriaxone sodium) of 1:25 (cyan). The assignments of the significantly perturbed residues are labeled on the right. (b) Superimposition of HSQC spectra of N-CTD in the absence (red) and the presence of ceftriaxone sodium at a molar ratio (N-CTD: ceftriaxone sodium) of 1:100 (purple). (c) Residue-specific chemical shift perturbations (CSPs) of N-NTD in the presence of ceftriaxone sodium at a molar ratio (N-NTD: ceftriaxone sodium) of 1:25. The significantly perturbed residues are defined as those with the CSP values at 1:25 > 0.08 ppm (average + STD) (dark dashed line). The surface representation of N-NTD is colored according to the CSP values in a white-to-red scale. Color intensities correlate with CSP values, with darker colors indicating larger values. (d) The structure of the NTD-ceftriaxone sodium complex with ceftriaxone sodium in sticks and N-NTD in cartoon (left) and electrostatic potential surface (right) are shown in two orientations. (e) Superimposition (based on Cs) of ceftriaxone sodium-bound form and two dsRNA-bound forms of N-NTD reveals a similar binding interface. For the ceftriaxone sodium-bound form of N-NTD, only ceftriaxone sodium is shown.
I320-N285. In addition, residues F286, T296, W301, A305, and Q306 from monomer 1 show hydrophobic interactions with residues A311, S312, and G316 from monomer 2 in other interaction regions (data not shown).

We overlaid our N-CTD structure with reported structures (PDB ID: 7CE0, 7DE1) [23,39]. Because of extensive hydrogen bonds and hydrophobic interactions, all structures show similar and stable dimer structures and characteristics with RMSD of 0.95–0.97 Å (Fig. 4e). Stable dimer structures are also essential elements in the formation of RNP.

4. Discussion and conclusion

As the pandemic of COVID-19 enters its third year, several vaccines and drugs have been approved. However, this public health crisis has not been completely controlled due to the rapid mutation of SARS-CoV-2, highlighting the urgent demand to find additional potential drug targets and new drugs. As one of the important structural proteins of SARS-CoV-2, N protein plays a crucial role in many physiological processes of the virus, especially when it combines with viral RNA to form an RNP complex [10,11]. In this study, we presented the crystal structures of N-NTD and N-CTD and compared them with previously published structures. The comparison results indicate that these structures are very similar to each other. NTD shows a right-handed shape, while CTD exists as a stable homo-dimer.

Based on the reported N-NTD structures of coronaviruses, it has been proposed that NTD binds to RNA through its positively charged pocket [26,30,37,38]. Our NTD-RNA complex structure, which was not reported before, reveals that NTD and dsRNA display a specific and different RNA binding mode. The involved residues include R88, Y109, and Y111 in our monomer 1, R92, and R107 in our monomer 2, which are different from R92, R107, and R149 in the docking model as reported before [38]. Residues inter-
acting with RNA in other coronaviruses’ N-NTDs have also been reported, such as R106, R107, and R117 in hCoV-OC43 (corresponding to R92, R93, and K102 in SARS-CoV-2) [30]. Although involved residues in different binding modes are not exactly the same, they include positively charged arginine, which may play a major role in RNA binding. Different binding modes indicate that N-NTD could bind to RNA in many different ways.

As one type of cephalosporin, cetrixazone sodium is often used to treat bacterial infections, such as wound infections, respiratory tract infections, and intra-abdominal infections. Currently, there is an ongoing clinical trial to verify the therapeutic efficacy of cetrixazone sodium in lung infections caused by COVID-19 [40]. Nevertheless, we first confirmed the inhibitory effect of this small molecule on the N protein of SARS-CoV-2. According to the process of RNP formation, there are two main antiviral strategies for the N protein: blocking the packaging of the viral RNA or inhibiting the oligomerization of the N protein. Drugs previously screened only inhibit the function of NTD, such as PJ34 [37,41].

Nevertheless, cetrixazone sodium we screened shows inhibitory effects both on NTD and CTD. According to the docking model of the NTD-cetrixazone sodium complex, cetrixazone sodium mainly affects 17 residues, including those binding to dsRNA (R92, R107, and Y109). Meanwhile, the SPR assay also detected that the Kd value of N-NTD and cetrixazone sodium was 8.3 μmol/L, and cetrixazone sodium could compete with RNA for N-NTD binding within a similar concentration range. Therefore, cetrixazone sodium blocks the interaction between NTD and RNA by occupying the RNA binding site, inhibiting the formation of RNP and further interrupting the lifecycle of SARS-CoV-2. Its mechanism of inhibiting CTD remains to be further explored. Recently OMICRON (B.1.1.529) has been reported as a variant of concern. However, mutations of the N protein (P131L, R203K, and G204R) only occurred in three IDR regions, not the conserved NTD and CTD domains [42,43]. Therefore, these sites that bind to RNA can serve as conserved drug targets, and cetrixazone sodium may also serve as an effective inhibitor for OMICRON.

In conclusion, we reported and analyzed the crystal structures of NTD and CTD from the N protein of SARS-CoV-2 as well as the crystal structure of the NTD-RNA complex. Our structures revealed a novel tetramer configuration of NTD and identified a distinct binding mode of RNA binding by NTD. We also identified cetrixazone sodium as an inhibitor that blocks RNA binding of NTD and inhibits the formation of RNP, which could interrupt the life cycle of SARS-CoV-2. This structural information presented in this paper could offer new insights into the follow-up studies of antiviral drug design targeting N protein.

Conflict of interest

The authors declare that they have no conflict of interest.

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Author contributions

Shuyang Zhang, H. Eric Xu, and Yi Xue designed and supervised the study. Xiaodong Luan performed the protein purification and structure analysis. Xinming Li performed NMR-related experiments and molecular docking. Yufan Li assisted in the work of Xiaodong Luan and finished the manuscript writing and figure preparation. Xinming Li, Gengchen Su, Wanchoa Yin, and H. Eric Xu participated in the manuscript and figure preparation. Other authors have provided facilities and funds to assist in this study. All authors read and approved the manuscript.

Appendix A. Supplementary materials

Supplementary materials to this article can be found online at https://doi.org/10.1016/j.scib.2022.10.021.

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