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Permalink https://escholarship.org/uc/item/5z17x94z

Journal Journal of the American Chemical Society, 145(18)

ISSN

0002-7863

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Publication Date

2023-05-10

DOI

10.1021/jacs.2c12240

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Engaging a Non-catalytic Cysteine Residue Drives Potent and Selective Inhibition of Caspase-6

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catalytic cysteine residue (C264) unique to caspase-6 (C6), an enigmatic and understudied caspase isoform. Starting from disulfide ligands identified in a cysteine trapping screen, we used a structureinformed covalent ligand design to produce potent, irreversible inhibitors (3a) and chemoproteomic probes (13-t) of C6 that exhibit unprecedented selectivity over other caspase family members and high proteome selectivity. This approach and the



new tools described will enable rigorous interrogation of the role of caspase-6 in developmental biology and in inflammatory and neurodegenerative diseases.

INTRODUCTION

Caspases are cysteine-dependent aspartyl-specific proteases involved in a range of cellular and disease processes, ranging from apoptosis to inflammation and neurodegeneration.¹⁻ Caspase family members are broadly classified as inflammatory (C1, C4, and C5) or apoptotic caspases. The apoptotic caspases include the so-called initiator caspases (C8 and C9) that cleave executioner caspases (C3, C6, and C7) which, thus activated, go on to cleave hundreds of proteins⁶ as part of the apoptotic cell death program. The importance of caspases in various cellular processes and disease states emphasizes the potential value of isoform-selective small-molecule inhibitors and probes of caspase activity. Historically, small-molecule probes (e.g., Ac-VEID-CHO) and drug molecules (e.g., emricasan) targeting caspases have been structural mimics of caspase substrates bearing an electrophilic functionality (e.g., aldehydes or aryloxymethylketones) to engage the catalytic cysteine residue (Figure 1). Although such tools are widely available and frequently used in biomedical research, their lack of significant isoform selectivity, particularly in cellular contexts, confounds the biological or pharmacological conclusions inferred from their use.⁷ Moreover, electrophilic substrate mimics have proved challenging to develop clinically,^{8,9} likely due to insufficient selectivity and/or suboptimal drug-like properties.

While often classified as an executioner caspase, the function of caspase-6 (C6) in physiology and disease has remained enigmatic, and it has been suggested that C6 may act upstream of the other executioner caspases.^{4,10} In addition to a putative amplifying role during apoptosis, C6 has been found to also possess nonapoptotic functions important for axon pruning and neuroinflammation.^{11–14} Finally, C6-mediated proteolysis has been implicated in the pathology of Huntington's^{15,16} and Alzheimer's disease,^{17–19} in neuroinflammation generally^{20–22} and in nonalcoholic steatohepatitis.¹⁴

There is a clear need for potent and selective C6 inhibitors and activity-based probes to validate and exploit the role of C6 in these disease states. To produce the first truly selective probe of C6 activity in cellular contexts, we leveraged a nonconserved cysteine residue (C264) that lies on a loop at the distal end of the active site (Figure 2). While a non-catalytic cysteine is required for C8 activity²³ and regulates the response of C9 to oxidative stress,²⁴ no endogenous role for C264 in C6 regulation is known, nor has C264 to our knowledge been

Received: November 18, 2022 Published: April 27, 2023





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Figure 1. Examples of substrate-like tool compounds (left) and drug candidates (right) that engage the catalytic cysteine of caspases.



Figure 2. Summary of a targeted approach to the discovery of potent and selective caspase-6 inhibitors and chemical probes.

