

Massively parallel, computationally guided design of a proenzyme

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Confining the activity of a designed protein to a specific microenvironment would have broad-ranging applications, such as enabling cell type-specific therapeutic action by enzymes while avoiding off-target effects. While many natural enzymes are synthesized as inactive zymogens that can be activated by proteolysis, it has been challenging to redesign any chosen enzyme to be similarly stimulus responsive. Here, we develop a massively parallel computational design, screening, and next-generation sequencing-based approach for proenzyme design. For a model system, we employ carboxypeptidase G2 (CPG2), a clinically approved enzyme that has applications in both the treatment of cancer and controlling drug toxicity. Detailed kinetic characterization of the most effectively designed variants shows that they are inhibited by ~80% compared to the unmodified protein, and their activity is fully restored following incubation with site-specific proteases. Introducing disulfide bonds between the pro- and catalytic domains based on the design models increases the degree of inhibition to 98% but decreases the degree of restoration of activity by proteolysis. A selected disulfide-containing proenzyme exhibits significantly lower activity relative to the fully activated enzyme when evaluated in cell culture. Structural and thermodynamic characterization provides detailed insights into the prodomain binding and inhibition mechanisms. The described methodology is general and could enable the design of a variety of proproteins with precise spatial regulation.

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Autoinhibited enzymes, known as proenzymes or zymogens, are used ubiquitously for the precise regulation of key biological processes (1). For example, the activities of blood clotting enzymes, digestive proteases, regulatory proteases, apoptotic proteases, and complement system proteins can be triggered by relevant environmental stimuli. Autoinhibiton is typically achieved by extending the active protein with a prosegment or prodomain that inhibits enzymatic activity or prevents adoption of the active conformation until it is cleaved by a tissue-specific activating protease that selectively recognizes a cleavage sequence in the linker connecting the prodomain to the active protein. To enable tissue specificity for an exogenously administered therapeutic or diagnostic enzyme (2), a general approach for redesigning enzymes into activatable, zymogen variants would be valuable. Surveys of naturally occurring zymogens have revealed that prodomains have diverse lengths and folds and exert their inhibitory effects by steric shielding of the substrate binding sites, or in the case of metalloproteases, by additionally coordinating active site metal ions (1, 3, 4). There have been successful efforts to design proenzymes (5-8), but they have relied on unique structural features of the enzyme under consideration. For example, the proximity of N and C termini in ribonuclease A allowed for building an inhibiting linker through the active site (5, 6, 8), while barnase allowed for an alternate frame folding-based design (7) which required proper folding of a chimeric construct containing a duplicated C-terminal segment. Furthermore, like most natural proenzymes, these designed zymogens have been limited to enzymes that target macromolecular substrates (5-8), rather than small molecule substrates that are much more difficult to sterically impede, and thus represent a significantly greater design challenge. The development of a generalizable, structure-guided approach for zymogen design, particularly one that could be applied to enzymes that target small molecule substrates, would allow spatial control over a broad range of enzymes.

In order to achieve an effective means of localizing the activity of any chosen enzyme to a target microenvironment, we envisioned the addition of an inhibitory prodomain that would be removable by specific proteases which are markers of the target tissue (Fig. 1*A*). For example, matrix metalloproteases (MMPs) are highly expressed in tumors and would facilitate targeting of enzymatic activity to a tumor site. Tumor-specific MMP activity has been employed previously to achieve site-selective activation of prodrugs (9, 10) and imaging contrast agents (11). We selected the Food and Drug Administration

Significance

Proteins have shown promise as therapeutics and diagnostics, but their effectiveness is limited by our inability to spatially target their activity. To overcome this limitation, we developed a computationally guided method to design inactive proenzymes or zymogens, which are activated through cleavage by a protease. Since proteases are differentially expressed in various tissues and disease states, including cancer, these proenzymes could be targeted to the desired microenvironment. We tested our method on the therapeutically relevant protein carboxypeptidase G2 (CPG2). We designed Pro-CPG2s that are inhibited by 80 to 98% and are partially to fully reactivatable following protease treatment. The developed methodology, with further refinements, could pave the way for routinely designing proteaseactivated protein-based therapeutics and diagnostics that act in a spatially controlled manner.

