Structure-Based Stabilization of Non-native Protein–Protein Interactions of Coronavirus Nucleocapsid Proteins in Antiviral Drug Design

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ABSTRACT: Structure-based stabilization of protein–protein interactions (PPIs) is a promising strategy for drug discovery. However, this approach has mainly focused on the stabilization of native PPIs, and non-native PPIs have received little consideration. Here, we identified a non-native interaction interface on the three-dimensional dimeric structure of the N-terminal domain of the MERS-CoV nucleocapsid protein (MERS-CoV N-NTD). The interface formed a conserved hydrophobic cavity suitable for targeted drug screening. By considering the hydrophobic complementarity during the virtual screening step, we identified 5-benzyloxygramine as a new N protein PPI orthosteric stabilizer that exhibits both antiviral and N-NTD protein-stabilizing activities. X-ray crystallography and small-angle X-ray scattering showed that 5-benzyloxygramine stabilizes the N-NTD dimers through simultaneous hydrophobic interactions with both partners, resulting in abnormal N protein oligomerization that was further confirmed in the cell. This unique approach based on the identification and stabilization of non-native PPIs of N protein could be applied toward drug discovery against CoV diseases.

INTRODUCTION

Small-molecule stabilization of protein–protein interactions (PPIs) is an extremely promising strategy in drug discovery. It can be used to treat cancers and viral infections.1–3 Stabilizing PPIs with small molecules may be allosteric or direct (also called orthosteric). This process alters the oligomerization equilibrium of the protein and enables small molecules to modulate its physiological function.4–7 The anticancer drug paclitaxel, for example, allosterically enhances microtubule structure assembly by binding to β-tubulin.8,9 On the other hand, rapamycin, another anticancer agent, binds directly to the interface between FKBP12 and mTOR and stabilizes the structure of the complex.10 The most well-characterized PPIs suitable as targets for drug development form natively under physiological conditions. However, non-native interactions, which may form under extreme circumstances such as inside a crystal lattice, are also potential drug targets. For example, nucleozin exerts its antiviral activity by stabilizing the non-native PPI interface between the two neighboring nucleoprotein trimers within the influenza virus, which results in abnormal protein oligomerization and loss of viral viability.11

Middle East respiratory syndrome coronavirus (MERS-CoV) belongs to the betacoronavirus (β-CoVs) family. It causes severe respiratory distress with a high mortality rate in humans.12–14 Recently, a closely related novel coronavirus, coronavirus disease 2019 (COVID-19), caused an outbreak of pneumonia in Wuhan, which further underscored the risk of CoVs to the global public health.15,16 However, there is no

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effective treatment for CoVs. Thus, there is an urgent need to
develop new antiviral agents against CoVs. MERS-CoV packages its genome in a nucleocapsid (N) protein and forms a
ribonucleoprotein (RNP) complex. The RNP is essential for
viral transcription and assembly. Several studies suggested that
the modulation of CoV N protein oligomerization by small
molecules is a feasible antiviral drug development strategy.
The CoV N protein is organized into the N-terminal domain
(NTD) and the C-terminal domain (CTD), with both
domains participating in RNA binding. All CoV N-NTD
structures are folded in a monomeric conformation. In
contrast, the CoV N-CTDs are always dimeric and are
responsible for N protein oligomerization via protein−protein
interactions.

Here, we report the crystal structure of MERS-CoV N-NTD by molecular replacement (MR) using
the structure of HCoV-OC43 N-NTD (PDB ID: 4J3K) as the search model. The final structure was refined to R-factor and
R-free values of 0.26 and 0.29, respectively, at a resolution of 2.6 Å (Table S1). Each asymmetric unit contained four N-
NTD molecules assembled into two identical dimers with an
overall RMSD of 0.28 Å between the dimers (Figure S1A,B).

Results

Structure of the N-Terminal Domain of the MERS-
CoV N Protein. We determined the crystal structure of
MERS-CoV N-NTD by molecular replacement (MR) using
the structure of HCoV-OC43 N-NTD (PDB ID: 4J3K) as the search model. The final structure was refined to R-factor and
R-free values of 0.26 and 0.29, respectively, at a resolution of 2.6 Å (Table S1). Each asymmetric unit contained four N-
NTD molecules assembled into two identical dimers with an
overall RMSD of 0.28 Å between the dimers (Figure S1A,B).

