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Hotspot residues and resistance mutations in the nirmatrelvir-binding site of SARS-CoV-2 main protease: Design, identification, and correlation with globally circulating viral genomes



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ABSTRACT

Shortly after the onset of the COVID-19 pandemic, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has acquired numerous variations in its intracellular proteins to adapt quickly, become more infectious, and ultimately develop drug resistance by mutating certain hotspot residues. To keep the emerging variants at bay, including Omicron and subvariants, FDA has approved the antiviral nirmatrelvir for mild-to-moderate and high-risk COVID-19 cases. Like other viruses, SARS-CoV-2 could acquire mutations in its main protease (M^{pro}) to adapt and develop resistance against nirmatrelvir. Employing a unique high-throughput protein design technique, the hotspot residues, and signatures of adaptation of M^{pro} having the highest probability of mutating and rendering nirmatrelvir ineffective were identified. Our results show that ~40% of the designed mutations in M^{pro} already exist in the globally circulating SARS-CoV-2 lineages and several predicted mutations. Moreover, several high-frequency, designed mutations were found to be in corroboration with the experimentally reported nirmatrelvir-resistant mutants and are naturally occurring. Our work on the targeted design of the nirmatrelvir-binding site offers a comprehensive picture of potential hotspot sites and resistance mutations in M^{pro} and is thus crucial in comprehending viral adaptation, robust antiviral design, and surveillance of evolving M^{pro} variations.

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1. Introduction

RNA virus evolution depends on regular recombination events, which are frequently accompanied by escape mutations and result in enhanced transmissibility and stronger adherence to host cell-surface receptors [1]. Moreover, certain mutations confer resistance to the available drugs and antibodies [2], which facilitate the evasion of the virus by the immune system, thereby developing uncertainty about the efficacy of the therapeutics [3]. Along with viral recombination, constant replication of the virus in immuno-compromised people or inter-species spill-over & spill-back

events may also enable the accumulation of mutations to develop viral fitness and immune system evasion [4]. Thus, investigating the virus evolution is crucial to help comprehend the viral evolutionary mechanisms which control its pathogenicity and transmission.

Despite the development and widespread administration of several potent vaccines, the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) continues to infect people with Coronavirus disease 19 (COVID-19) and claim lives, making it possibly the most lethal virus of this century. Pfizer's Paxlovid (a combination of nirmatrelvir [PF-07321332] tablets and ritonavir tablets) has been approved as the first oral antiviral for the treatment of COVID-19 by the US Food and Drug Administration (FDA). In December 2021, Paxlovid was granted emergency use authorization (EUA) for adults and pediatric patients. Nirmatrelvir is emerging as a useful drug for treating mild-to-moderate and high-risk COVID-19 cases [5]. It is a potent inhibitor of the SARS-CoV-2 main protease (M^{pro}) with an inhibition constant (K_1) of ~1 nM [6], an EC₅₀ value of

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~16 nM [7], and an IC₅₀ value ranging between 22 and 225 nM [8]. Using a reversible covalent mechanism, nirmatrelvir, via its cyano group, reacts with the catalytic Cys145 residue of the M^{pro}. SARS-CoV-2 is continually evolving and generating new sequence variants globally. While the emergence of new variants is inevitable, predicting their evolution, spread, and impact through intense research may help improve countermeasures [9.10]. However, as current reports suggest the emergence of resistance mutations in SARS-CoV-2 against nirmatrelvir [11-15], we envisioned carrying out a cohesive and comprehensive study to identify all the potential resistance mutations and hotspots in the M^{pro} of SARS-CoV-2. In the current study, we used a unique high-throughput ligand-based interface protein design protocol to identify potential mutational hotspots in the nirmatrelvir-binding site in the SARS-CoV-2 M^{pro}. Notably, we correlate our findings with the globally circulating viral genomes and show that our design methodology can correctly predict the hotspot residues and adaptable mutations in the M^{pro}.

