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Discovery of *N*-(benzo[1,2,3]triazol-1-yl)-*N*-(benzyl)acetamido)phenyl) carboxamides as severe acute respiratory syndrome coronavirus (SARS-CoV) 3CLpro inhibitors: Identification of ML300 and noncovalent nanomolar inhibitors with an induced-fit binding

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ABSTRACT

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Keywords: 3CLpro Severe acute respiratory syndrome SARS MERS Coronavirus Herein we report the discovery and SAR of a novel series of SARS-CoV 3CLpro inhibitors identified through the NIH Molecular Libraries Probe Production Centers Network (MLPCN). In addition to ML188, ML300 represents the second probe declared for 3CLpro from this collaborative effort. The X-ray structure of SARS-CoV 3CLpro bound with a ML300 analog highlights a unique induced-fit reorganization of the S_2-S_4 binding pockets leading to the first sub-micromolar noncovalent 3CLpro inhibitors retaining a single amide bond.

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Coronaviruses (CoV) are enveloped, large plus-strand RNA viruses associated with mild to severe respiratory symptoms, including the common cold and the Severe Acquired Respiratory Syndrome (SARS)-CoV.^{1–3} Identified as the etiological agent responsible for the global pandemic in 2003, SARS presents an atypical pneumonia that during the first major outbreak led to progressive respiratory failure in over 8000 individuals and about 800 deaths by July of that year.⁴ With the cooperation of leading nations, a rigorous public healthcare campaign was fortunately successful in controlling this outbreak. However, a reemergence of the SARS-CoV is considered a potential pandemic risk and new strains of human coronavirus continue to be identified. Since 2003, two additional human coronaviruses, NL63 and HKU1, have been identified in patients and the viruses have been characterized

* Corresponding authors. E-mail address: shaun.stauffer@vanderbilt.edu (S.R. Stauffer). and found to be significantly less lethal than SARS-CoV.^{5–7} Most recently in 2012, a new SARS-like virus, designated the Middle East respiratory syndrome coronavirus (MERS-CoV), has been identified in 144 patients so far, 54 of whom died.⁸ There is now evidence for person-to-person transmission of MERS-CoV.⁹ Now, nearly a decade later, the possibility of another SARS-like pandemic appears even more palpable based upon the lethality and properties of the newly identified MEV-HCoV strain. Effective vaccines and small molecule antiviral agents to prevent or treat SARS-like infections still do not exist, thus tailored antiviral therapies are urgently needed in order to treat potential future outbreaks of SARs and related human coronaviruses.

The SARS and MERS coronaviruses encode two proteases, a papain-like protease (PLpro) and a 3-chymotrypsin-like protease (3CLpro), in their genome that are essential for viral replication. The viral polyprotein is cleaved at three unique sites by PLpro and 11 unique sites by 3CLpro. Initial reports of 3CLpro inhibitors

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in the literature focused on peptidomimetics, often four to five residues in length, bearing a reactive 'warhead' group, such as an aldehyde, halo-methyl ketone, or Michael acceptor at the terminus with several demonstrating a covalent interaction with the active site Cys-145 residue.¹⁰⁻¹⁶ Until recently, the majority of efforts to develop nonpeptidic 3CLpro inhibitors also relied on 'warhead' based design strategies (Fig. 1, 1-5)¹⁷⁻²¹ and a number of these nonpeptidic inhibitors achieved sub-micromolar activity. In the case of pyridyl ester **4**,²⁰ this potent nanomolar mechanism-based enzyme inactivator led to cell based inhibition below 10 µM in SARS-CoV infected Vero E6 cells. Recently, we reported N-(tert-butyl)-2-(N-arylamido)-2-(pyridin-3-yl) acetamide 6 (Fig. 1, ML188) and its X-ray complex with 3CLpro (PDB: 3V3M) as a rare example of a noncovalent SARS-CoV 3CLpro inhibitor of moderate molecular weight with good enzyme and antiviral inhibitory activity.²² Herein, we describe the continuation of efforts to develop potent, noncovalent SARS-3CLpro inhibitors based upon a second chemical class of triazoles from our MLPCN screening campaign (7, Fig. 2) and progression of this lead series to a second generation probe ML300 (16e, Fig. 1) and beyond to arrive at sub-100 nM inhibitors. We propose from crystallography data that ML300 and related triazoles in this series inhibit 3CLpro via a novel mechanism of action and provide a new direction for additional noncovalent inhibitor design and refinement.

