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An activation-based high throughput screen identifies caspase-10 inhibitors†

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Caspases are a family of highly homologous cysteine proteases that play critical roles in inflammation and apoptosis. Small molecule inhibitors are useful tools for studying caspase biology, complementary to genetic approaches. However, achieving inhibitor selectivity for individual members of this highly homologous enzyme family remains a major challenge in developing such tool compounds. Prior studies have revealed that one strategy to tackle this selectivity gap is to target the precursor or zymogen forms of individual caspases, which share reduced structural homology when compared to active proteases. To establish a screening assay that favors the discovery of zymogen-directed caspase-10 selective inhibitors, we engineered a low-background and high-activity tobacco etch virus (TEV)-activated caspase-10 protein. We then subjected this turn-on protease to a high-throughput screen of approximately 100 000 compounds, with an average Z' value of 0.58 across all plates analyzed. Counter screening, including against TEV protease, delineated *bona fide* pro-caspase-10 inhibitors. Confirmatory studies identified a class of thiadiazine-containing compounds that undergo isomerization and oxidation to generate cysteine-reactive compounds with caspase-10 inhibitory activity. In parallel, mode-of-action studies revealed that pifithrin- μ (PFT μ), a reported TP53 inhibitor, also functions as a promiscuous caspase inhibitor. Both inhibitor classes showed preferential zymogen inhibition. Given the generalized utility of activation assays, we expect our screening platform to have widespread applications in identifying state-specific protease inhibitors.

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Introduction

Human caspases are a family of 12 cysteine proteases, which are canonically associated with cell death,^{1–3} but have also been tied to nearly all cellular processes, spanning activation,^{4,5} proliferation,^{6,7} differentiation,^{8,9} and cell migration.¹⁰ Therefore, pharmacological

manipulation of individual caspases represents an exciting opportunity to delineate caspase-specific functions and to intervene in human pathologies linked to dysregulated caspase activity, including metabolic¹¹ and immune disorders,^{12–15} cancers,^{16–20} and neurodegenerative diseases.²¹ Selective inhibitors are available for caspase-1,²² caspase-2,^{23,24} caspase-6,^{25,26} and caspase-8.^{27,28} Despite these advances, obtaining highly selective inhibitors remains challenging, likely due in large part to the high sequence and structural homology of all caspases.

Caspase-10 is one such caspase that, while particularly intriguing from a target perspective due to its important functions in immune cell apoptosis, still lacks selective inhibitors. In fact, caspase-10 is one of the only caspases that is not labeled by many conventional peptide-based caspase inhibitors.^{29,30} Caspase-10 is absent in rodents³¹ and shares high sequence homology with caspase-8. Efforts to delineate caspase-10's role in initiating apoptosis have been complicated by seemingly contradictory findings in immortalized cancer cells *versus* primary T cells.^{32,33} In cancer cells, caspase-8 rather than caspase-10 is the primary driver of extrinsic apoptosis.^{32–37} Caspase-10 has even been implicated as a dominant negative regulator of programmed cell death in some cancer types.^{1,38}

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In contrast, in primary T cells, caspase-10 contributes to the initiation of apoptosis;¹³ humans harboring inactivating mutations in caspase-10 experience autoimmunity and excessive T cell proliferation^{39,40} because of decreased apoptosis. These human phenotypes indicate non-functionally redundant roles for caspase-8 and -10 and highlight the likely value of selective inhibitors targeting each caspase, both for further characterizing the unique and overlapping activities of each protease and, more broadly, towards the production of new chemical tool compounds to manipulate adaptive immune cell function.

Type II kinase inhibitors, which target the inactive form of enzymes, exemplify one strategy to improve inhibitor selectivity.^{41–44} We took this approach in our prior work, which identified selective caspase-8 and caspase-2 inhibitors that function by targeting the zymogen, or uncleaved and inactive, precursor caspase proteoforms.^{24,27,45} However, in our previous study, developing selective inhibitors for caspase-10 that did not cross-react with caspase-8 proved elusive despite our best efforts.²⁷ Consequently, there is an unmet need for new approaches to caspase-10 inhibitor discovery.

Here, we develop and apply a tobacco etch virus (TEV) activation-based screening platform to discover pro-caspase-10 inhibitors. To build this platform, we first generated an engineered caspase-10 protein (proCASP10TEV Linker) in which the caspase cleavage sites were replaced with TEV recognition sequences. This engineered protease showed low background, high stability, and robust TEV-dependent activity. After TEV activation, proCASP10TEV Linker protease showed comparable activity to recombinant active caspase-10. Enabled by this assay, we conducted a ~100 000 compound screen, which had a hit rate of ~0.22% (calculated as compounds with a Z-score less than -3) and an average Z'-factor (a measure of how well positive and negative controls are separated)⁴⁶ of 0.58 (Table S1, ESI†). Our subsequent re-screening and counter-screening efforts validated hits and delineated TEV inhibitors from those targeting proCASP10. Resynthesis of a thiadiazine-containing hit compound (**SO265**) unexpectedly revealed that compound rearrangement was driving the observed caspase-10 inhibition. Additionally, mode-of-action studies revealed that the hit compound pifithrin- μ (**PFT μ**), a previously reported inhibitor of TP53,⁴⁷ is also a broadly reactive caspase inhibitor. Taken together, we expect that our hit compounds and innovative screening platform will advance efforts to discover potent and selective pro-caspase-10 inhibitors.

Establishing a TEV-protease activatable caspase-10

Guided by previous reports of TEV-activatable caspases,^{24,48,49} our first step was to engineer a TEV-activatable caspase-10 protein (Fig. 1(A)). For our first attempt at engineering a TEV-cleavable construct (proCASP10TEV), we replaced the aspartate cleavage site, D415, with the TEV recognition site (Fig. S1; ESI† D415ENLYFQG). While this protein did show substantial TEV-dependent increased activity, a high background in the absence of TEV protease was also observed (Fig. 1(B) and Fig. S2, ESI†). We ascribed this background activity to the formation of activated protease in our purified protein sample (Fig. S3, ESI†).

The presence of TEV-independent protease activity, and therefore, cleaved caspase, while somewhat varied between recombinant protein batches, was consistently observed in our purified protein after labeling with **Rho-DEVD-AOMK** (Fig. S3, ESI†). The background activity was also observed to increase markedly over time, even at nanomolar protein concentrations and with increasing concentrations of the sodium citrate kosmotrope (Fig. S4, ESI†). Therefore, we deemed this construct incompatible with HTS, given the absolute requirement for a highly stable and consistently performing protein for large-scale screening applications.

While caspase-10 is not known to harbor additional caspase cleavage sites, we postulated that D435 might also be recognized and cleaved, given the proximity to a likely caspase-recognition motif (PAED). Therefore, we inserted a second TEV motif to generate the proCASP2xTEV protein (Fig. S1, ESI†). Disappointingly, this enzyme showed low overall activity and negligible TEV-dependent activation, suggesting that sequence alterations at D435 are not well tolerated (Fig. 1(B)).

While caspases show high selectivity for aspartyl residues, we postulated that the glutamate residue in the TEV recognition sequence could be recognized and cleaved at a low level, thereby rationalizing the observed high TEV-independent background for the proCASP10TEV protein. To test this hypothesis, we next generated a caspase-10 construct (proCASP10TEV Linker) in which a two-alanine spacer was included to reposition this glutamate further from the remainder of the caspase recognition motif. We additionally optimized our expression and purification protocol to reduce self-activation, both by decreasing the induction time and by not freezing the cell pellets prior to purification (see methods). Pleasingly, the proCASP10TEV Linker protein exhibited both higher overall activity and reduced background compared to our initial construct (Fig. 1(B)). Further supporting the improved behavior of this protein, we detected no formation of cleaved caspase (~20 kDa) by gel-based analysis (Fig. S5, ESI†).