2. Methodology

The crystal structure of SARS-CoV-2 M^{pro} in complex with nirmatrelvir (PDB ID: 7TLL) was used to obtain the M^{pro}'s interacting residues with nirmatrelvir [6] and then prepared using Molecular

Operating Environment (MOE), v2022.03. Subsequently, a highthroughput protein design involving the resistance scan methodology of MOE was employed to identify single-point mutations of the nirmatrelvir-binding site in M^{pro} that exhibit the highest potential for adaptability and drug resistance [16,17]. To sample the mutations that are not lethal to the virus and more likely to evolve naturally, the nirmatrelyir-binding sites were mutated to single nucleotide polymorphisms (SNPs) of the native sequence only. From the nirmatrelvir-bound M^{pro} complex, 25 drug-binding residues of M^{pro} were designed, except the catalytic dyad (Cys145 and His41), because of their role in substrate cleave and processing [18,19]. For exhaustive sampling and designed site flexibility, the ensemble protein design protocol with the rotamer explorer option was utilized, followed by the root mean square deviation (RMSD) threshold of 0.5 Å. Other important parameters, including the "conformation limit", "fix residues farther than," and "energy window," were set to 25 K, 4.5 Å, and 10 kcal/mol, respectively. A total of 210 designed mutations were generated, and the relative binding affinity of the mutation to the native M^{pro} (represented by dAffinity, which is the Boltzmann average of the relative affinities of the ensemble) was computed for the designed M^{pro}-nirmatrelvir complexes. The Arpeggio webserver was used to analyze the intermolecular interactions between nirmatrelvir and M^{pro} in the

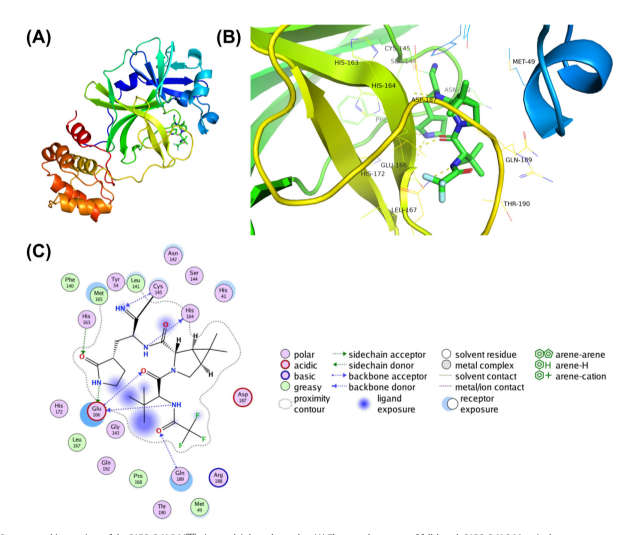


Fig. 1. Structure and interactions of the SARS-CoV-2 M^{pro}-nirmatrelvir bound complex. (A) The crystal structure of full-length SARS-CoV-2 Mpro is shown as a cartoon, and bound nirmatrelvir is shown as a stick. (B) The nirmatrelvir-interacting residues of M^{pro} are shown as lines along with hydrogen bond interactions as yellow dashed lines. (C) A 2D representation showing various types of interactions between M^{pro} and nirmatrelvir, where various types of intermolecular interactions are labeled in the legend. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Table 1 M^{pro} -nirmatrelvir interacting residues that were designed with corresponding SNPs of the native sequence to emulate the mutations that are more likely to happen naturally over the evolution of M^{pro} .

S. No	M ^{pro} -nirmatrelvir interacting and designed native residues ^a	Sampled SNPs in designs
1	Leu27	RQHIMFPSWV
2	Cys44	RGFSWY
3	Met49	RILKTV
4	Pro52	ARQHLST
5	Туг54	NDCHFS
6	Phe140	CILSYV
7	Leu141	RQHIMFPSWV
8	Asn142	DHIKSTY
9	Gly143	ARDCESWV
10	Ser144	ACLFPTWY
11	His163	RNDQLPY
12	His164	RNDQLPY
13	Met165	RILKTV
14	Glu166	ADQGKV
15	Leu167	RQHIMFPSWV
16	Pro168	ARQHLST
17	His172	RNDQLPY
18	Phe181	CILSYV
19	Val186	ADEGILMF
20	Asp187	ANEGHYV
21	Arg188	NCQGHILKMPSTW
22	Gln189	REHLKP
23	Thr190	ARNIKMPS
24	Ala191	DEGPSTV
25	Gln192	REHLKP

^a The catalytic dyad residues His41 and Cys145 of M^{pro} were not designed, as they carry out the acylation-deacylation reaction and cleavage of the substrates.