Using a designed expression construct which produces the post-proteolytic and authentic 3CLpro dimer, a screen against the NIH molecular libraries sample collection (~293 K compounds) at the Scripps Research Institute Molecular Screening Center (SRIMSC) was undertaken. In addition to the diamide acetamide series represented by ML188 (**6**, Fig. 1),²² a related diamide series, represented by SID 24808289 (**7**, Fig. 2), was identified demonstrating a 3CLpro IC₅₀ of 6.2 µM and good selectivity versus PLpro $(IC_{50} > 60 \,\mu\text{M})$ which is used as a control for cysteine-protease activity. Fortunately, guite early in the chemistry campaign an Xray crystal structure of diamide 7 bound to 3CLpro was determined to 1.85 Å resolution. A solvent accessible surface depiction of 7 in the 3CLpro active site along with a wall-eve stereo view with key contact resides and hydrogen bonding contacts in depicted in Figure 3. Interestingly, in contrast to the ML188-3CLpro crystal structure in which ML188 accommodates substrate sub-pockets in the enzyme active-site traditionally occupied by peptidomimetics, diamide 7 engenders an induced-fit complex resulting in a new surface dictated largely by a rearrangement of the Gln-189 and



Figure 1. Representative nonpeptidic 3CLpro inhibitors utilizing a warhead and noncovalent mechanism of inhibition (1–6).



Figure 2. Binding orientation and properties of MLPCN 3CLpro HTS hit 7 and evolved probe molecule ML300(16e).



Figure 3. (A) Solvent accessible surface view of **7**-3CLpro complex (PDB code: 4MDS, PubChem SID 24808289); (B) X-ray crystal structure of **7** (capped sticks in orange carbon) with SARS 3CLpro in wall-eye stereo view with key residues and hydrogen bonds.

Met-49 residue side-chains.²³ This induced fit accommodates the syn *N*-methyl pyrrole and anilido acetamide moieties of the

inhibitor within subpockets that can be characterized as S_2-S_4 and S_2-S_1' subpockets, respectively.

Figure 2 schematically illustrates the inhibitor-active site interactions oriented in a manner similar as depicted in Figure 3. In addition to the P_2-P_4 and P_2-P_1' groups the inhibitor partially occupies the S_3 subpocket with a terminating 2-methylbutylamide. Key hydrogen bonding interactions can be found near the catalytic site with His-163 and the benzotriazole *N*-(3) engaged in a key interaction, with an interatomic distance of 2.9 Å. In addition a backbone Glu-166 NH interaction is evident with the central acetamide oxygen (N–O distance 2.8 Å).

Flexibility of the diamide scaffold (RotBon ~7) coupled with the observed induced-fit within the active site of 3CLpro presents an added challenge with respect to in silico inhibitor approaches. Thus, our structure–activity-relationship (SAR) studies focused initially on three key areas within the diamide scaffold: (1) benzotriazole replacements with alternative hydrogen bond acceptor functionality to interact with His-163, (2) acetamide modifications within the P₂–P₁' region, and (3) minimum pharmacophore deletion studies of the P₃ 2-methylbutylmide. The P₂–P₄ group was held constant for this investigation and based upon HTS and reconfirmation results (data not shown) the *N*-methyl pyrrole was replaced with an equipotent 3-thienyl moiety.

In parallel with efforts to obtain the 3CLpro-7 crystal structure, synthesis of first generation analogs to survey diversity of the benzotriazole unit were initiated using a modified version of our 4CC-Ugi strategy (Scheme 1) to allow for late stage azole introduction. Thus, Ugi reaction using *t*-butyl isocyanide, chloroacetic acid, thiophene-3-carbaldehyde, and *N*-(4-aminophenyl)acetamide proceeded smoothly to give chloride **8**, which could be isolated in good yield after chromatography. Displacement of chloride **8** with azole NH heterocycles provided **9a–c**. Alternatively, displacement of **8** with sodium azide and subsequent Huisgen cycloaddition reaction with an appropriate acetylene furnished 1,2,3-triazoles **10a–c** in good overall yield.

Synthesis of P_2-P_1' amide analogs within the elaborated diamide were similarly prepared in an Ugi reaction using Boc-protected 4-(amino) aniline (Scheme 2) as the amine component. Deprotection of **11** using trifluoroacetic gave aniline **12** which was reacted with a variety of carboxylic acid derivatives under HATU coupling conditions or reacted with an acid chloride or sufonyl chloride in the presence of TEA to give final examples **13a–1**.