The structures showed that P3 induces abnormal full-length N protein
oligomerization in vitro and at the cell level. We also described
the structure of MERS-CoV N-NTD complexed with 5-
benzyloxygramine and revealed its stabilizing mechanism. Our
findings provide insight into the development of a new
therapeutic approach based on stabilizing a non-native protein
interaction interface. It may lead to the discovery and
development of new treatments for various infectious diseases.

Figure 1. Structure and sequence of MERS-CoV N-NTD. (A) Overall structure of the MERS-CoV N-NTD dimer containing monomers 1 and 2 is
depicted as a cartoon and colored yellow and green, respectively. Residues involved in dimerization are shown as sticks and highlighted in (B). (B)
Interactions among MERS-CoV N-NTDs. Dimerization is mediated mainly by vector-fusion residues interacting with the conserved hydrophobic
regions on the core structure (first area) along with the residues surrounding it (second area). Interacting residues on monomers 1 and 2 are
labeled in black and blue, respectively. Vector-fusion and conserved hydrophobic regions are colored cyan and red, respectively. The color of all
other interacting residues is the same as that for each monomer in (A). Polar contacts are indicated with red dashed lines. (C) Upper panel: close-
up of the interacting region of vector-fusion residues. The surface was colored according to the hydrophobicity level at the protein surface. Vector-
fusion residues (black) are shown as sticks to emphasize the hydrophobic pocket. Lower panel: 2D diagrams of the interaction between the
hydrophobic pocket and the vector-fusion residues. The latter are labeled in black. The hydrophobic contacts are indicated with black dashed lines.
(D) Sequence alignment of various CoV N proteins in the N-terminal region. Red letters indicate strictly conserved residues. Cyan indicates
conservative substitution sites. Hydrophobic regions involved in unusual dimerization are indicated by black triangles.
among the CoVs. In our structure, however, the loop connecting strands β2 and β3 protruding out of the core into other CoV N proteins was absent. Unlike the reported structures that have a monomeric conformation, our structure was atypically dimeric.

Figure 1B shows the details of the interactions in the MERS-CoV N protein dimer. We named the units monomer 1 and monomer 2 (Figure 1A). According to the amino acid composition of the binding site on monomer 2, we divided the dimeric interface into two areas: one located on the N-terminus flexible region and the other on the loop between β4 and β5 of the N protein. In the first area, W43, N66, N68, Y102, and F135 of monomer 1 generated a conserved hydrophobic pocket permitting the side chain of M38 of monomer 2 to enter this hole by a hydrophobic contact (Figure 1C,D). H37 and N39 of monomer 2 were packed against W43 and F135 of monomer 1, respectively, and contributed to the hydrophobic interaction. The side chains of N39 of monomer 2 formed one hydrogen bond with the N68 backbone in monomer 1 at a distance of 2.6 Å. The second area was relatively more hydrophilic. The main chain oxygens of G104, F135, and T137 of monomer 2 to enter this hole by a hydrophobic contact (Figure 1C,D). H37 and N39 of monomer 2 were packed against W43 and F135 of monomer 1, respectively, and contributed to the hydrophobic interaction. The side chains of N39 of monomer 2 formed one hydrogen bond with the N68 backbone in monomer 1 at a distance of 2.6 Å. The second area was relatively more hydrophilic. The main chain oxygens of G104, F135, and T137 of monomer 2 formed three hydrogen bonds with the side chains of Q73 and T134 of monomer 1 at distances of 3.8, 3.2, and 3.7 Å, respectively. The side chain of N139 on monomer 2 formed a hydrogen bond with the main chain oxygen of T137 on monomer 1 at a distance of 3.6 Å. The interactions of the first and second areas comprise buried surface areas (BSA) of 289 and 103 Å², respectively. The small surface area buried at the interface accounts for ∼5 kcal mol⁻¹ binding energy, which translates to a dissociation constant of ∼200 mM. Thus, the dimer described here is unique in that it is non-native and relies on vector-fusion residues (H37 and M38) to maintain its dimeric status. This property may also explain why the present structure has an oligomeric status different from previously reported structures for the CoV N protein. We used cross-linking experiments to analyze the oligomeric capacity of MERS N-NTD containing the vector-fusion residues in solution. MERS N-NTD had a dimeric conformation in solution. Our structure indicated that W43 played an essential role in forming the hydrophobic pocket accommodating the vector-fusion residues and, therefore, mediated the N-NTD dimer formation. The W43A mutation significantly reduced the oligomeric tendency of N-NTD (Figure S1C). This further supports that the “exogenous residues” encoded by the vector backbone mediated the formation of the non-native dimer. We also superimposed the previously published structure of MERS-CoV N-NTD (PDB ID: 4ud1) containing a native
N-terminal flexible region with our dimer structure. The side chain of N38 in the native structure could not interact with the hydrophobic pocket as the former is hydrophilic and short (Figure S1E). Thus, it may be possible to utilize small compounds to replace the vector-fusion residues and stabilize the PPI through hydrophobic interactions.