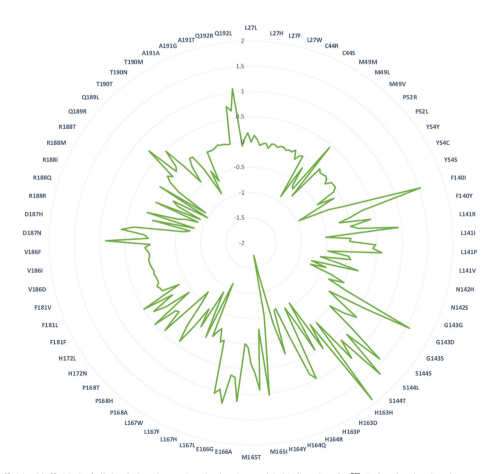


Fig. 2. Relative binding affinities (dAffinities) of all the designed mutations in the nirmatrelvir-binding site of M^{pro}. Radar plot showing the computed dAffinities for the most plausible resistant and adaptable single-point mutant designs of M^{pro} bound to nirmatrelvir. The mutated designs are labeled in the radar plot, and the corresponding dAffinity values are shown.

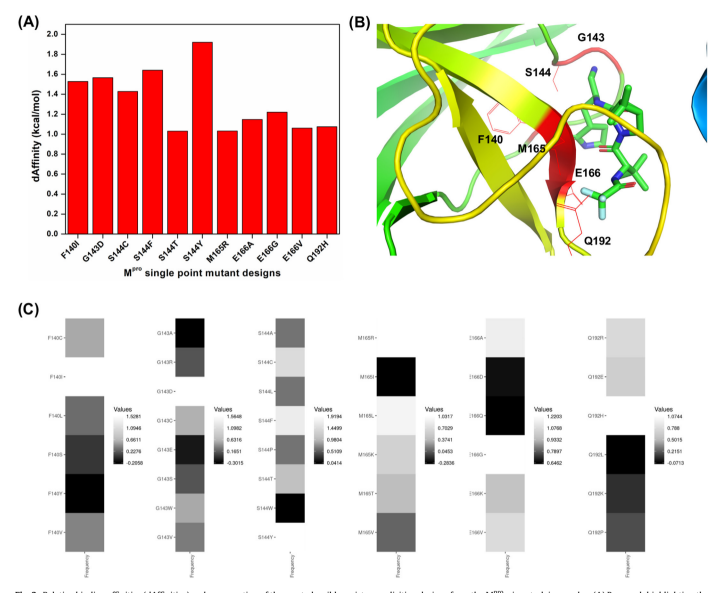


Fig. 3. Relative binding affinities (dAffinities) and conservation of the most plausible resistance-eliciting designs from the M^{pro}-nirmatrelvir complex. (A) Bar graph highlighting the dAffinities for the most plausible resistance-eliciting mutants of M^{pro} against nirmatrelvir, where designs exhibiting dAffinity > 1.0 kcal/mol suggest a decrease in affinity towards nirmatrelvir and hence their ability to become resistant easily. (B) hotspot residues having the greatest tendency to undergo resistance are shown within the protein. (C) heatmaps showing the dAffinity values of every designed mutation in each hotspot, thus highlighting their conservation within the protein. Here, the frequencies of the designed mutations in the hotspot residues are shown in black to white colors as *per* their dAffinity values, respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

low-affinity and high-affinity designs of the M^{pro}-nirmatrelvir complex [20]. Finally, to validate the protein design protocol and correlate the designed M^{pro} mutations with available experimental data, the designed M^{pro} residues that bind to nirmatrelvir were compared with globally circulating SARS-CoV-2 sequences deposited in the Global Initiative on Sharing Avian Influenza Database (GISAID) enabled CoV-GLUE-Viz database, after which their frequencies of occurrence were obtained to demonstrate design accuracy [21,22].