Synthesis of P_3 truncated analogs began with reductive amination using thiophene-3-carbaldehyde with either 4-bromoaniline



Scheme 1. Synthesis of P₁ analogs **9a–c** and **10a–c**. Reagents and conditions: (a) MeOH, 50 °C, 4 h, 95%, (b) (i) NaH, HetNH, DMF, (ii) **9**, DMF, 65–80%, (c) (i) NaN₃, DMF, 100 °C µwave 30 min, 95%, (ii) acetylene (R = Ph, TMS), DCE, 120 °C 16 h, 85–98%, (iii) R = TMS, TBAF, HOAc, 0 °C–rt, 45%. Final library compounds were purified by UV prep or mass-directed prep HPLC.



Scheme 2. Synthesis of P_2-P_1 analogs 12 and 13a–I. Reagents and conditions: (a) MeOH, 50 °C, 4 h, 94%, (b) TFA, 95% (c) (i) HATU, DIPEA, DMF, RCO₂H, 55–73% (d) RCO₂Cl or RSO₂Cl, TEA, DCM, 51–64%. Final library compounds were purified by UV prep or mass-directed prep HPLC.



Scheme 3. Synthesis of P_2-P_1' analogs **16a–k** and **17a–e** within truncated series. Reagents and conditions: (a) NaHB(OAc)₃, DCE, rt, 80% (b) benzotriazol-1-yl-acetic acid, HATU, TEA, DMF, rt, 74% (c) (i) **15a**, TFA, DCM, 95%, (ii) HATU, DIPEA, DMF, RCO₂H, 65–80%; RCO₂Cl or RSO₂Cl, TEA, DCM, 90–95%; NaHB(OAc)₃, RCHO, DCE, 45–95% (d) **15b**, Ar/HetB(OH)₂, 1 M aq Na₂CO₃, 5 mol % Pd(PPh₃)₄, THF, 30–65%. Final library compounds were purified by UV prep or mass-directed pre HPLC.

or Boc-protected 4-(amino) aniline to give intermediates **14a–b** in good yield. Amide coupling with HATU using benzotriazol-1yl-acetic acid installed the requisite P₁ groups to afford **15a–b**. Initial efforts focused on preparing the identical amide library prepared in the elaborated series **13** (Scheme 3). This was readily accomplished as before; Boc-deprotection of **15a** followed by amide coupling or acylation/sulfonylation, gave **16a–k**. Subsequent synthesis of a series of biaryls as amide replacements commenced



Figure 4. Representative azole replacements (9a-c and 10a-c).

using a Suzuki cross-coupling with **15b** and a variety of boronic acids to afford target molecules **17a–e**.

SAR for 1,3-azole P₁ replacements (**9a-c**, Fig. 4) indicated a strict requirement for the 1,2,3-triazole unit; benzimidazoles 9a-b and 2-methyl-1-imidozyl derivative 9c were uniformly inactive. Since the N-(3) nitrogen of 7 appeared to be involved in a hydrogen bond with His-163, it was somewhat surprising that 9a was not tolerated since the imidazole has the potential to maintain a N-(3)-His-163 hydrogen bond interaction. However, within the 3CLpro-7 structure the catalytic Cys-145 residue is located within 3.3 Å of the N-(2) nitrogen, indicating potential for a weak hydrogen bond and/or dipole-dipole stabilization interaction. This potential interaction may perhaps be responsible for the 1,2,3-triazole preference. Interestingly, 4-phenyl 1,2,3-triazole 10c was tolerated with an IC_{50} of 11 μ M, suggesting additional avenues for optimization. Accommodation of the phenyl moiety of **10c** within the active site S₁ subpocket is not entirely clear at this time. Based on the 3CLpro-7 structure, Glu-166, Phe-140, and Glu-166 are predicted to be within close proximity. Unsubstituted triazole 10a and trimethyl silyl triazole 10b were inactive, demonstrating the importance of maintaining a proper aromatic ring in this subpocket.

