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Engineered cellular gene-replacement platform for selective and inducible proteolytic profiling

Charles W. Morgan^{a,b,c}, Juan E. Diaz^b, Samantha G. Zeitlin^b, Daniel C. Gray^{a,b,c,1}, and James A. Wells^{b,c,2}

^aChemistry and Chemical Biology Graduate Program, University of California, San Francisco, CA 94143; ^bDepartment of Pharmaceutical Chemistry, University of California, San Francisco, CA 94143; and ^cDepartment of Cellular and Molecular Pharmacology, University of California, San Francisco, CA 94143

Edited by Benjamin F. Cravatt, The Scripps Research Institute, La Jolla, CA, and approved May 28, 2015 (received for review March 6, 2015)

Cellular demolition during apoptosis is completed by executioner caspases, that selectively cleave more than 1,500 proteins but whose individual roles are challenging to assess. Here, we used an optimized site-specific and inducible protease to examine the role of a classic apoptotic node, the caspase-activated DNase (CAD). CAD is activated when caspases cleave its endogenous inhibitor ICAD, resulting in the characteristic DNA laddering of apoptosis. We describe a posttranscriptional gene replacement (PTGR) approach where endogenous biallelic ICAD is knocked down and simultaneously replaced with an engineered allele that is susceptible to inducible cleavage by tobacco etch virus protease. Remarkably, selective activation of CAD alone does not induce cell death, although hallmarks of DNA damage are detected in human cancer cell lines. Our data strongly support that the highly cooperative action of CAD and inhibition of DNA repair systems are critical for the DNA laddering phenotype in apoptosis. Furthermore, the PTGR approach provides a general means for replacing wild-type protein function with a precisely engineered mutant at the transcriptional level that should be useful for cell engineering studies.

DNA damage | apoptosis | ICAD | site-specific proteolysis | TEV protease

Apoptosis, programed cell death, is crucial during devel-opment and for maintaining tissue homeostasis in an adult organism (1–3). Dysregulation of apoptosis can have drastic disease consequences, such as cancer, autoimmunity, and neurodegeneration (4–6). Apoptosis is a tightly controlled cellular process that leads to the contained demolition of individual cells and preparation for their noninflammatory disposal through engulfment by phagocytic cells (7, 8). Induction of apoptosis causes prominent morphological changes, such as cellular blebbing and DNA laddering, resulting from chromatin cleavage (9, 10).

A great deal is known about the signaling events that initiate the hard-wired program of apoptosis that leads to caspase activation (11–13), cysteine-class aspartate-specific proteases that deconstruct the cell. The number of caspase substrates continue to grow every year, with over 1,500 substrates identified from several complementary global proteomic approaches (14–18). The substrate cleavage rates vary over 500-fold in cells (19). Although substrate identification and kinetics can begin to shed light on the proteolysis process, they do not address the role of specific cleavage events in apoptosis. Our understanding of how single substrates can drive apoptosis or how they depend on coordination with other substrates, thereby forming nodes of synthetic lethality, is lacking. Gene disruption and overexpression of cleaved substrates have suggested the importance of particular substrates in driving the apoptotic process (reviewed in ref. 20). However, overexpression approaches cannot mimic the temporal aspects or N- and C-terminal proteolytic fragments resulting from caspase proteolysis. Thus, there is a need to develop approaches for inducible site-specific proteolysis in cells to assess the role of specific substrates in apoptosis.

Recently, our group has developed an enzyme called the SNIPer (Single Nick in Proteome) that is capable of cleaving individual substrates in cells (21). The SNIPer is a split-protein version of NIa tobacco etch virus protease (TEVP), which relies on chemical induced dimerization of FRB (FKBP rapamycin binding)

and FKBP in the presences of rapamycin for selective activation. The SNIPer was first applied to determine the role of selective proteolysis that activates individual caspases in apoptosis. By replacing the normal caspase cleavage sites in the executioner caspases-3, -6, and -7 with the unique TEVP recognition sequence [ENYLFQ/(G or S)], it was possible to generate caspase^{Te $\tilde{\sigma}$ S iso-} forms that could be cleaved and activated by TEVP upon small molecule addition. Interestingly, these experiments demonstrated that direct activation of a single substrate, caspase-3 or -7, lead to apoptosis, whereas caspase-6 does not (21) . Can other individual substrates do the same?