3. Results

Structural inspection of the nirmatrelvir-M^{pro} crystal structure revealed 25 interacting residues and were subjected to design (Fig. 1A & B). Employing the resistance scan module of MOE, the 25 nirmatrelvir-interacting M^{pro} residues were mutated to naturally evolving SNPs (Table 1). This led to 210 single-point mutants, of

which the relative binding affinities of the mutations to the native M^{pro} (dAffinity) were computed, where a higher positive dAffinity denoted lower affinity with nirmatrelvir, thereby implying easy tolerance and resistance of the mutant to nirmatrelvir. The computed dAffinity of 210 single mutants ranged from 1.91 to -1.75 kcal/mol, of which ninety-four M^{pro}-designed mutants demonstrated positive dAffinity values. These results implied tolerated and resistant mutants (Fig. 2). Moreover, to shortlist the most potential resistant mutants, when a strict cut-off of dAffinity >1.0 kcal/mol was used, it resulted in the identification of eleven M^{pro} mutants. The hotspots were mostly from 140, 143, 144, 165, 166, and 192 positions of the M^{pro}. Fig. 3 shows the hotspot residues and potential resistant mutants against nirmatrelvir. Interestingly, this analysis suggested that positions 144 and 166 of M^{pro} are crucial and of high susceptibility for tolerance, adaptability, and conferring resistance during evolution.

Next, a critical analysis was performed focusing on the accuracy

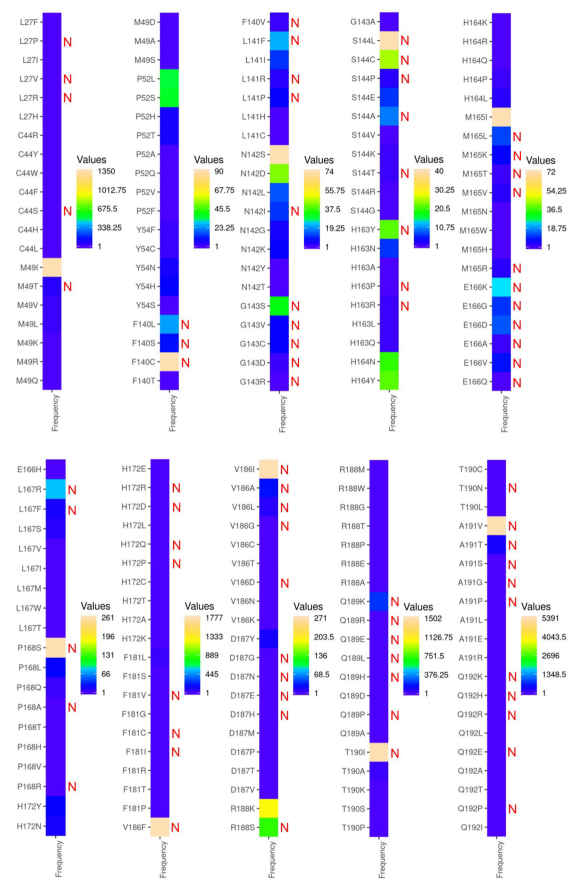


Fig. 4. Heat maps of the nirmatrelvir-bound M^{pro} mutations with their frequencies of occurrence retrieved from GISAID-enabled CoV-GLUE-Viz databases. Mutations at the nirmatrelvir-binding site of M^{pro} obtained from the GISAID-enabled CoV-GLUE-Viz databases are shown. Frequencies of the mutations in SARS-CoV-2 sequences from the COVID-19 pandemic ranged from lower to higher numbers and from blue to orange colors, respectively. The designed mutants that potentially developed resistance and adaptation towards nirmatrelvir are denoted as 'N' in red. It was found that out of 199 mutations, 78 mutations were predicted as positively selected and resistant in the design computations, therefore attaining ~40% correlation, already exist in the SARS-CoV-2 sequences and are currently circulating. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Table 2List of prevalent M^{pro} mutations and their corresponding SARS-CoV-2 lineages.