Assessing the suitability of proCASP10TEV Linker for high throughput screening (HTS)

To test whether the proCASP10TEV Linker protein would faithfully recapitulate proCASP10 activity and thus prove suitable for HTS, we further assessed the activity of this protein. Gratifyingly, upon the addition of TEV protease, we observed marked TEV-dependent increased catalytic activity (Fig. 1(B) and (C)). TEV activation occurred rapidly and required only a modest (~500 nM) concentration of TEV protease to achieve complete conversion of proCASP10TEV Linker (333 nM) to the cleaved proteoform (Fig. S6 and S7, ESI†). The proCASP10TEV Linker protein, once activated by TEV protease, showed near-comparable activity to activeCASP10 (Fig. 1(C)), with only a modest decrease in k_{cat} and K_m (Fig. 1(D)). We find that the protein labels robustly with our previously reported caspase-8/10 click probe **KB61** (Fig. S6, ESI†),²⁷ which corroborates that the proCASP10TEV Linker protein behaves similarly to proCASP10. The TEV cleavable construct was also inhibited by the dual caspase-8 and caspase-10 inhibitor **KB7** (10 μ M) and was not inhibited by the structurally matched inactive control, **KB62** (10 μ M) (Fig. 1(E)).²⁷ These findings



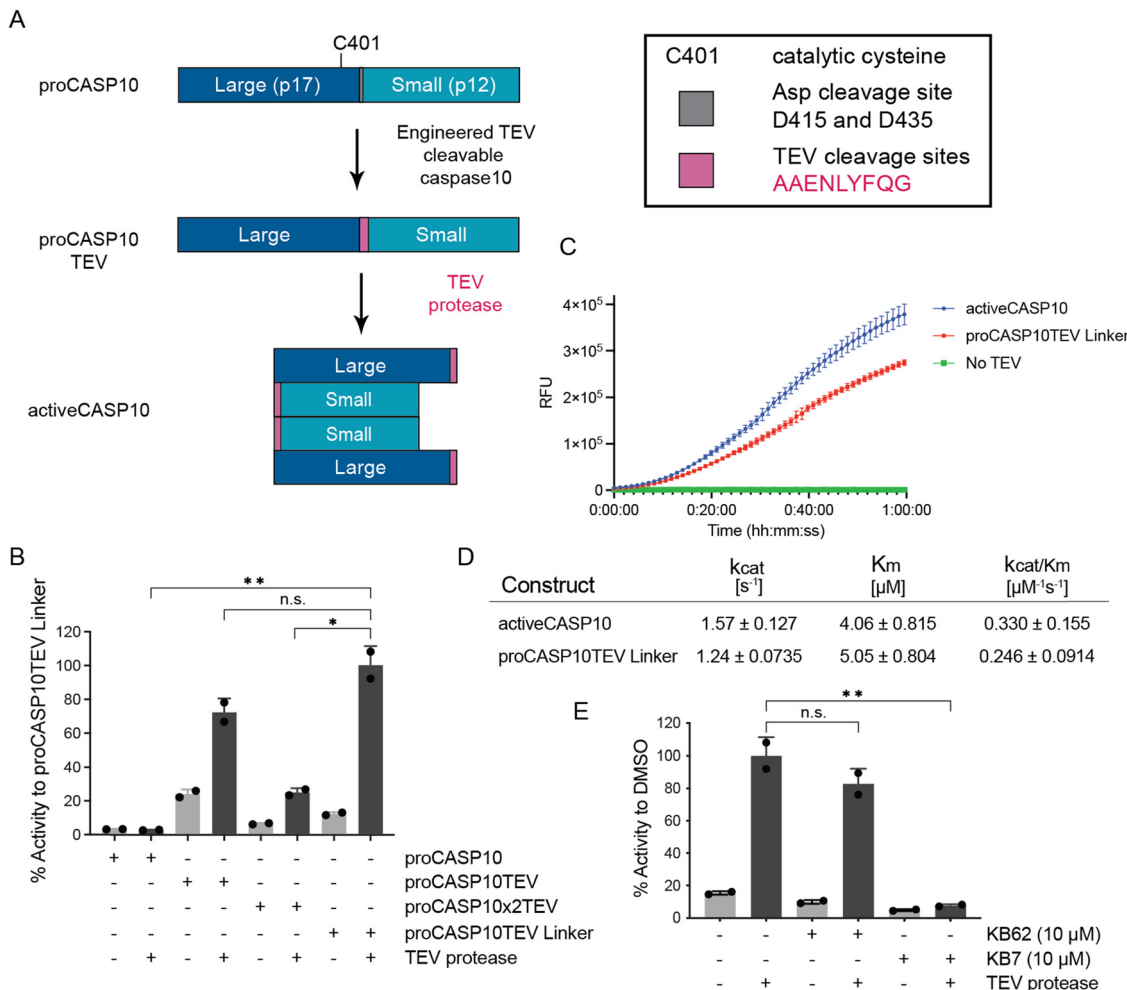


Fig. 1 Robust TEV-cleavable functional caspase-10 is only cleaved in the presence of TEV protease and can be inhibited with a dual caspase-8/-10 inhibitor (KB7).²⁷ (A) Design of engineered TEV cleavable caspase-10 containing TEV recognition site, ENLYFQG (pink). (B) Comparison of relative TEV-dependent and TEV-independent caspase activity for the indicated engineered proteins (333 nM) to cleave Ac-VDVAD-AFC fluorogenic substrate (10 μM) with/without TEV protease (667 nM). Sequences of engineered proteins can be found in Fig. S1 (ESI†). (C) Time course comparing activeCASP10 (blue) versus proCASP10TEV Linker with (red) and without (green) TEV protease (667 nM) in the presence of Ac-VDVAD-AFC fluorogenic substrate (10 μM). (D) Michaelis–Menten kinetics comparing proCASP10TEV Linker (333 nM) activity to activeCASP10 (333 nM) activity as measured by cleavage of the fluorogenic substrate over time. (E) Assessing the relative inhibition of proCASP10 Linker protein by dual pro caspase-8/-10 inhibitor KB7 (10 μM) and negative control compound KB62 (10 μM).²⁷ Assay conducted as in 'B'. For B and E, data represents mean value ± standard deviation for two biological replicates. For C and D, data represent mean values ± standard deviation for three biological replicates. Statistical significance was calculated with unpaired Student's *t*-tests, **p* < 0.05, ***p* < 0.001, ns, not significant *p* > 0.05. Relative fluorescence units, RFU.

provided further evidence that our engineered protein behaves similarly to proCASP10 and, additionally, demonstrated that the protein is well suited to assess pro caspase inhibition.

We next vetted our assay conditions for HTS compatibility, with the goal of ensuring both high activity and high stability over longer assay periods. Additionally, we found that the proCASP10-TEV Linker protein showed increased activity in the presence of the kosmotrope sodium citrate at increasing concentrations (37 mM, 111 mM, and 333 mM) (Fig. S8, ESI†). This added activity was restricted to the TEV-cleaved/activated protein; little TEV-independent activity was observed after chaotrope addition in the absence of TEV protease. Providing evidence of high enzyme stability, a favorable property for screening applications, proCASP10TEV Linker showed no substantial change in enzyme

activity after prolonged (18 h) incubations times for both 4 °C and ambient temperature (Fig. S9, ESI†). Additionally, substrate turnover was observed to progress in a linear manner for the initial ~2 h period, with increasing activity observed up to 6 h (Fig. S10, ESI†), which provides a wide time window for acquiring data. Thus, with high stability, low background, and high activity enzyme in hand, we turned to HTS implementation.