Scheme 1. Synthesis of Enantiomers 2a and 2b



Table 1. Measures of C264 Engagement, Biochemical IC₅₀ Values, and Inhibition Kinetics for Caspase-6 Inhibitors

F ₃ CO			Ar_NH H O ^S =0	Ar_NHUNCH	Ar, N, H, N,	Ar N H Ph A	N H Ph
	2a (3 <i>R</i>)	2b (3S)	3a (3 <i>R, 5</i> S)	3b (3 <i>S</i> , 5 <i>R</i>)	(±)- 3-c (cis)	(±)- 3-t (trans)	(±)- 10- <i>t</i> (<i>trans</i>)
Casp-6 IC ₅₀	0.19 µM	>100 µM	0.0081 µM	4.2 μΜ	> 100 µM	0.052 μM	>100 µM
SE of IC ₅₀	0.020 µM	N/A	0.0004 µM	1.2 µM	N/A	0.0017 µM	N/A
$C6^{sm} DR_{50}^{a}$	0.08 µM	13 µM	-	-	-	-	-
Ki	-	-	7.0 µM	-	-	12.5 μM	-
k _{inact}	-	-	3.1 x 10 ⁻² s ⁻¹	-	-	$1.67 \ge 10^{-2} \text{ s}^{-1}$	-
$k_{\rm inact}/K_{ m i}$	-	-	$4400 M^{-1} s^{-1}$	-	-	1,330 M ⁻¹ s ⁻¹	-
Casp-3 IC ₅₀	>500 µM	-	> 500 µM	-	>500 µM	>500 µM	-

^{*a*}DR₅₀ is the concentration of the inhibitor, producing 50% labeling of the Casp-6 small subunit (C6sm), as determined by MS.

leveraged for inhibitor discovery previously. To achieve this, we employed disulfide trapping (tethering²⁵), a mass spectrometry-based screening approach, to identify covalent probe ligands for surface-exposed/reactive cysteines in diverse proteins. The application of tethering to caspases was pioneered by Hardy and Wells,^{26,27} who showed that endogenous cysteine residues at the dimer interface of C3 and C7 could be trapped as disulfides with small molecule thiols to stabilize an inactive, zymogen-like conformation, thereby inhibiting the protease in biochemical assays. While these early thiol-based probes were not effective in the

reducing environment of cells, the work inspired subsequent efforts by us and others, leading to cell-active²⁸ probes of proC6 and proC8.^{16,29}

RESULTS AND DISCUSSION

C6 possesses a non-catalytic cysteine (C264) not present in other family members and is situated on a loop in the small subunit at the distal end of the active site (Figure 2 and supplementary Figure S1). The catalytic cysteine (C163) is found on the large subunit, while two additional cysteine residues appear to be buried and thus inaccessible for disulfide

formation. A disulfide tethering screen of C6 was performed employing our synthetic library of ~1,500 diverse disulfidebearing small molecules.^{30,31} In this technique³² proteindisulfide conjugates formed under reducing conditions are detected by automated HPLC/MS (supplementary Figure S2A). Notably, a large majority of screening hits labeled the small subunit, suggesting a preference for C264 over the catalytic cysteine C163 on the large subunit (supplementary Figure S2B). Moreover, labeling of the small subunit in MS experiments correlated with inhibition of C6 in biochemical assays (supplementary Figure S2C). The best C264-binding thiols contained a substituted piperidine scaffold with either a 3-amido (derived from nipecotic acid) or 4-amido linkage to the disulfide. The distal substituents varied, including substituted phenyls, 5,6-fused rings, and 6,6-fused rings (supplementary Figure S2D).

Among the thiol-bearing hits identified in the screen was (S)-nipecotic acid-derived analogue 1 bearing a distal paratrifluoromethoxy substituted aryl ring (Figure 2). As with the other hits identified in the screen, compound 1 labeled the small subunit almost exclusively. To derive a cell-active probe, we sought to replace the thiol side chain with a suitably positioned, electrophilic warhead. One successful foray in this vein produced analogue 2a (R = p-OCF₃), in which the nipecotic acid core was reoriented, with the piperidine nitrogen displaying a vinylsulfonamide. Analogue 2a, having the 3R configuration, exhibited markedly superior C264 engagement compared to its 3S enantiomer 2b, with a mass spectrometrybased dose-response-50 (DR₅₀) value of 0.08 μ M and a C6 inhibition IC₅₀ value of 0.180 μ M (Scheme 1 and Table 1), indicating that inhibition by 2a involved both molecular recognition and reaction with C264. Notably, among several cysteine-reactive warheads explored, the vinylsulfonamide alone provided potent C6 inhibition. Also fortuitously, the para-trifluoromethoxyaryl ring present in the original screening hit 1 proved superior to many differently substituted aryl rings explored and was thus retained during further optimization.