S. No	Prevalent M ^{pro} mutations ^a	Lineage
1	G15S	C.37 (Lambda)
2	T21I	B.1.1.318
3	L89F	B.1.2
4	K90R	B.1.351 (Beta)
5	P132H	B.1.1.529 (Omicron)
6	L205V	P.2 (Zeta)

^a The most prevalent M^{pro} mutations are not part of the nirmatrelvir-binding site.

of our ligand-based design protocol [17] and assessing how closely our designed single-point mutants of M^{pro}-nirmatrelvir corroborate with the experimentally determined SARS-CoV-2 sequences from the COVID-19 pandemic. As SARS-CoV-2 is actively mutating due to poor immunity, drug, and vaccine pressures, it resulted in the accumulation of >1000 unique mutations at the M^{pro}, thereby continuously being added into the GISAID and released through CoV-GLUE-Viz. We obtained all the deposited mutations and their frequencies of the nirmatrelvir-bound M^{pro} site and compared them with our designed mutants. The comparison showed that seventy-eight mutations (out of 199) were predicted as resistantcausing and hence crucial in conferring adaptability as obtained from the design calculations, thereby achieving ~40% corroboration with the clinical-sequencing data (Fig. 4). Several of the globally circulating, frequently found mutants such as A191V, T190I, V186I, V186F, P168S, S144L, and F140C already have high dAffinity values as obtained from our computational design. This implies that these mutants are adaptive and resistant to nirmatrelvir, even without significant drug or selection pressure (Figs. 3 and 4). However, it is important to comment that the dAffinities are not directly comparable with frequencies obtained from the databases and designed mutants (dAffinities >1.0 kcal/mol), for instance, S144C, which are being circulated as high-frequency mutants (Figs. 3 and 4).

Notably, while this manuscript was being prepared, a preprint article experimentally reported 11 mutants at the drug-binding site of M^{pro} that are resistant to nirmatrelvir ($K_i > 10$ -fold increase) [13]. This offered us an opportunity to critically assess the designed M^{pro} mutants with that of the experimentally reported resistant mutants. Upon examination, it was found that several of the designed mutants, such as S144A, S144F, S144Y, M165T, E166Q, and H172Q. were already found to be resistant and naturally occurring. These mutants are reported to be nirmatrelvir resistant, with K_i values increasing between 19.2 and 38.0-fold, therefore corroborating our findings and reflecting the accuracy of our design methodology [13]. Interestingly, the E166V mutant, experimentally shown to confer the strongest resistance (~100-fold) [23], was found to be one of the most plausible resistance-causing mutants in our design experiments (Fig. 3). Moreover, our targeted design of the nirmatrelvir-binding site in M^{pro} provided a comprehensive picture of other potential hotspot sites and resistance mutations that may emerge if the drug pressure keeps mounting and the pandemic endures its devastating course.

Since the spike protein of SARS-CoV-2 has acquired some permanent mutations, such as the D614G, which continues to be a part of the new variants, a similar quest was conducted for potential clues in M^{pro}. It was found that there were a few mutations in the M^{pro} that were highly prevalent in different lineages (Table 2). Interestingly, these mutations were not in the nirmatrelvir-binding site of the SARS-CoV-2 M^{pro} and hence were not designed (Table 1). Nonetheless, it is possible that there could be a presence of (i) higher-order mutants (such as double, triple, quadruple mutants, and so on) which may harbor some of the mutations from the nirmatrelvir-binding site and/or (ii) compensatory mutations that

are part of the nirmatrelvir-binding site and/or (iii) with mutations of other proteins, such as the spike protein. Although this information is not yet available to the best of our knowledge, resistance to nirmatrelvir may escalate by the simultaneous acquisition of two or more higher-order mutational combination events. A study is underway in this direction.