DNA fragmentation is a catastrophic event for cells and a hallmark of apoptosis. DNA laddering is the result of doublestranded breaks at accessible regions of DNA between nucleosomes, thereby producing discrete chromosomal fragments (multiples of ∼180 bp) (22). The inhibitor caspase-activated DNase (ICAD)–CAD complex is believed to be the central apoptotic node for this fragmentation (23). Caspase cleavage of the regulator ICAD results in release, dimerization, and activation of CAD. Splicing of the ICAD transcript produces two variants, a cytosolic localized short (ICADS) isoform and the nuclear localized long (ICADL) isoform (24). Additionally, ICADL serves as a mandatory folding chaperone for CAD (24). Cells deficient in ICAD lack the CAD protein and DNA fragmentation, suggesting CAD is required for DNA fragmentation (25). Two types of apoptotic DNA fragmentation have been previously described: a high molecular weight and lower molecular weight cleavage. CAD has been implicated in generating either high molecular weight or lower molecular weight, or both, depending on the specific cell line and method used (26). Is CAD activation sufficient to cause DNA fragmentation and to kill cells?

Significance

Apoptosis, a type of programmed cell death, is critical for the removal of diseased or superfluous cells and is essential in maintaining tissue homeostasis. At the molecular level, apoptosis leads to the highly coordinated selective proteolysis of more than a thousand cellular proteins. Until now, determining the individual molecular consequence of proteolysis has been largely out of reach. Our article reports biotechnological advances that enable rapid and reliable posttranscriptional gene replacement for robust inducible and selective proteolysis of the replacement allele. We go on to further show the utility of our technology in deciphering synthetic lethality resulting from coordinated proteolysis of caspase substrates that control the apoptotic hallmark of chromatin fragmentation.

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¹Present Address: Warp Drive Bio, LLC, Cambridge, MA 02139.

²To whom correspondence may be addressed. Email: [jim.wells@ucsf.edu.](mailto:jim.wells@ucsf.edu)

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Here, we present a system to test the cellular consequences for direct activation of CAD through site-specific proteolysis of ICAD using two small-molecule inducible TEVP strategies. This process required that we simultaneously deplete the endogenous transcripts for caspase sensitive ICAD and introduce TevS variants at physiological levels, an approach we call posttranscriptional gene replacement (PTGR). The recent advances in clustered regularly interspaced short palindromic repeats (CRISPR)-mediated genomic engineering are now routine for studies involving gene knockout and gene activation experiments; however, site-specific mutagenesis remains a substantial hurdle for efficient and robust Cas9 nickase-induced homology-directed repair (27). Using a single lentiviral vector for PTGR-mediated replacement and cellular engineering is a well-suited technology for experiments requiring replacement with site-specific mutations.

Combining selective and inducible proteolysis with substrate replacement, we discovered that CAD activation alone did not lead to substantial DNA fragmentation or cell death, but could only enhance fragmentation levels when combined with DNA-repair inhibitors or drug-induced chromatin relaxation. These data suggest that CAD cannot act alone in cancer cells, and that constitutive DNA damage response is sufficient to control CAD inflicted DNA damage (28). Moreover these studies show the power of combining a next generation of inducible TEVP strategies with a loss-of-function substrate through PTGR-mediated replacement for selective proteolysis of substrates in cells.