A small-scale screen of two pharmacologically active library compounds confirms assay compatibility with HTS

Toward large-scale library screening, our next step was to validate our TEV-mediated caspase activation assay and establish a miniaturized (384-well plate) semi-automated workflow. Following the workflow shown in Fig. 2(A), we first screened the



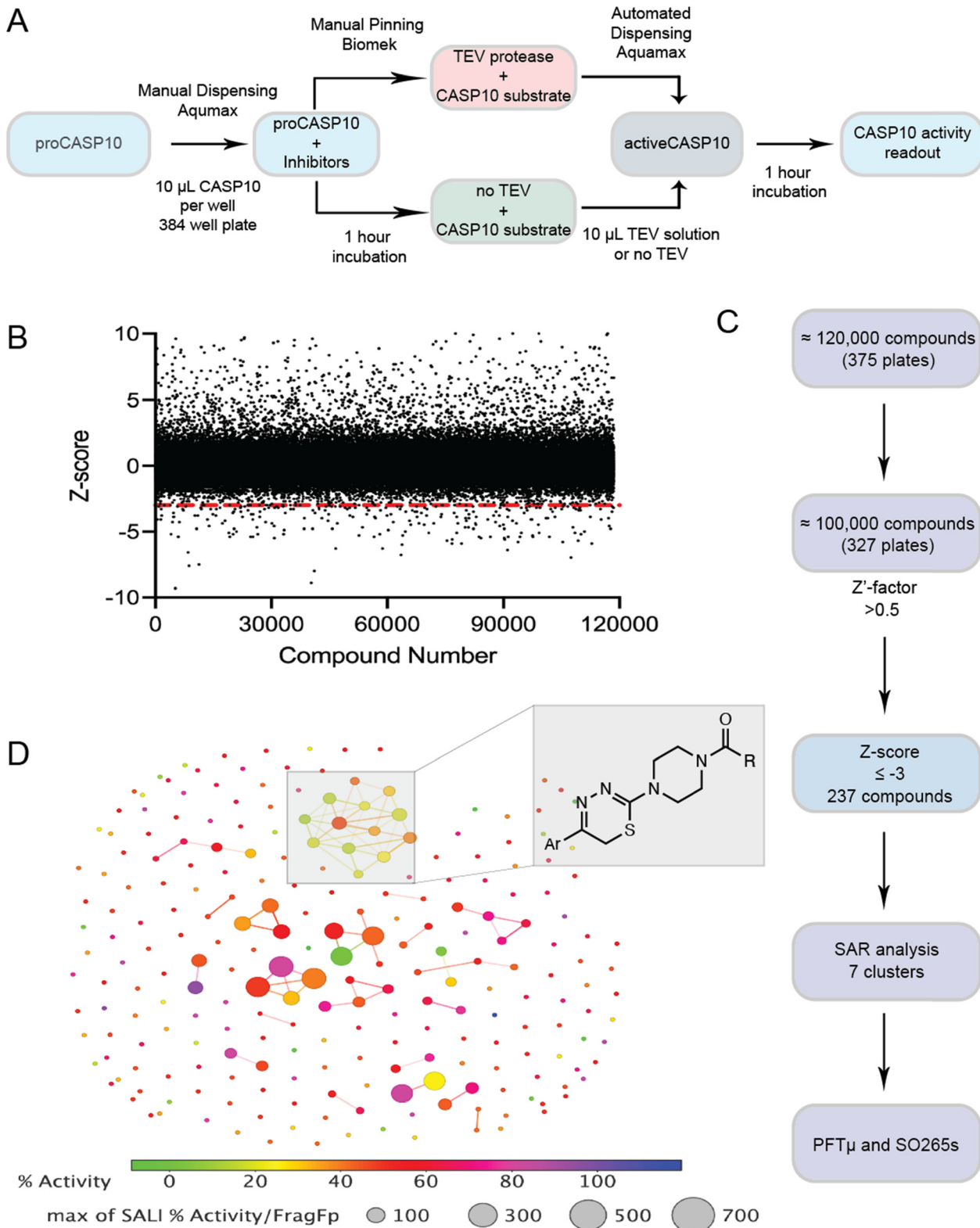


Fig. 2 TEV cleavable proCASP10TEV Linker enables a $>100\,000$ compound screen. (A) Scheme of HTS setup with proCASP10TEV Linker (333 nM concentration) incubated with screening compounds (10 μ M) for 1 h in a 384-well format followed by the addition of the fluorogenic substrate (Ac-VDAVAD-AFC, 10 μ M). Endpoint reads were measured and collected after 1 h incubation with the TEV substrate solution. (B) Z-scores of $>100\,000$ compounds (arbitrarily numbered) screened against proCASP10TEV Linker. The red line indicates hits below a Z-score of -3 . (C) Summary of filtering parameters for HTS screen. The first filtering step was to remove plates that were not within the desired range of Z'-factor (0.5–1.0), followed by a second filtering step that identified a total of 237 compounds with a Z-score of ≤ -3 . (D) SAR analysis of 237 hit compounds identified from HTS using the structure–activity landscape index (SALI) analysis as described in the DataWarrior SAR analysis methods section. The structure of recurring hit chemotypes identified by SAR cluster analysis (DataWarrior, v06.01.04)⁵¹ are shown for cluster 1. All screening data is in Table S1 (ESI†).



'Library of Pharmacologically Active Compounds' (LOPAC@1280), which is a widely utilized library of bioactive compounds that contain known promiscuous protease inhibitors such as E-64, along with a second library consisting of structurally diverse FDA approved drugs covering a broad spectrum of therapeutic areas.⁵⁰ Key features of our screen include (1) the pre-incubation of compound library members with the proCASP10TEV Linker construct to favor the detection of procaspase inhibitors and (2) the automated dispensing of the premixed TEV protease and caspase substrate solution followed by automated plate reading. These two automation steps were designed to ensure the consistent timing of our assay to minimize both plate-to-plate and library-to-library variability. Confirming the robustness of our approach, this initial screen had a Z' -factor above 0.5 across 8 screened plates, with a total of 30 out of 2569 compounds affording a >50% reduction in caspase-10 activity relative to DMSO control (Fig. S11, ESI[†]). Illustrating the robust performance of our screen, the Z' -factor comparing our positive control KB7 inhibitor-treated wells to DMSO-treated wells²⁷ was 0.90 (Fig. S12, ESI[†])—values above 0.5 are considered an acceptable range for HTS Z' -factor.⁴⁶

Large-scale screen identifies hit caspase inhibitors

Guided by the successful implementation of our pilot screen, we next deployed our platform to screen 118 498 total compounds (all at 10 μ M). In aggregate, this screen had Z' factor ranging from 0.4–1.0 and a hit rate of 0.81% for a total of 963 compounds, affording a 50% decrease in proCASP10TEV Linker activity (Fig. S13, ESI[†]), with 237 unique hits showing a Z -score ≤ -3 (Fig. 2(B)). The screen filtering steps are summarized in Fig. 2(C). Re-screen validation of the 237 compounds results in 38 compounds with <50% activity (Fig. S14A, ESI[†]). Z -Score analysis of the follow-up screen resulted in 97 (or approximately 41%) compounds below 3 standard deviations away from the mean of DMSO control samples (Fig. S14B, ESI[†]). Structural similarity clustering analysis using Datawarrior⁵¹ revealed that approximately 22% of the 237 hits (defined as Z -score ≤ -3 or those values that are three standard deviations lower than the mean of the DMSO controls calculated per plate) clustered together, providing evidence of *bona fide* inhibitors. The 7 major hit clusters consisting of over 50 compounds are shown in Fig. 2(D). One prominent and intriguing chemotype was a class of thiadiazine-containing compounds identified in cluster 1 (Fig. 2(D)). Upon closer inspection of the hits, several compounds with clear electrophilic groups stood out, including several organometallic species such as cisplatin (Table S1, ESI[†]), several cyanoacryl sulfones, such as **HTS-2** and the related **HTS-6**, which is a previously reported as non-specific κB kinase (IKK) inhibitor,⁵² and pifithrin- μ (**PFT μ**), a previously reported TP53 inhibitor⁴⁷ (Scheme S1, ESI[†]). As caspase-10 is a cysteine protease, we anticipated that some of these electrophilic compounds might react with the catalytic nucleophile.