Amide library 13a-I (Fig. 5) within the elaborated diamide series displayed a range of potency from moderate micromolar activity (13a, 13b-d, 13f-g) comparable to the HTS hit 7, to weakly active or inactive. Cyclic and acyclic acetamide congeners related to HTS lead 7 showed consistent activities below $10 \,\mu M$ with branched *i*-propyl derivative **13d** and cyclobutyl amide **13g** having the greatest activity below 5 µM. Modification to sulfonamide 13b resulted in a three-fold loss in inhibition relative to acetamide 13a. The smaller cyclopropyl (13f) or larger cyclohexyl (13h) cyclic amide generally resulted in loss of inhibition. Incorporating a sterically hindered *t*-butyl amide **13e** also led to a modest three-fold loss in activity. Lastly, aromatic and heteroaromatic amides (13ik) in addition to iso-butyl carbamate 131 were weak or inactive as 3CLpro inhibitors. Collectively, these data appear to be consistent with the 3CLpro-7 structure whereby a short helix-loop-helix motif (Val-42-Ile-43-Cvs-44-Thr-45-Ala-46) and a proximal β-turn

P2-P1' Exploration: amide, urea, sulfonamide ΗN NH_2 12 13a 13b 13c 13d IC₅₀ > 20 μM 7.72 µM IC₅₀ = 25.3 μM IC₅₀ = 6.9 µM IC₅₀ IC₅₀ poor curve fit 13 13f 13g 13h 9.1 μM 3.8 uM 23% inh (100 µM) 22.5 uN ΗŅ ΗN HN ΗN 13i 13j 13k 13 Inactive Inactive 15% inh (100 uM) IC 50 = 26 µM

Figure 5. 3CLpro activity from library 13.

element (Thr-24-Thr-25) define the edge of this pocket with minimal volume for larger groups beyond acetamide **7**.

With limited success from the above S_1 and S_2-S_1' studies we turned to P_3 truncation to potentially identify a minimum pharmacophore to reduce overall MW and improve ligand efficiency (LE).²⁴ Examination of the P_3 group within the **7**-3CLpro structure suggested this group was unfavorably solvent exposed relative to the *t*-butylamide– S_3 interaction found within the ML188-3CLpro structure.²² Initial efforts led to **16a–k** (Table 1). Gratifyingly truncated amides proved to have comparable activity versus the elaborated diamide counterparts (Fig. 5 see **13c–d**, **13f–g**, vs Table 1 see **16a–c**, **16e–f**). Interestingly, truncated series **16** appeared to better tolerate larger substituents, perhaps suggesting additional changes in the shape of the active site within this subpocket. For example cyclohexyl amide **16g** was found to be a weak inhibitor and similarly carbamate **16i** had moderate inhibitory activity of 10.3 µM while its related counterpart **13l** (Fig. 5) was inactive.

At this stage in the project with efforts focused on P₃ truncated analogs bearing a putative S_2-S_1' interaction, we elected **16e** for further characterization and probe declaration (ML300, Fig. 6). Relative to probe ML188 (**6**) and the equipotent diamide **13d**, ML300 proved to offer improvements in several areas (Fig. 6). SARS 3CLpro inhibitor ML300 is ~100 amu lower MW (MW = 431) relative to **13d** with moderate ligand efficiency (LE). Deletion of the lipophilic P₃ group reduces cLogP an order of magnitude (cLogP = 3.2) and thus greatly improves ligand-efficiency-dependent lipophilicity (LELP) versus ML188 and **13d**. Probe molecules ML188 and ML300 were evaluated in an in-house²⁵ in vitro DMPK panel including plasma protein binding, P450 enzyme inhibition, and



3CLpro activity **16a–k**, **17a–e**



Cmpd	R	IC ₅₀ ^a	Cmpd	R	IC ₅₀ ^a
16a	HN	2.9	16i		10.3
16b		3.6	16j	HN ^{CH} 3 	2.1
16c		13.3	16k	HN Ph	1.5
16d		3.4	17a		0.051
16e		4.1	17b		0.97
16f	HN 	8.1	17c	H ₃ CO	0.70
16g	HN 	22.1	17d	N	2.0
16h	0 HN 	18% (100 μM)	17e	$- N \rightarrow OCH_3$	12.5

 $^{\rm a}$ IC_{50} are the average of three independent determinations and represent a co-efficient of variation (CV) < 0.10.