Results

Simultaneous Protein Knockdown and Replacement Using PTGR. The CAD inhibitor, ICAD, is inactivated by caspase cleavage, resulting in a loss-of-function event. To place this event under proteolytic control of the SNIPer required concurrently reducing levels of the endogenous ICAD allele and introducing the TEVP sensitive (TevS) allele. The PTGR strategy allows simultaneous protein knockdown and replacement through the bicistronic expression of an engineered gene followed by a synthetic microRNA (miRNA) targeting the endogenous gene's UTR or coding sequence (Fig. 1A and [Fig. S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF1)). The coexpression of each cassette results in miRNA reduction of the endogenous gene while simultaneously expressing the engineered substrate (Fig. 1B). The native caspase cleavage site is precisely replaced with a TEVP cleavage sequence, resulting in the orthogonal *substrate*^{TevS}, which can no longer be

Fig. 1. Strategy for simultaneous protein knockdown and replacement using PTGR. (A) The PTGR vector uses a synthetic miRNA that is complementary to the endogenous target mRNA but is genetically orthogonal to the mutated region or UTR in the replacement allele. (B) The endogenous protein levels diminish as the orthogonal allele's expression levels increase as a result of cistronic expression of the microRNA and the orthogonal allele. (C) The three components of the PTGR vector are inserted in a lentiviral backbone for targeting a wide variety of cell lines. Multisite Gateway cloning enables quick assembly of the final vector from donor plasmids, synthetic microRNA, and the orthogonal replacement allele. Donor plasmids are assembled and screened independently. The third component consists of an IRES sequence for cap-independent initiation of a combination selectable marker including both drug resistance and GFP fluorescence, separated by a "self-cleaving" T2A peptide sequence.

cleaved by caspases. In addition, we have made a silent mutation in the replacement allele (*substrate*^{TevS}) to make it resistant to the miRNA. For robust and broad specificity, the PTGR cassette is placed in a lentiviral backbone, pHR (Fig. 1C) (29). To achieve optimal tuned expression, we used a dual-selection strategy that relied on drug selection followed by stringent flow cytometry sorting of GFP⁺ cells.

We tested functional knockdown and simultaneous replacement using a series of ICAD replacement alleles in HEK293 and HeLa cell lines (Fig. 2 and [Fig. S2](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF2)A). We observed a substantial reduction in ICAD protein levels in the ICAD knockdown cell line compared with empty vector HEK293 cells. As expected, there was decreased CAD expression in the absence of ICAD, consistent with its role as a folding chaperone. Upon ICAD replacement with engineered alleles, we observed expression of ICAD^{TevS} and corresponding rescue of CAD levels. Furthermore, CAD expression levels remained unchanged by PTGR-mediated replacement with WT or TevS alleles (Fig. 2). Taken together, these data suggest that (i) the ICAD^{TevS} allele was resistant to the orthogonal miR and (ii) both WT and LTevS alleles are able to act as chaperones for CAD folding during translation. We crossvalidated the immunoblot results using quantitative RT-PCR (RT-qPCR) to directly monitor ICAD transcript levels [\(Fig. S2](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF2)). The parental cell line expression level was set at 1.00, and mean value of expression of ICAD in PTGR ICAD knockdown cell line was 0.13 (+0.12/-0.05), corresponding to ∼85% reduction of ICAD expression.

Induced and Selective Cleavage of ICAD^{TevS}. We sought to preserve the natural pathway of proteolytic CAD activation and retain ICAD's chaperone and inhibitor function (Fig. 3A). To complete this objective we combined our PTGR strategy with inducible and selective proteolysis of TEVP. Before combining PTGR and split-TEVP strategies, we next optimized the inducible protease approach. The split-TEVP, SNIPer, relies on the heterodimerization of FKBP and FRB in the presence of cell permeable rapamycin to assemble the N- and C-terminal halves of TEVP and activate the protease. A low level of TEVP activity is detectable with the SNIPer in the absence of rapamycin. Furthermore, split-TEVP has reduced activity upon rapamycin activation, likely a consequence incomplete assembly or instability of the split-protein (30). Although the lower activity of the SNIPer is adequate for testing apoptotic nodes undergoing positive feedback (such as caspase activation), we found it was less optimal for the current application where we wanted rapid and complete proteolysis of the ICAD^{TevS} substrate to ensure depletion of ICAD pools while outpacing substrate resynthesis rates.