Hit prioritization by re-screening and counter-screening against activeCASP10

To further interrogate our hits and confirm on-target activity against procaspase-10, our next step was determining if our hits

were selective for proCASP10 over the active proteoform. Using our previously reported fluorogenic caspase assay,²⁴ we measured the relative inhibition of recombinant active CASP10 by each of our 237 prioritized hits. This screen had a Z' factor of 0.73, resulting in 9 of 237 compounds having $\leq 50\%$ active-CASP10 activity (Fig. S15A and Table S1, ESI[†]), with 78 compounds being 3 standard deviations below the DMSO control (Fig. S15B, ESI[†]). Notably, several of our aforementioned hits, including **HTS-6**, **SO265**, and **PFT μ** , showed substantially reduced inhibition of activeCASP10 compared to the pro-CASP10TEV Linker (Fig. S14A and S15A, ESI[†]). SAR clustering analysis of the counter screen showed ~ 20 different clustering types for activeCASP10, but overall, less reactivity towards active compared to the pro-form (Fig. S16, ESI[†]), consistent with more favorable inhibition of the proenzyme. Together, our prioritized compounds (Scheme S1, ESI[†]) included 10 total inhibitors that showed preferential activity against proCASP10TEV when compared to activeCASP10.

TEV assay identifies likely TEV inhibitors

As our proCASP10TEV linker screen requires the addition of TEV protease for caspase activation, we anticipated that some hits might be *bona fide* TEV protease inhibitors rather than caspase inhibitors. To test this hypothesis and to filter out such compounds, we established a TEV protease activity assay using a customized fluorogenic substrate (see methods) inspired by prior TEV protease assays.⁵³ Following the workflow shown in Fig. 3(A), we first optimized both the substrate concentration and time points to ensure activity was within the enzyme's linear range (Fig. S17, ESI[†]). We find that TEV protease was highly active towards our fluorogenic substrate, with complete consumption of substrate within five minutes (Fig. 3(B)). Using automated dispensing to ensure the rapid and equal delivery of substrate across assay conditions, we observed comparable Michaelis–Menten kinetic parameters to those reported previously for TEV protease⁵⁴ (Fig. 3(C)).

Using this optimized assay, we used freshly sourced compound stocks to evaluate the TEV protease inhibitory activity of ten prioritized hits (Fig. 3(D)–(G) and Fig. S18, S19, ESI[†]). Notably, while most compounds were sourced commercially, compound **SO265** required in-house synthesis (Scheme S2, ESI[†] for the synthetic route for **SO265**). We observed substantial dose-dependent inhibition of TEV protease for **HTS-1** (Fig. 3(D)), **HTS-2** (Fig. S18A, ESI[†]), and **HTS-3** (Fig. 3(E)). Some inhibition was also observed for **HTS-6** (Fig. S18B, ESI[†]), **HTS-4** (Fig. S18C, ESI[†]), and **HTS-5** (Fig. S18D, ESI[†]). Therefore, we excluded these likely TEV inhibitors from further analysis. Encouragingly, several HTS hits stood out as having no appreciable TEV protease inhibitory activity (Fig. 3(F) and Fig. S19A, B, ESI[†]). Notably, **PFT μ** afforded a slight, but not significant, decrease in TEV protease activity at the highest tested concentration (100 μ M) (Fig. 3(G)).

Caspase-10 inhibition is confirmed by activity assays

We next turned to confirm that our screening hits engage both recombinant and endogenous proCASP10 protein. We first rescreened a subset of the prioritized hits in our proCASP10TEV



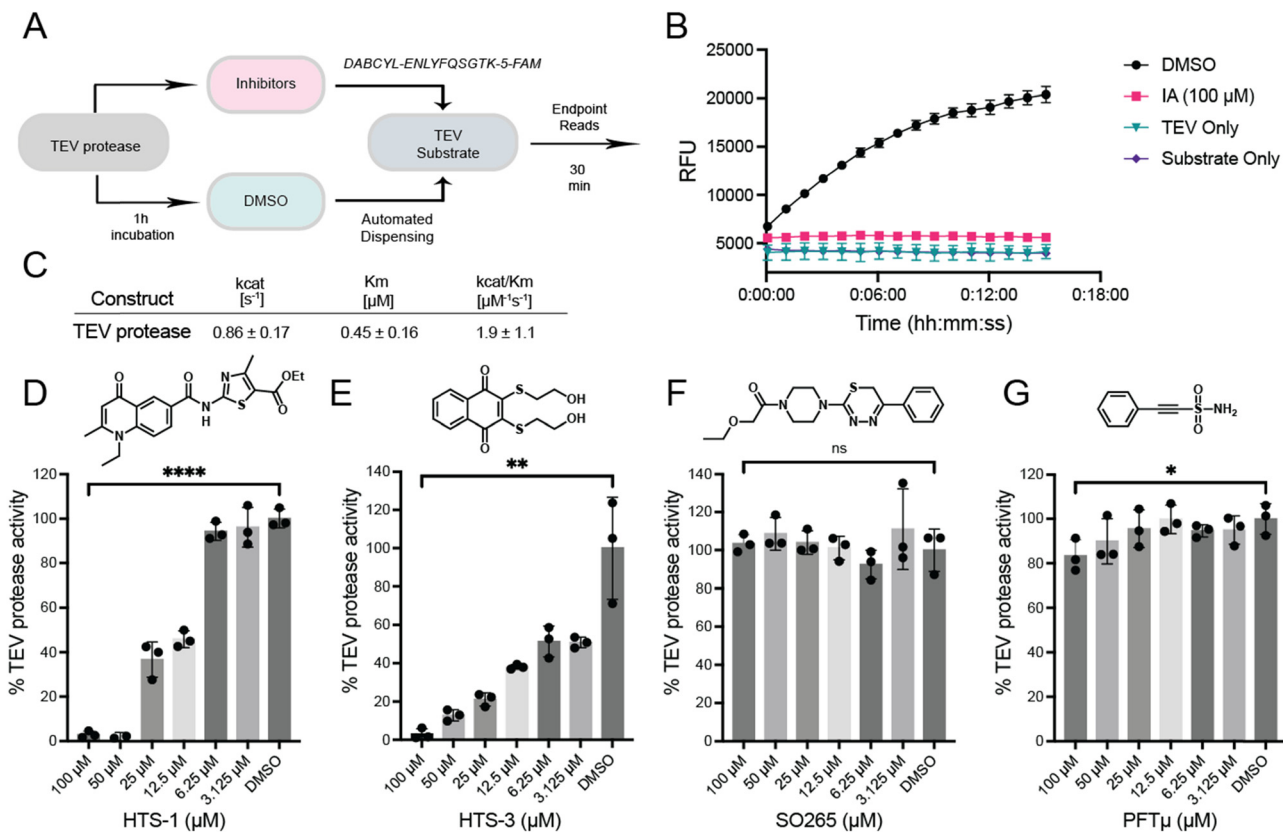


Fig. 3 Procaspase-10 HTS identified TEV protease inhibitors. (A) General scheme of TEV protease activity assay with our in-house TEV cleavable fluorogenic substrate, **DABCYL-ENLYFQSGTK-5-FAM**. (B) Shows relative fluorescent units (RFUs) for time-dependent increased fluorescence ($\lambda_{ex} = 495$ nm $\lambda_{em} = 550$ nm) for TEV protease (100 nM) exposed to either vehicle (DMSO) or iodoacetamide (IA). (C) Michaelis–Menten kinetics of TEV protease (667 nM) with TEV substrate. (D)–(G) TEV protease (50 nM) activity assay treated with the indicated screening compounds at the indicated concentrations for 1 h in PBS buffer under ambient conditions. For D–G, data represent mean values ± standard deviation for three biological replicates. Statistical significance was calculated with unpaired Student's *t*-tests, **p* < 0.05, ***p* < 0.01, ****p* < 0.0001, ns, not significant *p* > 0.05.