Direct Targeting of the Non-native Dimer Interface for Antiviral Screening. We performed a structure-based virtual screening by targeting W43 in the hydrophobic pocket of the N-NTD dimeric interface. H37 and M38 were removed from the template to identify compounds that could replace the vector-fusion residues and, therefore, contribute to the stabilizing effect (Figures 2A and S2A,B). We chose the highest-scoring hits, listed in Table S2, based on shape complementarity, the presence of aromatic moieties, and the ability to stack onto W43 of N-NTD. Because the formation of the non-native dimers was primarily mediated by hydrophobic interactions in our structure (Figure 1C,D), we next considered the hydrophobic complementarity between the acquired ligands and N-NTD in the form of the lipophilic match surface (S_L/L). We also took into account the ability of the drug to permeate cells by aiming for lower topological polar surface areas (TPSA). Based on the above criteria, three compounds were finally chosen for further study. Benzyl-2-(hydroxymethyl)-1-indolinecarboxylate (P1) and 5-benzyloxystyrylamine (P3) had higher S_L/L and docking scores and lower TPSA. The clinical drug etodolac (P2) had a comparable microenvironment surrounding the tryptophans of the protein increased in rigidity and hydrophobicity in the presence of P3. The result also indicated that P3 bound more tightly to the N protein than P1 or P2 by interacting with the W43 pocket (Figure 2B). Fluorescent thermal stability assays disclosed that the N-NTD denaturation melting temperature had increased from 42 to 45 °C when P3 was added. The sigmoidal melting curve for MERS-CoV N-NTD changed in the presence of P3. The delay in protein denaturation suggests that P3 had a favorable therapeutic index among the lead compounds tested in this study. Therefore, P3 is an excellent candidate inhibitor against MERS-CoV.

Structural Model of P3-Induced MERS-CoV N Protein Aggregation. We used SAXS to assess the effects of P3 on the full-length MERS-CoV N protein structure. The fitted distance distribution function of the protein with and without P3 are shown in Figure 3A. P3 increased the maximum distance distribution function of the protein with and without P3 (Figure S1E). Thus, it may be possible to utilize small compounds to replace the vector-fusion residues and stabilize the PPI through hydrophobic interactions.

| Compound | CC<sub>50</sub> (μM) | EC<sub>50</sub> (μM) | TI (CC<sub>50</sub>/EC<sub>50</sub>) |
|----------|---------------------|---------------------|-------------------------------|
| P1       | 459.69              | >100                | NA                           |
| P2       | 569.77              | >100                | NA                           |
| P3       | 805.32              | 32.1                | 25.1                         |