As a result, we explored a variety of transcriptional and translational strategies to enhance inducible TEV proteolysis (Fig. 3B and [Fig. S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF3). We first generated a single vector dualexpression system, using an internal ribosome entry site (IRES) sequence to drive the expression of the second, C-TEV, fragment; however, this suffered from low expression levels of the second fragment and poor activity upon dimerization. Next, we used a 2A "self-cleaving" peptide sequence to drive cistronic translation of N-TEV and $\overrightarrow{C-TEV}$ in two orientations (31). This 2A configuration yielded robust protein expression levels and inducible activity, with the C-Tev_T2A_N-Tev being the better of the two. However, background activity in the absence of rapamycin was still a concern as expression levels were increased ([Fig. S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF3)C). As a result of the limitations observed in the split-TEVP strategy, we explored tetracycline inducible promoters to drive full-length TEVP expression we call iTEVP. As expected, when $ICADL^{TevS}$ was expressed in cells containing either the SNIPer or iTEVP and activated with rapamycin or tetracycline, respectively, we observed a substantial decrease in full-length ICADLTevS. However, the iTEVP exhibited better inducible and selective cleavage of the target protein in the shortest period (Fig. 3C). We therefore moved forward with characterizing the biochemical consequences in cells engineered with the iTEVP system.

Fig. 2. Biochemical evidence that the PTGR strategy reduces the endogenous target while producing the replacement target. A series of stable cell lines were made using the PTGR vector and protein levels were analyzed by immunobloting against ICAD and CAD in HEK293 (A) and HeLa (B) cells. EV corresponds to cells engineered with an empty vector. ICAD knockdown (KD) was completed with a PTGR vector expressing Renilla in the position of the replacement ICAD allele. Three different ICAD replacement cells lines were created using ICAD long (L) and containing either WT caspase cleavage sequences or TevS sequences. In A, ICAD knockdown results in an absence of CAD protein, whereas PTGR mediated replacement rescues CAD levels. The addition of the larger TevS sequence at the two cleavage sites causes a slight upward molecular weight shift in the ICADL^{TevS} allele. The short splice ICADSTevS allele is the clear lower molecular weight band in the ICAD blot (Top, far right). GAPDH was used as a protein loading control.

ICADL^{TevS} is rapidly cleaved after doxycycline addition (Fig. 3C). To confirm our engineering approach results in an active CAD species, we undertook a series of in vitro CAD activity DNA fragmentation assays to confirm that our activation strategy worked in the engineered tissue culture cells. We expressed and purified recombinant heterodimers of ICAD:CAD and ICAD^{TevS}:CAD dimers and treated with activating protease, caspase-3 or TEVP, respectively. In both cases this resulted in an active CAD species [\(Fig. S4](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF4)A). As expected, caspase-3 did not activate ICAD^{TevS}:CAD and neither did TEVP activate ICAD^{WT}: CAD, showing the orthogonality of the systems [\(Fig. S4](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF4)B). We next determined if the ICADL^{TevS}:CAD complex produced directly in HeLa cells could be activated by TEVP. Coimmuno-
precipitation of the ICADL^{TevS}:CAD complex from HeLa cells followed by TEVP treatment showed activity against both plasmid followed by $I E V F$ in called showed density form these results we and chromatin substrates [\(Fig. S4](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF4) C and D). From these results we concluded that active CAD can be generated from an $ICADL^T$ CAD complex by TEVP but not by caspase-3.

Phenotypic Effects of Selective Activation of CAD by iTevP. Following biochemical confirmation that the iTEVP system was optimized, we sought to understand the phenotypic consequences for specific activation of CAD in HeLa cells. Remarkably, we observed no significant change in cell viability or caspase activity upon doxycycline treatment for up to 48 h, as measured by Cell Titer Glo or Caspase-3/-7 Glo (Promega) (Fig. 4A and [Fig. S5](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF5)A). This came as an initial surprise to us because we believed that difficult to repair double-stranded DNA breaks would feedback and activate the apoptotic program. This finding indicated that a more detailed cellular characterization of the pathway downstream of CAD activation was required.