Linker assay. While we observed that most compounds only showed modest caspase-10 inhibition (Fig. S20, ESI[†]), several compounds stood out, including **HTS-6** (Fig. S21, ESI[†]), the TEV protease inhibitor **HTS-2** (Fig. S22, ESI[†]), and **PFTμ** (Fig. S23, ESI[†]). We then turned to gel-based activity-based protein profiling (ABPP) analysis to further corroborate our inhibition data. Having previously established ABPP gel-based assays for procaspase-8 and -10 using the **KB61** click probe²⁷ in HEK293T lysates, we first deployed this ABPP assay to assess compound engagement of recombinant procaspase-10 at the catalytic cysteine residue C401 by concentration-dependent competitive labeling of the initial screen compounds against the dual caspase-8/10 click probe **KB61** (Fig. S24, ESI[†]), focusing on the compounds that contained obvious electrophilic moieties, namely **HTS-6** and **HTS-2** and **PFTμ**. We find that **PFTμ** shows similar potency when compared to established caspase-8/10 dual inhibitor **KB7**.²⁷ Both **HTS-6** and **HTS-2** exhibited high proteome-wide reactivity, indicating that their caspase engagement is likely driven by the high electrophilicity of the cyanoacryl sulfone moiety. Further illustrating the increased reactivity of these two compounds relative to **PFTμ**, we also observed increased competition of iodoacetamide-rhodamine (IA-Rho) (Fig. S25, ESI[†]), consistent with generalized cysteine reactivity rather than a caspase-10-

specific effect. Due to their high reactivity, we excluded **HTS-6** and **HTS-2**, from further analyses.

Compound rearrangement drives caspase inhibition for SO265, which shows preferential inhibition of the pro-form of caspase-10

As the **SO265** thiadiazine chemotype had shown pronounced procaspase-10 inhibition in our initial screen (approximately 80% proCASP10TEV Linker inhibition; Fig. 2(D) and Fig. S14, ESI[†]), we were surprised by the lack of activity using our resynthesized compound (20% proCASP10TEV Linker inhibition) (Fig. S26, ESI[†]). Substituted thiadiazine rings can isomerize to thiol imidazole moieties, gaining aromaticity.⁵⁵ Therefore, we hypothesized that such a rearrangement of **SO265**, which could be favored by lower pH, could be driving its inhibitory activity (Fig. 4(A)). To test this hypothesis, we subjected **SO265** to acidic conditions (10 mM HCl) prior to assessing caspase inhibitory activity. Consistent with our hypothesis, we observed increased inhibitory activity for stocks of **SO265** pretreated with acid (Fig. S26, ESI[†]). This activity was generally restricted to caspase-10, with a more modest increase in inhibition observed for TEV protease (Fig. S27, ESI[†]).

Guided by this increased activity, we next tested the putative isomer, **SO265i** (Fig. 4(A)), which we had isolated from the



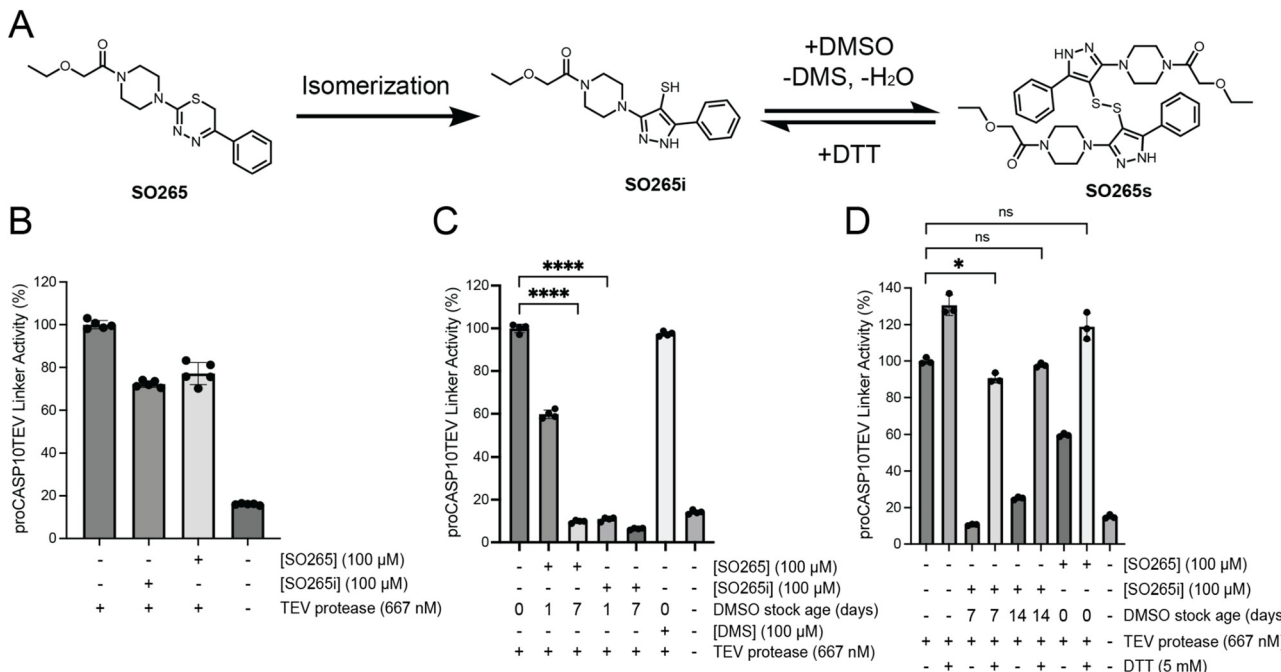


Fig. 4 The rearranged product of **SO265** inhibits proCASP10TEV linker activity. (A) Proposed scheme of **SO265** isomerization and formation of the disulfide product (B) Relative activity of proCASP10 Linker protein (333 nM) treated with the indicated compounds for 1 h followed by addition of TEV protease (667 nM) and fluorogenic substrate (Ac-VDVAD-AFC, 10 μM, and 333 mM sodium citrate) in PBS buffer. (C) The indicated compound stocks were subjected to either one- or seven-day incubation in DMSO at ambient conditions. Subsequently, compound-treated proCASP10 Linker protein was assayed as in 'B' with 1 h pre-exposure to either dimethyl sulfide (DMS) or the indicated compounds. (D) Compounds incubated for the indicated days in DMSO were further subjected to DTT (5 mM DTT) followed by evaluation for caspase inhibition as described in 'B'. For B-D, data represent mean values \pm standard deviation for four biological replicates. Statistical significance was calculated with unpaired Student's *t*-tests, $*p < 0.05$, $****p < 0.0001$, ns, not significant $p > 0.05$.

reaction mixture during the synthesis of **SO265**. While the masses of **SO265** and **SO265i** are identical, the two compounds can be distinguished *via* H-NMR analysis (Fig. S28A, ESI[†]) and LC-MS analysis (Fig. S28B-F, ESI[†]). Notably, the thiadiazine methylene signal was only present in **SO265**, with loss in the **SO265** spectra indicating that aromatization had indeed occurred. The **SO265i** spectra uniquely featured a signal for a thiol proton, consistent with the proposed isomerization. The presence of a thiol in **SO265i** was further corroborated by IR analysis (Fig. S29, ESI[†]). However, when tested in our activity assay, **SO265i** exhibited equivalent caspase-10 inhibition (30% inhibition) to **SO265**, indicating that the thiol imidazole was not the active species (Fig. 4(B)).