The authors declare no competing interest.

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Fig. 1. Concept and design strategy for protease-activatable forms of CPG2. (*A*) Mechanism of activation of Pro-CPG2. A protease (yellow) cleaves the linker that connects the prodomain (blue) to the inactive enzyme (red), activating it (green). The enzyme can then process small molecules (teal and purple) which can have a biological effect. (*B*) Activated CPG2 can cleave and detoxify the drug methotrexate (MTX), rescuing the MTX-sensitive cells (black to orange). (*C*) The Pro-CPG2 design strategy ensured incorporation of a zinc (gray sphere)-coordinating glutamate (sticks representation) on the designed prodomain (blue). The CPG2 scaffold is shown in wheat. (*D*) The design strategy ensured that the N terminus of the prodomain is sufficiently close to the C terminus of the circular permuted enzyme to incorporate a short linker containing a protease cleavage sequence (yellow scissors). (*E*) Successful designs were tested for incorporation of a prodomain-scaffold disulfide bond, designed by Rosetta. All structural figures are Rosetta design models.

(FDA)-approved enzyme carboxypeptidase G2 (CPG2) as the target for zymogenization. CPG2 is a well-studied and functionally versatile enzyme with a variety of therapeutically beneficial activities (SI Appendix, Fig. S1). Specifically, CPG2 efficiently inactivates the anticancer drug methotrexate to reduce systemic toxicity (Fig. 1B, SI Appendix, Fig. S1B) (12-15), for which it has received FDA approval. Moreover, CPG2 activates nontoxic nitrogen mustard prodrugs, such as N-{4-[(2-chloroethyl)(2mesyloxyethyl)amino]benzoyl}-L-glutamic acid, to their active, toxic forms in directed enzyme prodrug therapy (DEPT) (SI Appendix, Fig. S1C) (16, 17). A strategy that exploits the natural properties of tumors to express elevated levels of specific proteases (18, 19) would be broadly applicable for tissue-specific drug delivery. Designed activatable forms of CPG2 which respond to tissue-specific proteases could therefore be useful diagnostic and therapeutic reagents for both delivering and ablating toxicity in a tissue-specific manner (20).

Here, we describe a computationally driven protein design approach to develop protease-activatable zymogens and employ this strategy to develop proenzyme forms of CPG2. Combining next-generation sequencing (NGS) and a computational design approach implemented in the Rosetta software suite, we have developed a massively parallel enzyme design strategy to screen thousands of computationally designed pro-CPG2 variants in parallel. Using this approach, we designed artificial proenzymes that target a small molecule substrate. We used small-angle X-ray scattering (SAXS) to elucidate the structural basis of inhibition, demonstrating that key domain movements are critical for the catalytic activity of CPG2. A designed, disulfide-linked pro-CPG2 variant significantly reduces the protection of the PC-3 human prostate cancer cell line from methotrexateinduced cytotoxicity relative to the uninhibited variant. These CPG2 variants are promising starting points for further optimization for anticancer therapeutic development, and the design methodology may also enable facile zymogenization of other enzymes with therapeutic value.

Results

Overview of Design Strategy. In natural zymogens, prodomains are fused by a protease-cleavable short linker to one of the protein termini. Inspection of the wild-type CPG2 crystal structure (21) reveals that both termini are >30 Å distant from its active

site. We previously designed active circular permutations of CPG2 (CPG2_{CP-N89}) with termini proximate to the active site, creating the possibility of attaching a terminal prodomain that can access and effectively inhibit the active site (22). The CPG2_{CP-N89} variant (in which the polypeptide chain is reorganized such that the N-terminal residue is wild-type residue 89) served as the starting point for our design efforts. As CPG2 is a zinc-containing metalloenzyme, we reasoned that using a zinc-coordinating sidechain in the prodomain, such as glutamate, to guide prodomain placement toward the catalytically important zinc ion would effectively block the core active site region when the prodomain is bound. We thus envisioned appending prodomains to either terminus such that both a metal-chelating residue and a chosen activating protease cleavage site sequence in the connecting linker are incorporated (Fig. 1 C and D).