To determine the extent of cellular response to activation of CAD, we investigated the DNA damage-response pathway in greater detail, using ELISA-based detection strategies. Cytoplasmic nucleosomes were detected in cells treated with doxycycline (Cell Detection Plus, Roche) (Fig. 4B). Initially, immunoblotting for H2AX-Ser139 phosphorylation (H2AXP) as a marker for double-stranded DNA breaks, showed no change over an extended time course of treatment [\(Fig. S6\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF6). However, a sensitive ELISAbased detection system (Trevigen) revealed a marked increase in H2AXP levels following the addition of doxycycline in as little as 6 h (Fig. 4C).

To better understand the apparent discrepancy between the immunoblotting and ELISA-based detection methods, we hypothesized that the DNA damage signal might be coming from a subset of the cells exhibiting a strong response. To explore this idea, we looked for evidence of DNA damage and response at the single-cell level. Direct labeling of fragmented DNA by TUNEL staining showed only a small increase in TUNEL intensity of individual cells (Fig. $4D$ and E). The TUNEL assay was significantly less sensitive in detecting early apoptotic cells as determined by a side-by-side comparison with H2AXP staining when using positive control DNA damage-inducing agents ([Fig. S7\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=SF7). This is likely a result of the endogenous cellular amplification as the phosphorylation of H2AX spreads extensively around a site of DNA breakage while TUNEL directly labels a single DNA break (32, 33). High-content immunofluorescence quantitation of H2AXP levels showed clear enrichment after 12 h of doxycycline treatment over the vehicle control (Fig. $4F$). Although the mean values were relatively consistent over time, a log plot of individual cell intensity of H2AXP staining demonstrates a time-dependent increase in H2AX phosphorylation (Fig. 4H). This finding indicates that over time CAD continues to cleave chromatin, thereby causing increased H2AXP signaling and we presume genotoxic stress in the cell.

To directly monitor DNA fragmentation following iTEVP induction, we used a single-cell gel electrophoresis (comet) assay. We observed an increase in comet tail moment in a small subset of treated cells compared with cells without iTEVP activation. The increase in comet tail moment indicates damaged chromatin migration following electrophoresis (Fig. 4G). The tail moment of individual cells is plotted using a box-and-whisker plot for each time point (Fig. 4I). Whereas the median remains nearly constant across the time course, there is a clear increase in heavily damaged cells starting at 6 h after iTEVP activation.

From these assays, it was apparent that although DNA damage signaling was taking place, many of the cells were still surviving. Two hypotheses emerged to explain the lack of robust apoptosis in the HeLa human cancer line upon selective CAD

Fig. 3. Evidence of induced and selective cleavage of ICADL^{TevS}. (A) ICAD is a mandatory folding chaperone and inhibitor of CAD. ICAD tightly inhibits CAD activation until it is cleaved which leads to dimerization and activation of CAD. To selectively activate CAD we used an orthogonal activation allele that that serves as a folding chaperone and is cleavable by TEVP (arrowheads). (B) Different strategies for inducible TEVP were constructed to maximize ease of use, expression levels, and protease activity. A dual vector version resulted in minimal decrease in full-length substrate, tagged with the FLAG epitope. The tetracycline inducible full-length TEVP results in near complete full-length substrate cleavage within 12 h. (C) Combining PTGR ICADL^{TevS} and iTEVP in a stable cell line results in robust inducible expression of TEVP (Myc-tagged) and complete and sustained proteolysis of $\textsf{ICADL}^\mathsf{TevS}$

Fig. 4. Phenotypic effects of selective CAD activation by iTEVP. (A) Cell viability, measured by Cell Titer Glo (detected by luciferase signal absorption at 405 nm) following iTEVP activation by addition of doxycycline in PTGR ICAD-engineered cells. (B) Cytoplasmic enrichment of free histone measured by ELISA following activation of iTEVP. (C) Phosphorylation of H2AX monitored by ELISA. All error bars in A–C represent SD for three biological replicates. Significance was determined by one-way ANOVA Dunnett correction. (D) Quantitation of TUNEL staining at 24 and 48 h following iTEVP activation (1 µg/mL Doxycycline) (red bar, mean; blue, SD). (*E*) Representative images
from TUNEL staining following selective ICADL^{Tevs} cleavage. (*F*) Representative images from immunofluorescent staining for phosphorylation of serine-139 H2AX, following selective ICADL^{TevS} cleavage induced by addition of doxycycline. (G) Representative images of single cell gel electrophoresis (comet assay) following addition of doxycycline. (Scale bars in $E-G$, 100 μ m.) (H) Single cell high content image analysis of H2AXP levels plotted on a log scale; image intensity of entire population is shown (red bar, mean value; blue, SD; $n = 2,870$ cells). Significance was determined by one way ANOVA with Sidak correction. (I) Opencomet automated analysis of comet tail moment. Data are plotted on a Tukey boxplot ($n = 43$ cells per time point). * $P \le$ 0.05; *** $P \leq 0.001$.