The crystal structure of **2a** bound to C6 confirmed the covalent modification of residue C264, whose α -carbon atom moved ~2.4 Å toward the substrate-binding groove, compared to the structure of C6 bound to a VEID peptide substrate analogue (Figure 3). Informed by this structure, we explored new piperidine substitutions, seeking to engage additional



Figure 3. Structures of 2a, 3a, and Ac-VEID-CHO bound to caspase-6. (A) Superposition of crystal structures of 2a (cyan) and 3a (green) bound in the active site of caspase-6. The catalytic cysteine C163 is on the large subunit (at right), while inhibitor-modified C264 is on the small subunit (at left). (B) Superposition of 3ou for your a (green, light green protein) and substrate analogue Ac-VEID-CHO (violet, light violet protein). The side chain of R220 moves 4.2 Å to accommodate the trifluoromethoxyphenyl substituent on 3a. PDBIDs: 8EG6 (2a/C6); 8EG5 (3a/C6); 3OD5 (VEID/C6).

pockets in the active site. Initial modeling suggested this might be achieved with a bulky C-5 substituent bearing a *cis* relationship to the benzamide side chain at C-3. Contrary to expectations, however, it was in fact the *trans* diastereomer (\pm) -3-*t* that realized potent C6 inhibition, while (\pm) -3-*c* was surprisingly without detectable biochemical activity (Table 1).

To assign the eutomer and distomer of (\pm) -3-t, we separated the enantiomers of the synthetic precursor (\pm) -8-t (Scheme 2) and from these prepared analogues 3a (3R, 5S) and 3b (3S, 5R). When tested in a biochemical assay, enantiomer 3a potently inhibited C6 activity with an IC₅₀ value of 8 nM, while its enantiomer **3b** exhibited an IC_{50} at least 100fold weaker (Table 1). A complex X-ray crystal structure of 3a showed similarities and differences as compared to the binding of 2a (Figure 3A). Both analogues engaged C264 in a covalent bond as expected, while the trifluoromethoxyaryl side chain of both analogues was buried in the S1 pocket, where the aspartic acid residue of the C6 substrate is usually bound (Figure 3A). To accommodate this large and hydrophobic ring system in S1, arginine 220, which formed hydrogen bonds with the aspartic acid in VEID peptide-type inhibitors, is moved by 4.2 Å (ε nitrogen) to create a hydrophobic cavity complementary in shape to the sidechain of 3a (Figure 3B). The accessibility of this pocket could explain the higher hit-rate for the smallsubunit vs the active-site cysteine in the primary screen. However, the location and orientation of the piperidine ring and sulfonamide function were quite different in the two analogues. The phenyl substituent present in 3a made few, if any, contacts with the protein, but by occupying an equatorial position in the bound conformation, it energetically stabilized the required axial orientation of the trifluoromethoxy benzamide side chain of 3a, as bound in the S1 pocket.

To evaluate caspase isoform selectivity, we tested 3a for the inhibition of recombinant C2, C3, and C6 WT and the C264A mutant protein in a biochemical activity assay in the presence of the fluorogenic substrate (Figure 4A). The potent low-nM inhibition of C6 WT protein by 3a was completely lost with the C264A mutant, further establishing that 3a engages C264. Also consistent with this requirement was the lack of activity for 3a against either C2 or C3 (Figure 4A). Next, human caspases C1–C10 were challenged with 3a at 10 μ M (>100fold its C6 IC_{50}). In this experiment, **3a** had little, if any, effect on any of the other isoforms (Figure 4B). The potency and exquisite isoform selectivity of 3a, together with the lack of C6 activity for its enantiomer 3b, strongly suggested that 3a inhibits C6 by first forming a pre-covalent complex that positioned the electrophilic vinylsulfonamide for subsequent reaction with C264. The acrylamide 10-t, a direct congener of 3-t (Scheme 2), lacked any measurable activity against C6, further supporting the notion that inhibition was highly sensitive to proper electrophile placement and orientation.