Following a 2-day incubation at room temperature in DMSO, repeated NMR analysis revealed that the **SO265i** stock had converted into a second species. This species was distinguished by the loss of the thiol proton and an increase in a signal matching the expected shift of dimethylsulfide (DMS) (Fig. S28A, ESI[†]). We hypothesized that DMSO-mediated oxidation of the **SO265i** thiol to a disulfide (**SO265s**) was occurring; this is a reported redox reaction that produces DMS as a byproduct and is promoted by low thiol pK_a values.^{56,57} Therefore, to further assess whether **SO265s** was the active species responsible for caspase-10 inhibition, we incubated 50 mM DMSO stocks of **SO265** and **SO265i** at room temperature for one, seven, and 14 days to promote the formation of **SO265s**. We then tested the activity of each of these compound stocks alongside DMS as a

control (Fig. 4(C)). Consistent with **SO265s** as the active species, we observed a time-dependent increase in inhibition of proCASP10TEV Linker for both **SO265** and **SO265i**; DMS had no effects on proCASP10TEV Linker activity. Providing further evidence of the likelihood that the rearranged disulfide structure is the active compound, the addition of 5 mM DTT to the compound stock completely abolished inhibitory activity (Fig. 4(D)). **SO265s** also exhibited increased reactivity with glutathione when compared to **SO265** and **SO265i** (Fig. S30A, ESI[†]), indicative of the likelihood that **SO265s** functions as a cysteine-reactive covalent inhibitor. Gel-filtration of **SO265s**-modified protein failed to recover proCASP10TEV Linker activity, which further corroborates that **SO265s** acts as a covalent inhibitor (Fig. S30B, ESI[†]) that shows reduced potency compared to **PFT1** (Fig. S30C, ESI[†]). Competitive gel-based ABPP analysis provided initial evidence that **SO265s** likely functions by labeling non-active site cysteine residue in caspase-10 (Fig. S30D, ESI[†]). Taken together, these data provide evidence that **SO265s** is a cysteine-reactive compound that blocks proCASP10TEV Linker activity with preferential inhibition of the zymogen (Fig. S31, ESI[†]). More broadly, we find that **SO265s** show some caspase selectivity, also inhibiting active caspase-8 (Fig. S32, ESI[†]), whereas only a slight decrease in activity was observed for active caspase-3 (Fig. S33, ESI[†]) and caspase-9 (Fig. S34, ESI[†]).

CETSA confirms caspase-10 labeling

As all our data, thus far, had been assessed for recombinant protein, we next opted to extend our analyses to assess the



compound engagement of endogenous procaspase-8 and -10. For this, we turned to Cellular Thermal Shift Assay (CETSA)⁵⁸ to measure binding-induced changes to protein thermal stability. To validate our assay, we subjected Jurkat lysates spiked with recombinant hexahistidine-tagged proCASP10 to CETSA analysis, comparing the thermal stability of the proCASP10 construct to that of the inactive catalytic cysteine mutant construct (proCASP10 C401A) with and without addition of **KB7**, as a positive control. We observe a marked decrease in thermal stability upon **KB7** treatment for both the endogenous (anti-caspase-10 signal) and recombinant (anti-His signal) caspase-10 proteins (Fig. 5(A) and complete blots in Fig. S35, ESI[†]). The C401A mutant protein did not show a similar thermal shift (Fig. S36, ESI[†]), which is consistent with the covalent modification of the C401, which is the catalytic nucleophile, by **KB7**. Extension of this analysis revealed that, like **KB7**, **PFTμ** induced destabilization of

the wildtype spiked proCASP10 and endogenous proCASP10 (Fig. 5(B) and complete blots in Fig. S37, ESI[†]) but not the C401A mutant protein (Fig. 5(C) and complete blots in Fig. S38, ESI[†]), which further confirms that **PFTμ** labels C401. Unexpectedly, the disulfide product (**SO265s**) caused some protein stabilization, suggesting an alternate mode of action compared to the two active site-directed inhibitors (Fig. 5(D) and complete blots in Fig. S39, ESI[†]). Corroborating an alternate mode of engagement by **SO265s**, we observed no competition in gel-based ABPP analysis using the **KB61** click probe against procaspase-10 (Fig. S40A, ESI[†]), unlike **PFTμ** that labeled both procaspase-10 (Fig. S40A, ESI[†]) and procaspase-8 (Fig. S40B, ESI[†]). Disappointingly, we observed no similar compound-induced thermal stability shift for endogenous procaspase-8 upon **KB7** treatment (Fig. S35, ESI[†]) nor **PFTμ** treatment (Fig. S37, ESI[†]), indicating that the CETSA assay was not suitable for evaluation of caspase-8