induction: (i) DNA repair machinery was so robust that it was quickly able to repair CAD-mediated damage and (ii) chromatin was only accessible to CAD in a subset of cells, possibly because of changes in chromatin accessibility known to vary during the cell cycle.

Caspase substrate profiling, both in our laboratory and others, previously demonstrated that many proteins involved in DNA repair are indeed targets of caspases, such as poly(ADP-ribose) polymerase (PARP), DNA-PK, and ATM/ATR among others (16, 34, 35). Moreover, a number of proteins involved in maintaining chromatin's compact structure like the N-CoR/Smrt complex that contains histone deacetylase (HDAC) 3 and 7 are also targets (16).

Therefore, we tested if available small-molecule inhibitors of these processes would synergize with CAD activation to impact cell death, caspase activation, H2AXP, and TUNEL staining. We thus tested the effects of combining sublethal doses of each of these inhibitors in the context of CAD activation, including: the DNA-PK inhibitor, Nu7441; the ATM inhibitor, Ku60019; the HDAC3 specific inhibitor, RGFP966; and the broad-spectrum HDAC inhibitor, TSA. Indeed we found each inhibitor, to varying degrees, resulted in a small increase in caspase activation (Fig. 5A) and TUNEL staining (Fig. 5B). However, there was no significant change in viability between inhibitor $+$ doxycycline compared with inhibitor alone (Fig. $S5B$). There was a greater increase in phosphorylation of H2AX when cells were cotreated with doxycycline and various DNA pathway inhibitors (DNA repair and chromatin relaxation) (Fig. $5C$ and time-course detailed in Fig. $S8B$). These studies suggest that inhibiting a single arm of the DNA damage response synergizes with CAD activation, yet is still insufficient to trigger full-blown apoptosis.

We hypothesized that chromatin accessibility to CAD should peak during S-phase, because DNA is stripped of many of its chromatin binding proteins. To test this idea, we synchronized a population of cells in early S-phase using a double thymidine block and released them following a 3 h preactivation of iTEVP with addition of doxycycline. This regime induced approximately twice the number of apoptotic cells, compared with synchronized cells in the absence of doxycycline as assessed by staining with annexin V and propidium iodide (PI) (Fig. 5E).

To assess the consequence of extended CAD activation in different stages of the cell cycle, we tracked the percentage of cells by PI staining using flow cytometry. We observed a marked arrest of cells in G_2/M phase, a doubling of the percentage of cells $(21-41\%)$ in G_2/M compared with cells without doxycycline (Fig. 5F). The arrest in G_2/M was stable out to 72 h after addition of doxycycline. Overall, our data suggest that direct activation of CAD does lead to DNA cleavage, especially in S-phase, and that the DNA repair process once initiated can efficiently antagonize CAD activation.

Discussion

DNA fragmentation of is a classic hallmark of apoptosis. A key event in this process is believed to be the activation of CAD by caspase cleavage of its inhibitor and folding chaperone, ICAD. We engineered an inducible and selective proteolysis system to closely mimic caspase proteolysis of ICAD without parallel cleavage of over 1,500 other caspase substrates. The system preserved the critical role of ICAD as a CAD folding chaperone, by replacing the endogenous ICAD at physiological levels with one sensitive to TEVP. This process was enabled by designing the replacement ICAD allele to be resistant to the expressed synthetic miRNA, allowing the endogenous ICAD to be knocked down while the exogenous ICAD^{TevS} rescued endogenous function. This posttranscriptional gene-replacement strategy is simple, rapid to use, and genetically encoded. Recent advances in genome engineering using transcription activator-like endonucleases and CRISPR show great promise for gene ablation studies; however, the ability to use these techniques for site-specific biallelic replacement is not guaranteed and requires a large investment in reagents and screening time (36). We believe the PTGR strategy may be more broadly appropriate to many problems where one wishes to simultaneously replace endogenous biallelic expression with an engineered allele variant and screen for emergent functions.