(CC<sub>50</sub>: Half maximal toxicity concentration. EC<sub>50</sub>: Half maximum effective concentration. TI: Therapeutic index. NA: Nonavailable.)
Figure 3. P3-induced abnormal aggregation on the full-length MERS-CoV N protein. (A–E) SAXS analysis of the full-length MERS-CoV N protein. (A) Normalized results from GNOM showing pairwise distance distribution $P(r)$ and maximum distance. The radius of gyration fitted to 207 and 230 Å for the N protein and the N-P3 complex, respectively. "$r$" represents pairwise distances. (B, C) Scattering profiles of the N protein (B) and the N-P3 complex (C) and normalization fitting with GNOM (dashed lines). (D, E) Representative models of the N protein (D) and the N-P3 complex (E) generated by CRYSOL simulations of the SAXS data. Only $\alpha$ carbons are shown. NTD (yellow), CTD (green), and disorder region (cyan). (F, G) Conformation (F) and stability (G) analyses based on FL spectra of the MERS-CoV N protein (1 μM) incubated with P3 (10 μM) for 1 h in a buffer consisting of 50 mM Tris-HCl, 150 mM NaCl (pH 8.3). (H) Schematic of the P3 inhibition mechanism. Left panel: in the absence of RNA, N proteins organize as a dimeric building block contributed by N-CTD dimerization. Middle panel: P3 promoted the dimerization of N-NTDs from different building blocks, by which the distance between CTD cuboids was shortened and N protein aggregation occurred. Right panel: octameric conformation of building blocks buried in the RNA-binding surface of N-CTDs. It hindered the formation of filamentous ribonucleocapsids.
to confirm our SAXS findings. Immunofluorescence microscopy (Figure 4C) showed condensation of the intracellular N protein fluorescence signal in infected Vero E6 cells treated with 50 \( \mu \)M P3. Thus, P3 may induce intracellular N protein aggregation. At 100 \( \mu \)M, P3 suppressed N protein expression in most cells. However, a few presented with intense N protein signals. P3 may have restrained the MERS-CoV N proteins inside the infected cells that promoted the formation of new virions that could not be released. In this way, the adjacent cells could not be infected with MERS-CoV. The data, therefore, suggest that P3 may inhibit MERS-CoV by inducing abnormal aggregation of the N protein inside the cell. This finding is consistent with the results of our structure-based assays.

Crystal Structure of MERS-CoV N-NTD Complexed with Potent Compounds. We attempted to obtain crystals of MERS-CoV N-NTD in complex with compounds P1, P2, and P3 by cocrystallization or ligand-soaking. With the exception of P2, the complex structures of N-NTD with P1 and P3 were solved at resolutions of 3.09 and 2.77 Å, respectively (Table S1). The overall structures of the complexes resembled that of apo-MERS-CoV. Both complexes revealed well-defined unbiased densities in the dimer interface and permitted detailed analysis of the interactions between the compounds and MERS-CoV N-NTD (Figure 5). The interactions between the N protein and each compound were calculated with the Discovery Studio Client (v19.1.0.18287). Most interactions were hydrophobic contacts, which were consistent with our selection rationale. In the P1 complex, N68, F135, and D143 on monomer 1 and V41, G106, P107, and T137 on monomer 2 packed against P1 to create a dimer (Figure 5A). In addition, two nonbonding interactions were detected between P1 and the monomers. There was a \( \pi \)-anion interaction between the benzene ring of the P1 indoline moiety and D143 of monomer 1. There was also a \( \pi \)-donor hydrogen bond between the other P1 benzene ring and the T137 side chain of monomer 2 (Figure 5B). Relative to P1, P3 bound more deeply into the dimer interface and interacted with a larger number of residues on both N-NTD monomers. The amino acid composition of this binding region was W43, N66, N68, S69, T70, N73, and F135 on monomer 1 and V41, G104, T105, G106, A109, and T137 on monomer 2. These residues along with P3 generated a massive hydrophobic driving force allowing the proteins and ligands to pack against each other and stabilize the dimeric conformation of the N protein (Figure 5C). Several nonbonding interactions were also observed at the P3-binding site. These included the interaction between the P3 benzene ring and N68 of monomer 1 and A109 of monomer 2 via \( \pi \)-lone pair and \( \pi \)-alkyl interactions. The dimethylaminomethyl moiety of P3 was a major source of nonbonding interactions. Three \( \pi \)-cation interactions formed between this moiety and the aromatic groups of W43 and F135 in monomer 1. This moiety also formed a \( \pi \)-lone pair interaction with N66 and a \( \pi \)-sigma interaction with W43 of monomer 1 (Figure 5D). The structural analyses explain the comparatively stronger binding of P3 to N-NTD (Figure 2B) and corroborated the thermal stabilization effects (Figure 2C) and antiviral activities (Table 1) of the compounds.