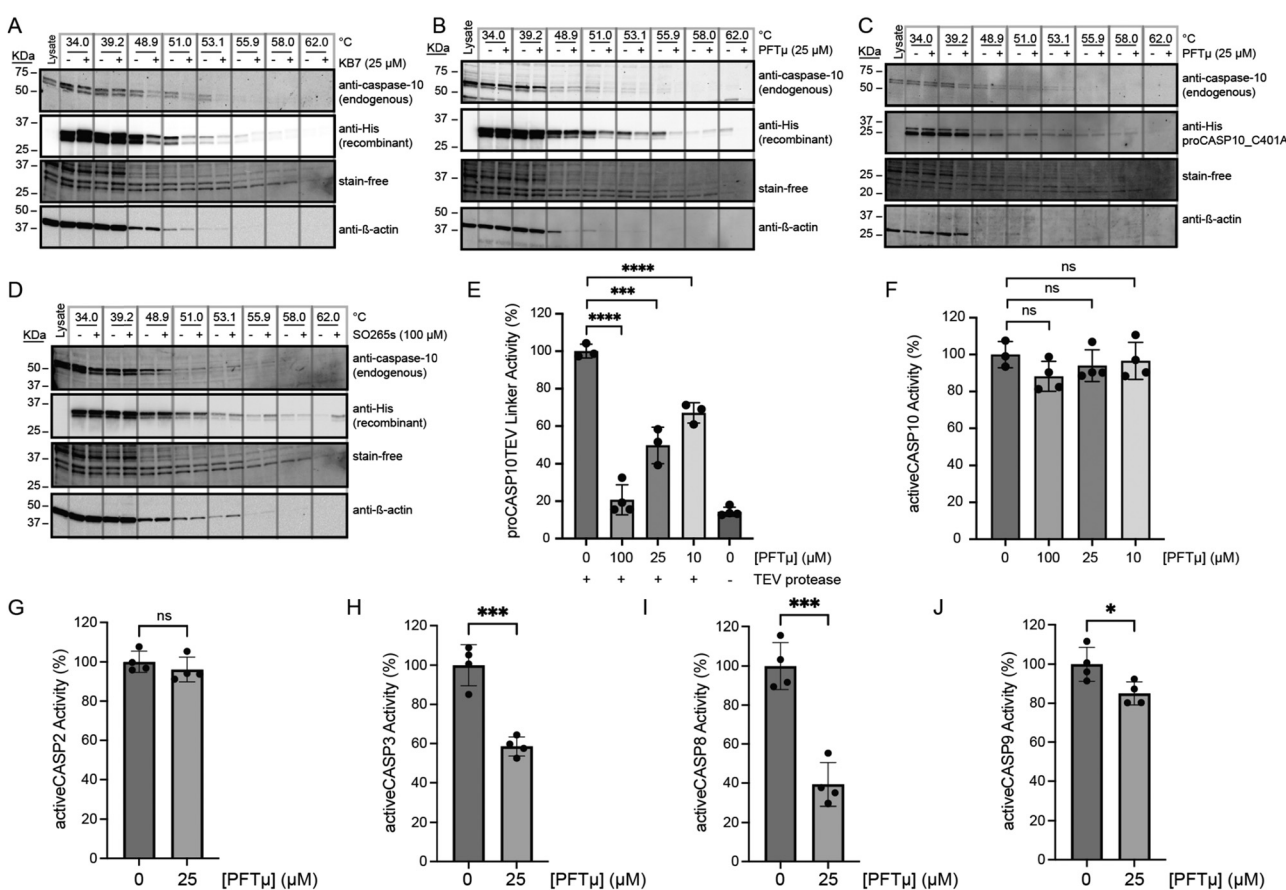


Fig. 5 **PFTμ** and **SO265s** impact the stability of both recombinant and endogenous procaspase-10. (A)–(D) Cellular thermal shift assay (CETSA)⁵⁸ analysis blot of Jurkat cell lysates spiked with the indicated recombinant caspase-10 proteins featuring C-terminal hexahistidine tag and treated with the indicated compounds at the indicated concentrations or vehicle (DMSO) and subjected to immunoblot analysis for using both anti-His and anti-caspase-10 antibodies to visualize recombinant and endogenous protein, respectively. Loading was visualized both with the BioRad Chemidoc Stain-Free imaging technology⁵⁹ and with anti-β-actin. For 'A', 'B', and 'D', recombinant proCASP10 and for 'C', recombinant proCASP10_C401A were analyzed. (E) Relative activity of proCASP10TEV Linker protein (333 nM concentration in PBS buffer) exposed to **PFTμ** at the indicated concentrations for 1 h followed by TEV protease (667 nM) and analysis with Ac-VDVAD-AFC fluorogenic substrate (10 μM) in PBS supplemented with 333 mM sodium citrate. (F)–(J) Relative activity of recombinant active caspases in PBS buffer analyzed with Ac-VDVAD-AFC fluorogenic substrate (10 μM) after treatment with the indicated concentrations of **PFTμ** for 1 h. For activeCASP10, fluorogenic substrate (10 μM) in PBS was supplemented with 333 mM citrate. Recombinant proteins used in 'F' activeCASP10 (333 nM), 'G' activeCASP-2 (0.3 μM), 'H' activeCASP-3 (0.3 μM), and 'I' activeCASP9 (1 μM). For A–D, data is representative of two biological replicates. For E–J, data represent mean values ± standard deviation for four biological replicates. Statistical significance was calculated with unpaired Student's *t*-tests, **p* < 0.05, ***p* < 0.01, ****p* < 0.0001, ns, not significant *p* > 0.05.



target engagement. Therefore, we returned to enzyme activity assays for broader target analysis.

Pifithrin- μ is a promiscuous caspase inhibitor that prevents FasL-mediated apoptosis

In contrast with the complex activation mechanism of **SO265s**, which likely could complicate assessing in-cell activity, we expected **PFT μ** to retain caspase inhibitory activity in complex cell environments. This expectation is further supported by **PFT μ** 's well-documented anti-apoptotic activity, which has been ascribed to its function as a TP53 inhibitor.⁴⁷ Therefore, we prioritized **PFT μ** for further analysis. Activity assay analysis revealed that **PFT μ** showed some preferential inhibition across a panel of analyzed caspases; consistent with our screening data, we observed preferential inhibition of procaspase-10 when compared to active caspase-10 (Fig. 5(E), (F) and Fig. S14, S15, ESI[†]). Similarly, **PFT μ** also preferentially labeled procaspase-2, as analyzed using our caspase-2 directed click probe²⁴ and ABPP analysis (Fig. S41, ESI[†]). No detectable inhibition of active caspase-2 was observed, either by ABPP gel (Fig. S41, ESI[†]) or activity assay (Fig. 5(G)). **PFT μ** also significantly inhibited active caspase-3 and active caspase-8, with slight, albeit significant, inhibition observed for active caspase-9 (Fig. 5(H)–(J)). These data provided evidence that **PFT μ** is a promiscuous caspase inhibitor.

We next turned to chemoproteomics to more broadly assess whether **PFT μ** labels endogenous caspases. We deployed our established cysteine chemoproteomic platform⁶⁰ in which covalent labeling sites are identified in a competitive manner using the pan cysteine-reactive probe iodoacetamide alkyne (**IAA**) and isotopically enriched “light” and “heavy” biotin-azide capture reagents. Following the workflow shown in Fig. 6(A), out of 8925 total cysteines quantified, we find that 1070 total unique cysteines showed $\log_2(H/L)$ values greater than 1.5, indicative of labeling by **PFT μ** . Included in this list were several known targets, including PARP1, HSP70 (HSPA1A), and PRDX proteins (PRDX1, PRDX2, PRDX5), which were previously identified *via* chemoproteomics using a clickable **PFT μ** analog,⁶¹ providing evidence of the robustness of our approach (Table S2, ESI[†]). Quite strikingly, several caspases stood out as having high $\log_2(H/L)$ ratios (Fig. 6(B)), including caspase-8, which aligns with our aforementioned activity assay and gel-based ABPP analysis (Fig. 5(I) and Fig. S40, ESI[†]). Aligning with our prior discovery²⁴ of structurally-related phenylpropionate molecules that label caspase-2 at C370, we also observe that C370 is highly sensitive to **PFT μ** , which is also consistent with our gel-based analysis (Fig. S41, ESI[†]). The noncatalytic cysteine residue, C264, in caspase-6,²⁶ was also observed to be labeled by **PFT μ** , as was the near-active site cysteine (C170) in caspase-3. We did

