The Remarkable Selectivity of Nirmatrelvir

Damien Y. Duveau and Craig J. Thomas*

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ABSTRACT: The SARS-CoV-2 main protease is among the most attractive targets for the development of therapeutic interventions for COVID-19. Successful candidate agents will not only possess potent on-target activity versus SARS-CoV-2 Mpro but also minimal polypharmacology versus human cysteine proteases. This Viewpoint explores the activity profile of the first approved SARS-CoV-2 Mpro inhibitor (Nirmatrelvir) versus a panel of cysteine proteases and considers the therapeutic implications of the data.

KEYWORDS: SARS-CoV-2, cysteine proteases, SARS-CoV-2 Mpro, Caspase 1, Nirmatrelvir, polypharmacology

The marshalling of rapid and comprehensive medical advances in response to the COVID-19 pandemic has been astonishing. Among the most promising new tools in the therapeutic armament is Nirmatrelvir (1, PF-07321332) (Figure 1A).1 Nirmatrelvir is a potent inhibitor of the SARS-CoV-2 main protease (Mpro), which is responsible for cleavage of two viral polyproteins (pp1a and pp1ab) into smaller effector protein members of the multiprotein replicase–transcriptase complex (RTC) required for viral RNA replication.2,3 SARS-CoV-2 Mpro is a cysteine protease, a protein class with significant precedent for being druggable as evidenced by multiple published inhibitors including several investigational drugs.

Among the best characterized examples of cysteine protease inhibitors are agents targeting earlier Mpro variants associated with previous SARS outbreaks. Scientists at Pfizer were at the forefront of these efforts with the development of the SARS-CoV-1 Mpro inhibitor PF-00835231 (2) which was later noted to have cross activity versus SARS-CoV-2 Mpro (Ki = 0.271 nM).4 Limited oral bioavailability for PF-00835231 sent the Pfizer team back to the drawing board in hopes of identifying an easily administered and efficacious therapy for COVID-19. The extraordinary effort that followed produced Nirmatrelvir which boasted remarkable clinical efficacy when combined with Ritonavir (i.e. Paxlovid) with an 89% reduced risk of hospitalization or death in the EPIC-HR study. Emergency use authorization was granted by the FDA in December of 2021.

Cysteine proteases are a conserved proteolytic enzyme class utilized across the entire spectrum of living organisms.5,6 This includes over 100 characterized cysteine proteases in the human genome. The success and failure of all drugs is manifestly tied to issues of tolerability which, generally speaking, can be improved by eliminating unwanted cross-

activity versus related (and unrelated) protein/enzyme classes (i.e., polypharmacology). For Nirmatrelvir this surely involved a nominal level of cross-activity against human proteases including cysteine proteases. Owen et al. provided key selectivity data for Nirmatrelvir by demonstrating a lack of activity in a panel of seven proteases including selected cysteine (3), serine (3), and aspartyl (1) proteases.7

Potent inhibition of SARS-CoV-2 Mpro and lack of activity at other proteases is based on the chemical structure of Nirmatrelvir. Nirmatrelvir’s chemical structure represents a modified peptide sequence designed to enhance binding at the target of interest. The Nirmatrelvir structure incorporates a hybrid proline and Z-L-tert-butyl glyceride dimer. A similar structural motif was employed by Vertex Pharmaceuticals in the design of the caspase 1 inhibitor Belnacasan (3, VX-765) (Figure 1A).7 Belnacasan is a prodrug that utilizes a cleverly designed 5-ethoxydihydrofuran-2(3H)-one moiety which is susceptible to intracellular esterase cleavage to reveal the active aldehyde drug form VRT-043198 (4). The aldehyde functionality represents a pharmacological warhead that forms a covalent interaction with the active site cysteine residue. Such “warheads” are a common feature shared by most cysteine protease inhibitors. Nirmatrelvir utilizes a nitrile group as a covalent reversible warhead which aided both SARS-CoV-2 Mpro activity as well as improved physicochemical properties that supported oral bioavailability. Previously,
Boxer et al. incorporated a similar nitrile functionality onto the Belnacasan peptide scaffold to generate highly active caspase 1 inhibitors NCGC00185682 (5) and ML132 (6, NCGC00183434) (Figure 1A).9

The structural similarity between these agents and Nirmatrelvir certainly allowed for a reasonable anticipation of cross-activity versus caspase 1. Further, previous evaluation of VRT-043198 (4), NCGC00185682 (5), and ML132 (6) across a panel of cysteine proteases highlighted potent activity versus caspases 4, 5, 8, 9, and 14 (Figure 1B). The profile released by Owen et al. included data for only three cysteine proteases (caspase 2, cathepsin B, and cathepsin L). Interestingly, inhibitory activity versus caspase 1 would not necessarily be undesirable. Caspase 1 plays a key role in inflammasome activation and pyroptotic cell death that can be associated with an overactive immune response.10,11 In advanced COVID-19 cases, exacerbated inflammation in the lungs is a significant contributor to respiratory failure and death.12,13 Given Nirmatrelvir’s structural relationship to well-studied caspase 1 inhibitors and its remarkable clinical activity in a setting where dampened inflammation would be therapeutically beneficial, it was of interest to further probe the selectivity of Nirmatrelvir versus a broader panel of human cysteine proteases.

To satisfy this curiosity, we submitted Nirmatrelvir to a commercially available panel14 of 20 human cysteine proteases as well as SARS-CoV-2 MPro and SARS-CoV-2 PLpro. We included NCGC00185682 (5) in this submission as an active control. The results confirmed Nirmatrelvir’s strong activity versus SARS-CoV-2 MPro (IC50 = 4 nM) while demonstrating an extraordinary level of selectivity versus the remainder of the panel with submicromolar activity being noted for only cathepsin K (IC50 = 231 nM) (Figure 1B) (Table S1).

![Figure 1](https://doi.org/10.1021/acsptsci.2c00065)

**Figure 1.**
speculate on the role of the lactam, the 6,6-dimethyl-3-azabicyclo[3.1.0]hexane, or the trifluoroacetamide as structural features that hone the drug’s binding to only SARS-CoV-2 Mpro. What is not left to conjecture is the remarkable skill, effort, and professionalism demonstrated by the Nirmatrelvir team to whom we all owe a debt of gratitude.

**ASSOCIATED CONTENT**

**Supporting Information**
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acspptsci.2c00065.

Profile of Nirmatrelvir (PF-07321332) and NCGC00185682 versus a panel of 23 cysteine proteases (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**
Craig J. Thomas — Division of Preclinical Innovation, National Center for Advancing Translational Sciences, National Institute of Health, Rockville, Maryland 20850, United States; Lymphoid Malignancies Branch, Center for Cancer Research, National Cancer Institute, National Institutes of Health, Bethesda, Maryland 20892, United States; orcid.org/0000-0001-9386-9001; Email: craigt@mail.nih.gov

**Author**
Damien Y. Duveau — Division of Preclinical Innovation, National Center for Advancing Translational Sciences, National Institute of Health, Rockville, Maryland 20850, United States

Complete contact information is available at: https://pubs.acs.org/10.1021/acspptsci.2c00065

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**ABBREVIATIONS**
SARS-CoV-2: severe acute respiratory syndrome coronavirus 2
COVID-19: coronavirus disease 2019
SARS-CoV-2 Mpro: severe acute respiratory syndrome coronavirus 2 main protease
EPIC-HR: evaluation of protease inhibition for COVID-19 in high-risk patients
FDA: U.S. Food and Drug Administration
RTC: replicase-transcriptase complex

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