DISCUSSION AND CONCLUSIONS

The use of small molecules to stabilize PPI interfaces has been shown to be a viable approach for the development of new therapeutics. Most compounds created by structure-based
design to manipulate the PPI depend on detailed knowledge of the native interacting interfaces. In contrast, most of the compounds with therapeutic potential that stabilize non-native PPI interfaces were discovered by chance alone. The anti-influenza compound nucleozin was initially discovered using a chemical genetics approach, and its ability to stabilize non-native nucleoprotein oligomers was elucidated much later. Swinholide A, a cytotoxic marine macrolide, has been known to disrupt the actin cytoskeleton and act as an anticancer agent, but it took ten years to discover that it stabilized G-actin as a non-native homodimer complex. However, these examples did underscore the importance of hydrophobicity as a crucial factor stabilizing protein–protein and protein–ligand associations.

Here, we demonstrated the possibility of using the hydrophobic interactions on non-native interfaces as targets for virtual screening. Combined with suitable selection criteria focusing on both shape and hydrophobic complementarities between the ligand and the receptor, small-molecule compounds that stabilize non-native PPIs may be identified in a rational manner. Using the above approach, we successfully identified a compound P3 that affected the biochemical activity of our target protein and showed efficacy against our target pathogen MERS-CoV (Figures 2 and 4). P3-mediated non-native MERS-CoV N-NTD dimerization induced abnormal N protein aggregation by influencing the oligomeric behavior of N-CTD and eventually halting its function in RNP formation. To the best of our knowledge, this structure-based strategy for targeting non-native interfaces has never been proposed for therapeutic design. Thus, non-native interaction interfaces in proteins may comprise a new drug development target class. For β-coronaviruses such as MERS-CoV, the amino acids comprising the non-native interaction interface on N-NTD are relatively conserved. The hydrophobic pocket surrounding W43 and F135 on monomer 1 is shared among other β-coronaviruses. The interacting surface on monomer 2, which includes G104, T105, G106, and A109, is also highly conserved (Figure S4). This conservation may provide certain advantages when developing compounds with broad-spectrum activity against a target pathogen family, including COVID-19. When we tested P3 on mouse hepatitis virus (MHV), we observed a reduction in viral titer, indicating that P3 may also inhibit MHV replication (Figure S5). On the other hand, targeting non-native interaction interfaces is not trivial. As interface formation is induced by an external agent, computations aimed at predicting native PPI structures may not be able to identify potential non-native interfaces. Nevertheless, several stratagems may assist in the identification of potential non-native interaction interfaces. One strategy is to
search for target protein structures with crystal packing contacts known to be biologically irrelevant. Another approach is to identify weakly interacting sites through NMR chemical shift perturbation and hydrogen-deuterium exchange MS.35 Once these potential non-native interaction interfaces are identified, standard screening and then functional characterization may be conducted for small compounds that bind to the interface.

To summarize, we demonstrated that non-native interaction interfaces might form when proteins abnormally oligomerize. Using the MERS-CoV N protein as an example, we showed that non-native interfaces might serve as targets for small-molecule, structure-based screening. We also proposed several stratagems to rationally identify potential non-native interface targets for screening. We believe that we have discovered and tested an alternative drug discovery paradigm that could help expand the repertoire of lead compounds against various pathogens.

## EXPERIMENTAL SECTION

### Chemicals

The compounds P1, P2, and P3 were purchased from Maybridge Chemical Company, TCI Chemicals, and Sigma-Aldrich Corporation, respectively. The reagents used in this study were purchased from Sigma Chemical Co. (St. Louis, MO). The purity of all compounds is higher than 95% and was used without further purification.