To better understand C6 inhibition by **3a**, we determined kinetic parameters (k_{inact}/K_i values) of inhibition (supplementary Figure S3). These data indicated that indeed, both noncovalent and covalent interactions contributed to the binding, with a K_i value of 7.0 μ M, providing an estimate of the intrinsic pre-covalent binding affinity of **3a** for C6. Moving to cellular studies, **2a** and **3a** were evaluated for their ability to inhibit cleavage of the C6-specific substrate lamin A.³³ This assay measured lamin A cleavage under staurosporine-induced caspase activation; while staurosporine activates all executioner caspases, lamin A cleavage monitors only the activity of C6. After 1 h of preincubation with the compound and 3 h of

Scheme 2. Synthesis of (\pm) -3-t and (\pm) -3-c, and the Corresponding Non-Racemic Stereoisomers 3a-d^a



^{*a*}Compounds 3a-d were prepared from the individual stereoisomers 9a-d, which were separated by silica gel chromatography, followed by chiral-SFC. The acrylamide analogue (\pm)-10-*t* was synthesized from (\pm)-8-*t*, as shown at lower right



Figure 4. Biochemical activities of **3a**. (A) **3a** inhibits wild-type (WT) caspase-6 (C6) but not C264A C6, caspase-2 (C2), or caspase-3 (C3). Error bars represent the standard deviation of thre replicate measurements. (B) Inhibition at 10 μ M **3a** of ten human caspases (C1–C10). Activity measured by the cleavage of aminofluorocoumarin-labeled peptides (see the Supporting Information for sequences). C = caspase.

incubation with staurosporine, lamin A cleavage was measured by in-cell western. Lamin A cleavage was inhibited by **2a** with an IC₅₀ value of 14.2 \pm 2.4 μ M and by **3a** with an IC₅₀ value of 170 \pm 4 nM (Figure 5). Importantly, **2a** and **3a** inhibited lamin A cleavage in a manner that tracked well with their biochemical potencies, with **2a** roughly 50-fold less potent than **3a**. These data suggested that **2a** and **3a** function as C6-selective inhibitors in complex cellular environments containing glutathione and other cellular reductants, as well as numerous cysteine-bearing proteins.

Having established that 3-t/3a exhibited caspase isoform selectivity and is active in cells, we sought to evaluate selectivity across the cellular proteome. We synthesized (\pm) -trans-13 (13-t), a direct congener of 3-t bearing a propargyl ether substituent on the aryl side chain (Scheme 3). After confirming that 13-t displayed similar potency to 3-t (IC₅₀ = 48 ± 2 nM, supplementary Figure S4), we evaluated the proteome-wide selectivity of this analogue in HEK293 cells using stable isotope labeling by amino acids in cell culture (SILAC) and quantitative liquid chromatography/mass spectrometry (LC/MS). Since active C6 levels were below our limit of detection for LC/MS in several cell lines (data not



Figure 5. Cell-based activity of **2a** and **3a**. Inhibition of lamin A cleavage in SK-N-AS cells after staurosporine induction of caspases. Cells were treated with the indicated compound for 1 h before the addition of 10 μ M staurosporine for 4 h. Error bars indicate the standard deviation from n = 4 replicate samples.