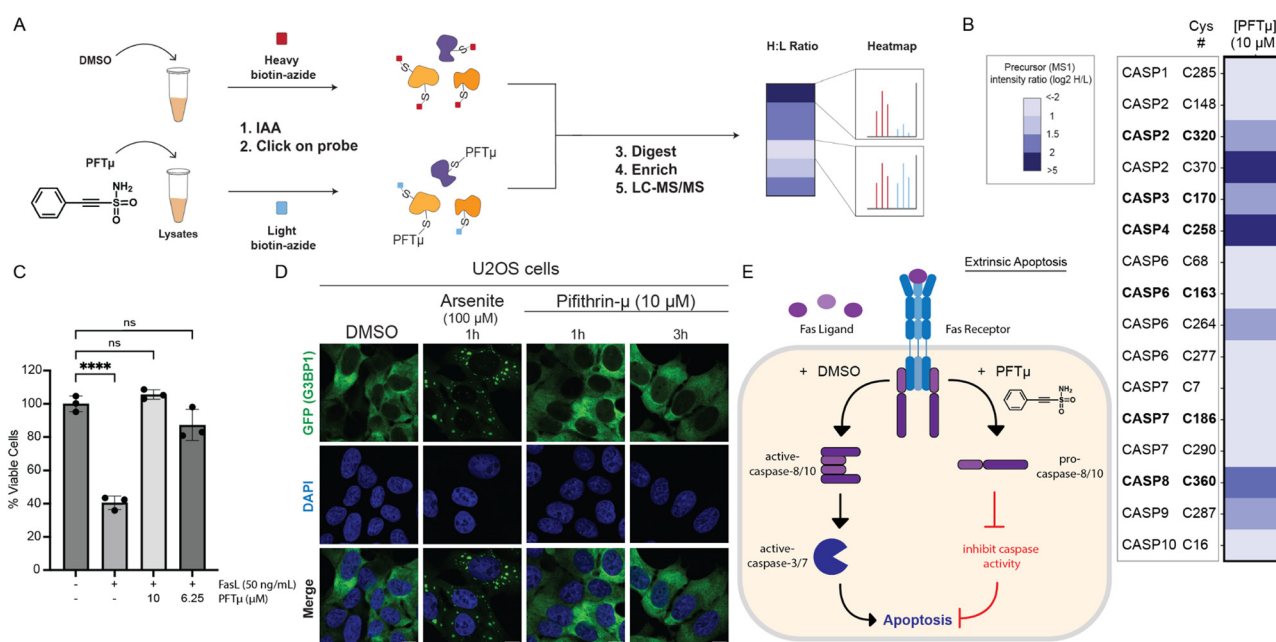


Fig. 6 **PFT μ** labels initiator caspases and can protect Jurkat cells from FasL-mediated apoptosis. (A, B) Cysteine chemoproteomic analysis identifies protein targets of **PFT μ** . 'A' shows the general workflow in which Jurkat cell lysates were treated with either **PFT μ** (25 μ M) or DMSO for 1 h, followed by IAA (200 μ M) cysteine capping, conjugated *via* click chemistry to isotopically differentiated biotin-azide reagents,^{62,63} with on-resin tryptic digestion, enriched, and analyzed by LC-MS/MS. Heatmap in 'B' shows the mean quantified precursor intensity ($\log_2(H/L)$) for DMSO (H, heavy) *versus* compound treatment (L, light) for all caspase cysteines identified. Catalytic cysteine residues are annotated in bold. (C) CellTiter-Glo[®] measurement of relative viability of Jurkat cells subjected to **PFT μ** at the indicated concentrations for 1 h followed by FasL (50 ng mL⁻¹, 3 h)-induced apoptosis. (D) Fluorescent microscopy of U2OS cells expressing GFP-G3BP1 and treated with **PFT μ** (10 μ M) or positive control sodium arsenite. All scale bars = 10 μ m. (E) Proposed model where **PFT μ** functions as a promiscuous initiator caspase inhibitor (labeling caspase-2, -8, and -10) and blocks FasL-mediated extrinsic apoptosis. For B, experiments were conducted in four biological replicates, with two samples additionally analyzed as technical replicates. For C, data represent mean values \pm standard deviation for three biological replicates. All MS data can be found in Table S2 (ESI[†]). Statistical significance was calculated with unpaired Student's *t*-tests, *****p* < 0.0001, ns, not significant *p* > 0.05.



not detect the catalytic cysteine nucleophile of caspase-3 (C163). Beyond these pro-apoptotic caspases, the catalytic cysteine in the pro-inflammatory caspase-4 (C258) was also labeled by **PFT μ** . The catalytic cysteine in caspase-10 was not identified, likely due to the long tryptic peptide that flanks this residue. Taken together, these data provide further evidence that **PFT μ** promiscuously labels many human caspases in addition to its previously reported targets.

Guided by these findings, we next asked whether **PFT μ** could protect cells from extrinsic apoptosis induced by Fas ligand (FasL). While **PFT μ** 's anti-apoptotic activity has been reported in other contexts,^{47,66–69} we selected FasL-mediated apoptosis as a model system due to the central role that caspase-8 and -10 play in this process^{12,70,71} and the less central role of TP53,^{72,73} with the goal of helping to define better the biologically active target(s) of **PFT μ** 's anti-apoptotic activity. Using the CellTiter-Glo assay, we find that **PFT μ** affords near-complete protection from FasL (Fig. 6(C)) without inducing cytotoxic effects (Fig. S42, ESI[†]). These data align with caspase inhibition as contributing to **PFT μ** 's activity.

As our recent work has shown that electrophile stress can lead to the formation of stress granules,⁷⁴ we also opted to rule out **PFT μ** -induced stress granules as a potential confounding variable. Using a $\Delta\Delta G3BP1/2$ KO cell line that stably expresses GFP-G3BP1,^{75,76} we find that **PFT μ** does not induce G3BP1 condensates using concentrations that are protective from apoptosis (Fig. 6(D)) and are consistent with prior studies using **PFT μ** as a putative TP53 inhibitor.⁴⁷ Thus, we put forth a model (Fig. 6(E)) that caspases are likely biological targets of **PFT μ** that contribute to the reported anti-apoptotic activity.

Discussion

To enable high throughput procaspase-10 inhibitor discovery, here we developed and applied an engineered TEV-activatable caspase-10 protein to high throughput screening. Our high quality (an average Z' of 0.58) $\sim 100\,000$ compound semi-automated screen yielded 237 total hits with Z -score <-3 . Subsequent rescreening and counter-screening delineated *bona fide* procaspase-10 inhibitors from those with activity against active caspase-10 or TEV protease. For screening hit **PFT μ** , orthogonal mode of action studies confirmed target engagement for both recombinant and endogenous procaspase-10. More broadly, we also expect that our and related activation screening platforms should prove broadly useful in the discovery of inhibitors targeting precursor proteases.

For such future activation-based caspase-10 high throughput screening campaigns, our work highlights intriguing opportunities. As TEV protease is a cysteine protease, we found that several of our hits functioned *via* engaging TEV rather than procaspase-10. We expect that these newly identified TEV protease inhibitors could serve as useful tools for synthetic biology studies that rely on TEV protease activity to control engineered circuits. To favor caspase rather than TEV inhibitor discovery, future studies could also consider using alternative

non-cysteine protease enzymes for activation, which we expect would prove more substantially orthogonal to caspase cysteine protease activity. In addition to uncovering candidate TEV inhibitors, our screen also revealed that the thiadiazine chemotype is likely prone to disulfide rearrangement, as exemplified by our characterization of **SO265s**, which we found was the active species. Thus, our work both highlights the value of resynthesis in confirming screening hits and hints at the likely amenability of procaspase-10 to a disulfide trapping/tethering-style inhibitor discovery^{77–79} using additional disulfide-containing compounds.

Beyond the **SO265** thiadiazine chemotype, **PFT μ** , a reported TP53 inhibitor, proved to be a key hit from our screen. Comparing active and procaspase-10 inhibitor activity, we found that **PFT μ** inhibited both proteoforms to some degree, with increased inhibition observed for the proenzyme. Looking beyond caspase-10, the *in vitro*, cell-based, and proteomic analysis together revealed that **PFT μ** also engages caspase-8, -3, -10, -5, and -6. These data allow us to put forth a model whereby **PFT μ** 's reported activity as an anti-apoptotic agent is likely driven by caspase inhibition in addition to the previously reported TP53 inhibitory activity.⁴⁷ Therefore, we anticipate that the cysteine-reactive ethynesulfonamide chemotype could prove broadly useful as a starting point for caspase inhibitor development campaigns. Further medicinal chemistry efforts will be needed to determine whether the reactivity and potency of ethynesulfonamide-containing compounds can be tuned to a level suitable for a selective chemical probe. We also expect our work to aid in phenotype interpretation for studies that have employed **PFT μ** to block p53 activity.

We also recognize some limitations of our work. Despite our extensive screening efforts, we failed to obtain potent, selective caspase-10 inhibitors, and, as such, we were unable to further inform the still cryptic functions of caspase-10. Due to the comparatively modest potency and low selectivity of our initial screening hits, we opted to proceed conservatively and restrict our mode-of-action studies to lysate- rather than cell-based analyses, so as to exclude possible electrophilic stress-mediated confounding activity, as has recently been reported for other classes of cysteine-reactive electrophiles.^{74,80} Our ABPP and CETSA analysis of **SO265s** suggest that caspase-10 harbors additional non-catalytic and possibly unique, ligandable cysteine residues. However, we have yet to pinpoint the positions and tractability of these putative cysteines. Future efforts to more fully characterize these still cryptic cysteines will benefit from more in-depth chemoprotomic and mutational analysis of caspase-10, together with solving the atomic resolution structure of caspase-10, which remains unresolved in the PDB.

In sum, our study sheds important light on the activity of two classes of cysteine-reactive electrophiles and provides a useful screening platform for future caspase-10 inhibitor discovery campaigns. We remain optimistic about the feasibility of developing a caspase-10 selective chemical probe, and hope that our screening hits, particularly those that we have yet to subject to counter screening analysis, will enable the discovery of potent and selective caspase-10 inhibitors.



Author contributions

J. O. C. and K. M. B. conceived of the project. J. O. C., C. Y., B. H., K. H. A., and A. R. J. collected data. K. H. A. and S. O. provided reagents. J. O. C., R. D., and L. M. B. performed data analysis. L. M. B., M. F. P., and N. P. wrote software. K. M. B. and R. D. provided guidance and funding. J. O. C. and K. M. B. wrote the manuscript and all authors revised and edited the manuscript.

Data availability

The MS data have been deposited to the ProteomeXchange Consortium (<https://proteomecentral.proteomexchange.org>) via the PRIDE partner repository with the dataset identifier PXD053315.

Conflicts of interest

K. M. B. is a member of the advisory board at Matchpoint Therapeutics. All remaining authors declare no conflicts of interest.

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