### Cloning, Protein Expression, and Purification

The MERS-CoV N proteins were prepared according to previously described methods.36 In brief, the cDNA fragments of MERS-CoV N proteins were cloned into a PET-28a expression vector (Merck, Darmstadt, Germany) containing a histidine tag-encoding sequence. Vectors encoding the single mutants N39A, N39G, and W43A were generated using the QuikChange site-directed mutagenesis protocol with the primers listed in Table S3. The vectors were transformed into Escherichia coli BL21 (DE3) pLyS5 cells. The cells were grown to an optical density range of 0.6–0.8 at 600 nm at 37 °C and protein expression induced with 1.0 mM isopropyl β-D-1-thiogalactopyranoside (IPTG), followed by incubation at 10 °C for 24 h. The cells were harvested by centrifugation (6000g, 12 min, 4 °C) and resuspended in lysis buffer (150 mM NaCl, 50 mM Tris-HCl, 15 mM imidazole, and 1 mM PMSF; pH 7.5). The cells were lysed by sonication and centrifuged (10000g, 40 min, 4 °C) to remove debris. The supernatant was purified by injection into a Ni-NTA column (Merck, Darmstadt, Germany) and eluted with buffer containing imidazole at a gradient range of 15–300 mM. Pure protein fractions were collected, dialyzed with low-salt buffer, concentrated, and quantified by the Bradford method (BioShop Canada Inc., Burlington, ON, Canada).

### Crystallization and Data Collection

MERS-CoV N-NTD crystals were grown as previously described.36 MERS-CoV N-NTD was crystallized at room temperature (−25 °C) by the sitting-drop vapor-diffusion method. A protein solution (2 μL; 10 mg mL⁻¹) was mixed with an equal volume of crystallization solution consisting of 75 mM ammonium sulfate, 2 mM NaBr, and 29% PEG 3350 (Sigma-Aldrich Corp., St. Louis, MO) and equilibrated against a 300 μL solution. MERS-CoV N-NTD:P3 co-crystals were obtained using a crystallization solution containing 2 mM P3. MERS-CoV N-NTD crystals in complex with P1 were obtained by soaking native MERS-CoV N-NTD crystals for 90 s at room temperature in a crystallization solution containing 2 mM P1. Diffraction datasets for MERS-CoV N-NTD alone and in complex with P1 were collected at beamline 13B1 of the Taiwan Light Source (TLS) of the National Synchrotron Research Center (NSRRC; Hsinchu City, Taiwan). Diffraction of the MERS-CoV N-NTD:P3 complex was performed at the beamline SP44XU of SPring-8 (Hyogo, Japan).

### Structural Determination and Refinement

Diffraction data were processed and scaled with HKL-2000 software. The structures were solved by molecular replacement (MR) in Phenix37 using HCoV-OC43 N-NTD (PDB:7fj3k) as the search model.38 The initial models were rebuilt and refined with Coot39 and Phenix. Structures were visualized using PyMOL (The PyMOL Molecular Graphics System, version 2.3.0).40

### Chemical Cross-Link Assay

Protein solutions containing 40 μM of wild-type or mutated MERS-CoV N-NTD were incubated with glutaraldehyde at a final concentration of 1% v/v. The reaction was conducted at room temperature for 10 min and quenched with the addition of 1 M Tris-HCl (pH 7.5). The samples were then stored on ice and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE).

### Discovery of Orthosteric PPI Stabilizers for MERS N Protein

To screen for compounds that induce hydrophobic PPI between MERS N-NTDs, a model of dimeric MERS-CoV N-NTD without the H37 and M38 residues was used in virtual drug screening. The Sigma-Aldrich, Acros Organics, and ZINC drug databases were screened with LIBDOCK molecular docking software to obtain compounds acting on the N protein. The N protein binding pocket was represented by a set of spheres. Each compound in the database was docked in a pocket comprising W43. The hydrophobic complementarity between ligands and receptors was calculated with PLATNUM.31 Compounds with higher docking scores are listed in Table S2.

### Fluorescence Measurements

Fluorescence assays were performed in a buffer consisting of 50 mM Tris-HCl (pH 7.5) and 150 mM NaCl. One micromolar N protein was incubated either with the control buffer or each compound (10 μM) at 4 °C for 1 h. Tryptophan fluorescence was acquired with a Jasco FP-8300 fluorescence spectrometer (JASCO International Co. Ltd., Tokyo, Japan) at an excitation wavelength of 280 nm and an emission wavelength range of 300–400 nm.