A further inspection of the high-frequency mutants A191V. T190I, V186F, P168S, S144L, and F140C with respect to the lineages revealed that these mutants pertain to some of the globally distributed lineages that are found in nearly all the COVID-19 prevalent countries. More specifically, the F140C mutant is detected in six, S144L in 23, P168S in 69, V186F in 102, V186I in 41, T190I in 126, and A191V in 285 different lineages (a total of 652 lineages), out of which 421 lineages (64.5%) were found to be common across all the seven high-frequency mutants (Supplementary Table 1). This suggests that the SARS-CoV-2 strains with certain residuespecific mutations of M^{pro} are already circulating in humans, even without significant nirmatrelvir pressure. Remarkably, this result validated our ligand-based design protocol and signifies that several of our M^{pro}-designed mutants (having high dAffinity values) may materialize to be highly valuable in drug resistance and surveillance of newly evolving mutants and viral strains as SARS-CoV-2 encounters more immunogenicity, drug, and vaccine pressure, and as the high-throughput sequencing information keeps deposited.

Finally, the intermolecular interactions between nirmatrelvir and M^{pro} in the low-affinity and high-affinity designs were visualized and obtained. While the design with high affinity had 431 interactions, the design with lower affinity had only 407 interactions. The proximal and van der Waals interactions were decisive in influencing the affinity between nirmatrelvir and M^{pro}. The interaction diagrams between the high-affinity and low-affinity design demonstrated that losing a critical hydrogen bond between nirmatrelvir and Cys145 reduces the affinity and develops resistance and adaptability (Supplementary Figure 1).

4. Discussion

A recent phase 2-3 double-blind, randomized, controlled trial showed that the administration of Paxlovid to symptomatic COVID-19 patients resulted in an 89% lower risk of disease progression to severe conditions and quickly reduced viral load without evident safety issues [24,25]. However, the rapid and regular emergence of new SARS-CoV-2 variants highlights the fatal nature of the virus to mutate and illustrates that the COVID-19 pandemic will stay in the foreseeable future. Continuous surveillance and prediction of residues prone to mutations that can render therapies ineffective can emerge as a crucial repository for managing the spread of emerging mutants and controlling the pandemic. Currently. > 3 million SARS-CoV-2 genome sequences have been submitted to the GISAID website. The use of high-throughput technologies can considerably extend the insights gathered during drug development, improve drug efficacy and safety, predict the challenging multiple variants, and strengthen the countermeasures. Therefore, one may either wait for the new resistant and adaptable variants to develop and then battle it or anticipate it coming and expand the treatment arsenals in advance. With the currently available data and the emergence of new SARS-CoV-2 variants, the scientific community should be prepared to tackle the potential nirmatrelvir-resistant variants and intensify the development of additional interventions.

Nonetheless, we recognize certain limitations of the study, firstly, the requirement of the experimental and functional validation of the nirmatrelvir-resistant designs. Secondly, it is also reported that mutations distal from the catalytic or active site (e.g., HIV-1 protease) can also cause drug resistance, which is not

addressed in this work. In conclusion, despite our recent results identified critical residues in the RNA-dependent RNA polymerase (RdRp) that could render SARS-CoV-2 resistant to remdesivir, molnupiravir, and favipiravir [26,27], a more systematic study to develop a unifying methodology is in progress to address some of these challenges. Nonetheless, our present study is significant with information on the potential structural and residue-specific sites in the M^{pro} that are susceptible to mutation under drug pressure and lead to viral adaptation with fitness advantages.

Author contributions

AKP carried out all the design experiments, data generation, and analysis. AKP and TT conceived the study, participated in its design and coordination, and drafted the manuscript. Both authors read and approved the final manuscript.

Data availability statement

Data will be made available upon reasonable request to the corresponding authors.

Declaration of competing interest

The authors declare no competing interests.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.bbrc.2022.09.010.

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