In vitro DMPK and ancillary pharmacology:

 ML188

 PPB Fut (h, r): 3.3%, 2.6%

 CL_{hep} (h, r): 20, 69 mL/min/kg

 CYPs (µM):

 >30 (1A2), 9.1 (2C9), >30 (2D6), 1.7 (3A4)

 Eurofins Profiler Screen:

 No significant activity

ML300 PPBFu (h, r): 4.8%, 10.6% CL_{hep} (h, r): 20, 67 mL/min/kg CYPs (μM): 7.7 (1A2), 7.2 (2C9), 8.4 (2D6), 4.6 (3A4) Eurofins Profiler Screen: Melatonin MT1 75% inhibition (10 μM)

Figure 6. Profiles of 3CLpro inhibitors 6 (ML188), 13d, and 163 (ML300).

intrinsic clearance using liver microsomes (Fig. 6). Both ML188 and ML300 possess good free fraction with ML300 being superior (1.5 and 4.0-fold improved human and rat fraction unbound, respectively); however, intrinsic clearance (CL_{HEP} normalized to liver blood flow, $Q_h = 21 \text{ mL/min/kg}$, $Q_r = 70 \text{ mL/min/kg}$) indicates ML188 and both ML300 are both predicted to be highly cleared. ML188 and ML300 possess modest P450 enzyme inhibition, with ML300 maintaining 5–10 μ M activity across four major CYP enzymes (Fig. 6). Probe ML300 was found to be highly selective in a Eurofins lead-profiling screen²⁶ with only modest activity for melatonin MT₁ receptor in a radioligand binding assay.

Based on the absence of key hydrogen bonding interactions of the P_2-P_1' amide of **7** with the 3CLpro active site, in addition to the poor metabolic instability and CYP profile of ML300, we opted to explore more diverse amide replacements as a means to improve metabolic stability, P450 activity, and 3CLpro inhibitory potency. Initial efforts identified representative N-methyl (16j) and N-benzyl (16k) anilines with potency comparable to probe ML300 (16e). The lack of activity for benzamide 16h versus the reduced benzylamine 16k is striking and indicates the enhanced flexibility of the N-benzyl group is permitting a productive interaction to occur where previously aromatic amides were not tolerated (see Fig. 5, 13i-j). A subsequent survey of biaryls was explored (Table 1, 17a-e) and on the basis of the 3CLpro-7 X-ray, we targeted 3-pyridyl (17b-c) and 4-pyrimidyl (17e) heterocycles as means to potentially engage a side-chain interaction from the hydroxyl groups of Thr-24 or Thr-25. These modifications afforded inhibitors with micromolar activity and in the case of 2-methoxypyridyl **17c** submicromolar activity (IC₅₀ = 700 nM). Unexpectedly the parent simple phenyl biaryl 17a proved to have a major impact on activity with a ~7-10-fold increase relative to 17b-c. 3CLpro inhibitor 17a represented the first sub-100 nM inhibitor for the series and to our knowledge one of the most potent nonwarhead based SARS 3CLpro inhibitors to date. At this time inhibitor 17a is relatively unoptimized and thus current efforts are focused on targeted biaryl congeners to understand DMPK, cellular activity, as well as potential broad spectrum activity against other coronavirus strains including MERS-CoV.

In summary, we have described the identification and binding orientation and interactions for a second class of diamide SARS 3CLpro inhibitors, culminating in probe ML300 and subsequently improved inhibitors such as **17a**, which possess LE > 0.3 and an LELP approaching 10. The X-ray crystal structure of HTS hit **7** bound to 3CLpro²⁷ was instrumental in guiding optimization and

the induced-fit of this inhibitor 3CLpro complex illustrates the challenges of divergent SAR and the limitations of virtual based screens. The four component Ugi reaction was utilized once more to rapidly generate SAR for the putative P_2-P_1' and P_1 subgroups. Importantly, P_3 truncation was possible for this triazole series of 3CLpro inhibitors, allowing for significant MW reduction without diminishing potency. Collaborative efforts in these laboratories continue towards the identification active inhibitors within the truncated biaryl class. Integrated efforts with DMPK assessment continue in order to improve intrinsic clearance and diminish P450 activity, which are issues to be addressed within the series prior to in vivo proof-of-mechanism studies. ML300 is an MLPCN probe and is freely available upon request.

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- 26. For information on MLPCN's probe compound ancillary screen see Eurofins
- LeadProfilingScreen[®]: www.eurofinspanlabs.com.
 27. The protein-ligand X-ray structure of 7-bound SARS-3CLpro has been deposited in PDB. RCSB ID code rcsb081783 and PDB ID code 4MDS.