### Thermostability Measurements

Thermostability assays were conducted in a buffer consisting of 50 mM Tris-HCl (pH 7.5) and 150 mM NaCl and with a Jasco FP-8300 fluorescence spectrometer (JASCO International Co. Ltd., Tokyo, Japan). One micromolar N protein was incubated either with the control buffer or each compound (10 μM) at 4 °C for 2 h. UV absorbance vs temperature profiles were acquired by ramping the temperature from 4–95 °C at a 1 °C min⁻¹ and recording the absorbance at 280 nm every 0.5 min.

### Determining CC₅₀ and EC₅₀ of Hit Compounds

Vero E6 cells were infected with MERS-CoV with M.O.I = 0.1 and treated with lead compounds for 48 h. Cell viability was determined by the neutral red uptake assay. CC₅₀ and EC₅₀ were determined by % cell viability. CC₅₀ was determined for cells treated with drugs only. EC₅₀ was determined for cells treated with N-CoV N-NTD crystals in complex with P3.

### Small-Angle X-ray Scattering (SAXS) Experiments

SAXS experiments were performed at the BL23A SAXS beamline at the TLS of NSRRC, using a monochromatic X-ray beam (λ = 0.828 Å), with an integrated HPLC system of an Agilent-Bio SEC-3 300 Å column (Agilent Technologies, Inc. Santa Clara, CA). Protein samples (44 μM MERS-CoV N and MERS-CoV N:P3 complex prepared by incubating the 44 μM native protein with 440 μM P3) were prepared in a buffer consisting of 50 mM Tris-HCl (pH 8.5) and 150 mM NaCl on ice for 1 h. Then, a 100 μL aliquot was injected into the column at a flow rate of 0.02 mL min⁻¹. After passing through the column, the sample solution was directed into a quartz capillary (2 mm dia.) for subsequent buffer and sample SAXS measurement at 288 K. The sample-to-detector distance of 2.5 m used covered a scattering vector q range of 0.01–0.20 Å⁻¹. Here, q is defined as q = (4π/λ) sin θ/2θ with the scattering angle 2θ. Thirty-six frames were collected for each sample elution with an X-ray frame exposure time of 30 s. Frames of good data overlapping (namely, of low radiation damage effects) were merged for improved data statistics and analyzed to determine initial R₁ using PRIMUS (version 3.1). The P(r) distance distribution and Dmax were calculated from the experimental scattering curve using GNOM (version 4.1). An ensemble optimization method (EOM) analysis was performed through the EM BL Hagenburg web interface.41 Modeling of the rigid body crystal structure was calculated and generated using CRYSTALS (ATSAS Program Suite v. 2.8.2).31 The
crystal structures of MERS-CoV NTD (PDB ID: 4UD1) and MERS-CoV NTD:P3 (solved in this study) and the CTD domain of MERS-CoV N protein (PDB ID: 6G13) were used as rigid bodies in EOM analysis. With the EOM analysis, 1000 models were generated in the beginning as a structural pool. Selected from the SAXS profiles of the structural pool was an ensemble of models that could fit the experimental scattering curve with their linear combination. Tetrameric MERS-CoV NP conformations and 16-mer MERS-CoV:P3 conformations were selected because their ensemble generated curves fit best to the experimental SAXS results.

Viral Infection. Vero E6 cells (ATCC No: CRL-1586) were seeded onto culture plates with complete Dulbecco’s modified Eagle’s medium (DMEM) and incubated overnight prior to infection. MERS-CoV (HCoV-EMC/2012) at a multiplicity of infection (M.O.I.) of 0.1 was added to the cells and incubated at 37 °C for 1 h, followed by washing thrice with phosphate-buffered saline (PBS) to remove the unattached virus. Fresh complete culture medium was then added to the plates.