We investigated two systems for inducing targeted proteolysis by TEVP, a split protein and full-length version. Although the SNIPer was able to cleave ICAD^{TevS}, we found the rate of proteolysis to be low and insufficient to deal with the steady-state
levels of ICAD^{TevS}, probably because synthesis of new ICAD^{TevS} kept pace with split-TEVP proteolysis. Thus, we investigated a
TetON full-length TEVP system for cleaving ICAD^{TevS}. Although there is a lag phase in TEVP activity as a result of transcriptional/ translational delay in producing the full-length TEVP, the intact TEVP is about five-times more active than the split construct.

Fig. 5. Pharmacological sensitization of DNA damage with selective CAD activation. (A) Fold-change of caspase activity in engineered cells cotreated with doxycycline and DNA repair or HDAC inhibitors for 24 h. Significance determined by unpaired t test. (B) Single-cell quantitation of TUNEL staining with pharmacological inhibitors and cotreatments with 1 μg/mL doxycycline (48 h). The following inhibitors were used in treatments for samples for TUNEL and H2AXP staining: 1 μM Nu7441 for DNA-PK; 3 μM Ku60019 for ATM; 10 μM RGFP966 for HDAC3; 400 nM Trichostatin A (TSA) for HDAC. (C) Single-cell quantitation of H2AXP levels following treatment with 1 μg/mL doxycycline for 12 h. Significance determined by unpaired t test. (D) Single-cell quantitation of H2AXP levels with cotreatments with DNA repair inhibitors (Nu7441 and Ku60019) and doxycycline (12 h). Significance determined by unpaired t test. (E) Single-cell quantitation of H2AXP levels cotreated with HDAC inhibitors (RGFP966 and TSA) and doxycycline (12 h). Colored bars represents mean value and error bars are SD. Significance determined by unpaired t test. (F) Flow cytometry based assessment of apoptotic cells by PI/annexin V staining following S-phase synchronization induced by double thymidine block followed by 24-h treatment with doxycycline. Doxycycline-dependent increase is observed in synchronized cells from 2.3 to 5.0% annexin V positive and 8.3 to 18.5% annexin V and PI double-positive cells. (G) Cell cycle distribution of cells as measured by PI staining using flow cytometry. PTGR ICADL^{TevS} cells were treated for an extended time with doxycycline to activate CAD. * $P \le 0.05$; $**P \le 0.01$; $***P \le 0.001$; $****P \le 0.0001$.

Thus, we favored the iTEVP for the loss-of-function studies where rapid induction is not as important as complete proteolysis.

TEVP was capable of cleaving ICAD^{TevS} both in vitro and in cells and trigger hallmarks of DNA damage, such as H2AX phosphorylation. Chromatin fragmentation also significantly increased following selective ICAD^{TevS} cleavage, as demonstrated by increases in TUNEL staining, nucleosome release, and DNA tail moment in comet assays. Remarkably, we detected no significant change in cell death as measured by cellular ATP levels (Cell Titer-Glo) or caspase activity (Caspase-Glo).

In aggregate, our data suggest that DNA repair can largely out-compete DNA cleavage by CAD and preserve the cell. Among the many substrates that caspases cleave are proteins involved in DNA repair processes (Fig. 6). Indeed we observed increases in all DNA damage reporters upon pharmacological inhibition of DNA repair pathways. The accumulation of DNA damage in S-phase or with HDAC inhibition further suggests that exposed chromatin is more sensitive to CAD. Thus, our data suggest that the combined action of multiple caspase cleavages, DNA repair, and chromatinmodifying enzymes, work in coordination with CAD activation to facilitate DNA fragmentation in human cells during apoptosis.