Scheme 3. Synthesis of (\pm) -13-*t*, a Prototype Activity-Based Probe of Caspase-6, for Cellular and Chemoproteomic Studies



shown), we instead transfected HEK293 cells with an autoactivating form of caspase-6. After 24 h of transfection, 1 μ M probe 13-*t* was added to cells for 60 min, with or without 60 min of pre-treatment with 10 μ M 3a as the competitor. Cells were lysed, and heavy and light SILAC lysates were



Figure 6. Targets of (\pm) -13-*t* $(1 \mu M)$ in HEK293 cells transfected with caspase-6. (A) In gel fluorescence showing that a protein the size of the caspase-6 small subunit is a primary target of 13-*t*, and this target is competed off by the presence of 3a. (B) Volcano plot showing 336 total proteins identified, 141 of which were statistically significant (p < 0.05) targets of 13-*t* that were competed for binding with 3a. Red circles, competition > log 2 and *p* value <0.05; blue diamond, a small subunit of caspase-6. Keratin proteins are not highlighted because these proteins are likely present due to contamination. Data shown are representative of two independent experiments.

combined in a 1:1 ratio and subjected to CuAAC reaction with azido-biotin to biotinylate probe-labeled proteins for subsequent capture on streptavidin-coated beads. Bead-bound proteins were split for analysis by in-gel fluorescence and tryptic digestion on-bead for LC–MS/MS analysis. The in-gel fluorescence scan showed a dominant band around 15 kDa, consistent with the size of the caspase-6 small subunit that contains C264, the labeling of which was robustly competed by **3a** co-treatment (Figure 6A). There were also several weakly labeled bands that, in contrast to the 15 kDa band, were not significantly competed by **3a** and likely reflected weak non-specific interaction with **13-t** and/or with the surface of the streptavidin beads.

The list of LC/MS-enriched protein targets from 13-t/3atreated cells was compared to that from cells not treated with 13-t or 3a. The data revealed more than 25 proteins that were significantly (p < 0.05) enriched by more than log 2 fold from the 13-t pulldown. The small subunit of caspase-6 was the second most highly enriched and significant protein identified in the 13-t pulldown (supplementary Figure S5). Of the 25 significant interactors, only eight proteins were significantly enriched in the presence of 13-*t* (enrichment \geq twofold) while also being competed for by 3a, suggesting saturable labeling by 13-*t* (Figure 6B). Notably, the caspase-6 small subunit was the most significantly 13-t enriched and 3a-competed protein target. None of the remaining seven putative off-target proteins were proteases; all are membrane-associated proteins, including two heme proteins (CYB5B and HMOX2) and one cysteine-dependent enzyme (PTGES2). These proteomic data indicate that 3a exhibits meaningful, proteome-wide selectivity for caspase-6, in addition to its exquisite caspase isoformselective inhibition.

CONCLUSIONS

Here, we used disulfide tethering and structure-based design to produce caspase-6 inhibitors and activity-based probes of unprecedented isoform selectivity and useful proteome-wide selectivity. Compared to peptidic electrophiles currently in use, compounds **3a** and **13**-*t* much better reflect the high bar set for chemical probes⁷ useful in biomedical research. Beyond their utility as probes, the compounds described herein represent suitable starting points for the development of potent and isoform-selective caspase-6 inhibitors as therapeutic leads. Efforts in this direction are underway and will be reported in due course.

ASSOCIATED CONTENT

9 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c12240.

A full list of proteomic targets of 13-*t* (XLSX)

Additional experimental details, materials, and methods, including scans of ¹H NMR spectra for 3a-d and chromatograms for the separation of 9a and 9b, list of caspase reporter peptides, crystallography data, conservation of C264, disulfide tethering screen, Ki/kinact measurement of 3a, caspase-6 biochemical IC₅₀ data, and SILAC quantitative proteomics (PDF)

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Author Contributions

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Notes

The authors declare the following competing financial interest(s): M.R.A. and A.R.R. are cofounders and advisors to Elgia Therapeutics.

ACKNOWLEDGMENTS

We gratefully acknowledge Jie Liu, Yanli Wang, Yanlong Zhao, and Panpan Fan (Viva Biosciences) for re-collection and submission of the structures 8EG5 and 8EG6 to the protein data bank. We thank Jamie Byrne for mass spectrometry analysis and Sarah Lively and Jeff Neitz for discussions. This research was supported by CHDI (M.R.A. and A.R.R.), the Alzheimer's Association DVT-14-322219 (M.R.A. and A.R.R.), and the National Institutes of Health 1F32AG072822-01 (A.J.A.).

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