Plaque Assay. Vero E6 cells were seeded in 12-well plates and incubated overnight before the assays. Samples containing MERS-CoV were serially diluted 10x with MEM, added to the wells, and incubated for 1 h with agitation every 15 min. After incubation, the inocula were removed and washed with PBS. An overlay medium comprising 2% MEM and 1.5% (w/v) agarose (1:1) was added to the cells followed by incubation at 37 °C and 5% CO2 for 3 days. The plaques were fixed with 10% (v/v) formalin containing 0.2% (w/v) crystal violet, and the plaques were counted.

RT-qPCR. Total RNA of infected Vero E6 cells was extracted with the RNeasy Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. Reverse transcription and PCR amplification were performed with an iQ™ Universal One-Step RT-qPCR Kit (Bio-Rad Laboratories, Hercules, CA). Real-time PCR was performed in a StepOnePlus Real-Time PCR System (Applied Biosystems, Foster City, CA). The primer pairs used to amplify the viral RNA were as follows: GAPDH-F: 5’-GAAGATGGTGATGGGATTTC-3’; GAPDH-R: 5’-GGGATTTCACCTGGTCAATAC-3’. The MERS RNA levels were normalized to that of GAPDH and compared between MERS-CoV groups at 24 h.p.i. and at 48 h.p.i.

Immunofluorescence Assay. Vero E6 cells were seeded in eight-well chamber slides and incubated overnight prior to infection with MERS-CoV at M.O.I. = 0.1. The cells were fixed with 4% (v/v) paraformaldehyde for 20 min at 4 °C, followed by permeabilization in 0.1% (v/v) Triton X-100 for 10 min. Then, 7.5% (v/v) BSA was used as a blocking buffer for 30 min at 37 °C. Anti-MERS-CoV N primary antibody (1:500 dilution; Thermo Fisher Scientific, Waltham, MA) for 1 h at room temperature. 4’,6-Diamidino-2-phenylindole (DAPI) was added during the PBS wash. MERS nucleocapsid expression was examined under a confocal microscope (LSM-700; Carl Zeiss AG, Oberkochen, Germany).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/10.1021/acs.jmedchem.9b01913. Dimeric conformation of MERS-CoV N-NTD (Figure S1); LibDock pose of dimeric MERS-CoV N-NTD with potent compounds (Figure S2); P3-induced aggregation of the MERS-CoV N protein sequesters the RNA-binding region on the N-CTD dimer (Figure S3); potential target sites for broad-spectrum antiviral compound development (Figure S4); antiviral activity of P3 against MHV (mouse hepatitis virus) (Figure S5); crystallographic data collection and refinement statistics (Table S1); pW43 docking pose with chemical structures, docking scores, and biochemical properties of 17 potential hits (Table S2); primers used for mutagenesis in this study (Table S3); molecular formula strings (PDF)

Molecular Formula Strings. JMC, revision (CSV)

Accession Codes

For PDB codes 6KL2 (native), 6KL5 (N-P1 complex), and 6KL6 (N-P3 complex), authors will release the atomic coordinates and experimental data upon article publication.

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Jia N. Hsu, Shih C. Lin, Yong S. Wang, and Ming H. Hou designed research; Jia N. Hsu, Shih C. Lin, Shan M. Lin, Ching M. Chien, and Yong S. Wang performed research; Hung Y. Wu, U S. Jeng, K. Kehn-Hall, and Ming H. Hou contributed new reagents or analytic tools; Shih C. Lin, Shan M. Lin, Ching M. Chien, and Ming H. Hou analyzed the data; Shih C. Lin, Shan M. Lin, Chung K. Chang, Ching M. Chien, and Ming H. Hou wrote the paper.
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ABBREVIATIONS
PPIs, protein–protein interactions; MERS-CoV, Middle East Respiratory Syndrome Coronavirus; NTD, N-terminal domain; CTD, C-terminal domain; N, nucleocapsid; β-CoVs, betacoronaviruses; RNP, ribonucleoprotein; SAXS, small-angle X-ray scattering; MR, molecular replacement; BSA, buried surface areas; TPSA, topological polar surface areas; CC50, cytotoxic concentration; EC50, effective concentration; Δrmax, maximum dimension; rgy, radius of gyration; MHV, mouse hepatitis virus

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