There have been reports that selective CAD activation can lead to DNA fragmentation and cell death in limited contexts, yet to our knowledge, our report is the first to achieve in situ activation in mammalian cells. Previous studies demonstrated the ability to activate CAD via addition of TEVP to ICAD^{TevS}/CAD in cytosolic extracts and utilization of a galactose-inducible TEVP in Saccharomyces cerevisiae (37, 38). Because yeast lack a traditional apoptotic pathway, caspases, and the ICAD/CAD node, the galactose inducible TEVP strategy only proved useful to study chromatin structure in this context. More recently, Samejima et al. described a cellular strategy that relied on an auxin-inducible degron system for inducing rapid and complete proteasomal degradation of huICAD in double-knockout DT40 chicken cells (39). It remains unclear how authentic their strategy is for selective CAD activation because it does not recapitulate both the spatial and endogenous site-specific mechanism of activation. Our approach of site-specific proteolysis closely mimics the endogenous mechanism. Our data are consistent with the view that apoptosis results from the synthetic lethality set up by coordinated caspase cleavage of multiple target pathways. Such a system provides useful safeguards for inadvertent triggering of cell death from a single node.

We believe the strategy of substrate replacement coupled with inducible and selective proteolysis provides a highly generalizable approach for assigning the functional consequences of selective cleavage of caspase substrates in cells. In addition, we envision the approach to be beneficial in characterizing the cross-talk between phosphorylation and apoptotic proteolysis by including either phosphorylation mimetic or nonphosphorylatable residues in the TevS allele series (40, 41). The approach outlined with the PTGR and SNIPer technology may prove particularly beneficial in profiling site-specific proteolysis throughout cell biology: for example, during cell-cycle progression mediated by separase during anaphase (42), differentiation, and intercellular signaling by Notch proteolysis (43), and mediating the unfolded protein response by ATF6 proteolysis (44).

As the proteomic-generated map of caspase substrates continues to grow, our understanding of the functional consequence resulting from individual cleavage events remains in its infancy. The present studies offer a strategy to precisely investigate substrate proteolysis and can provide new insight to the complexity and cooperativity between caspase cleavage events during apoptosis.

Fig. 6. Summary model of synthetic lethality and pharmacological sensitization through simultaneous activation CAD and inhibition of DNA repair. CAD activation through caspase or TEVP cleavage cleaves exposed regions of DNA (Center), which is opposed by either DNA repair (Right) or chromatin compaction (Left) promoted by HDACs. These opposing processes can be targeted by additional caspase cleavages (shown in red) or by pharmacological inhibitors (shown in light brown).

Materials and Methods

SNIPER, iTEVP, and the PTGR backbone was assembled from cassettes obtained from Kaveh Ashrafi (University of California, San Francisco, CA) and Addgene. PTGR- and iTEVP-engineered HeLa cells were plated in 96-well microclear plates (Greiner). Cells were allowed to attach and were treated for the indicated amount of time with the indicated small molecules: doxycycline (Sigma), Nu7441 (Torcis), Ku6009 (Torcis), RGFP966 (Selleckchem), TSA (Cells Signaling). Cells were fixed and permeabilized with 1% PFA in PBS +0.1% Triton X-100 for 10 min. H2AXP immunofluorescence protocol was developed from Zeitlin 2009 (45). High-content image analysis included cell identification by nuclear staining (DAPI), and segmentation was refined by watershed clump breaking and sieve algorithms to eliminate false targets. Using DAPI signal as a threshold area, H2AXP signal was recorded for each target cell in a field of view. Batch processing was then used to analyze

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>2,000 cells per well, across a 96-well plate. Target and pixel intensities were then exported to Microsoft Excel for numerical review and final graphical representation and statistical analysis using Prism (Graphpad). Statistics were performed using Prism as indicated in the figure legends. Additional experimental details are described in [SI Materials and Methods](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=STXT). See [Table S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1504141112/-/DCSupplemental/pnas.201504141SI.pdf?targetid=nameddest=ST1) for selected primer sequences used.

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