Helical Prodomain Design. As an initial proof of concept for the autoinhibition of CPG2, we designed a series of proenzymes featuring a short helical prodomain. We used Rosetta to dock a 12or 17-residue helix into the active site of the CPG2_{CP-N89} circular permutant (22) while allowing substitutions on both the prodomain and CPG2 sides of the interface. The resulting poses were then subjected to loop closure, fusing the helix to either terminus using a linker containing the tobacco etch virus (TEV) protease cleavage sequence. We identified multiple designs in which a Zn²⁺-coordinating residue could be added such that it would interact with the catalytic dinuclear zinc site. When considering the mutations introduced in CPG2, a single substitution, K177A (wild-type residue numbering), was present in many of the designs and appeared to decrease steric restrictions in the formation of prodomain-enzyme interactions. This substitution only resulted in a ~20% loss of activity relative to the CPG2_{CP-N89} circular permutant (SI Appendix, Fig. S2D), which we have previously demonstrated has equivalent activity to the wild-type enzyme (22). Eight designs were selected for activity characterization with either the original TEV cleavage sequence or by substituting the TEV sequence with the MMP-2 cleavage sequence, and two designs (SI Appendix, Fig. S2 B, G, and H and Table S1) were experimentally determined to be modestly autoinhibited (SI Appendix, Fig. S2). Incubation of proenzyme with the relevant protease for 24 to 48 h resulted in an enzyme that regained comparable activity to the uninhibited scaffold, and reversion of the K177A mutation resulted in a smaller degree of inhibition when comparing the proenzyme and the base variant while only moderately impacting base variant activity. These findings validate the design of a true proenzyme that is both inhibited and reactivatable, although the degree of inhibition is modest.

To gain insight into the structural basis for the observed inhibition, we solved the crystal structure of one of the designed proenzymes (Protein Data Bank identifier [PDB ID] 7M6U; *SI Appendix*, Table S2) and compared it to the structure of wildtype CPG2 (PDB ID 1CG2). We could not locate the density for the appended helical prodomain in the structure, a finding that is consistent with the weak inhibition observed for this design. Nonetheless, we were able to determine that this variant maintains the overall tertiary structure of the enzyme (*SI Appendix*, Fig. S3, C α RMSD of 0.26 to 0.52 Å when comparing all 1CG2 chains to all 7M6U chains), indicating that the circular permutation does not perturb the structure or function of CPG2, as expected from our previous results (22).

Massively Parallel Pro-CPG2 Computational Design. Building on the modest, proof-of-principle inhibition achieved in the helical pro-CPG2 designs, we pursued a massively parallel design approach to develop more strongly inhibiting prodomains. Three characteristics present in the initial designs were used to guide our approach. First, we included an interaction between the catalytic zinc ion and a glutamate residue on the prodomain as the "hotspot" interaction (Fig. 1*C*). This was achieved by directing prodomain placement using a library of free glutamate conformations that coordinate the zinc residue; backbone coordinates of these glutamate conformations were used to position the prodomain. Second, only C-terminal prodomains were considered, as they proved to be more effective in our initial designs (Fig. 1*D*, *SI Appendix*, Fig. S2*B*). Finally, the K177A mutation, which diminished the enzyme's activity by ~20% (*SI Appendix*, Fig. S2*D*) while providing the benefit of decreased steric restrictions at the expected prodomain binding site, was retained in all designs. A set of 3,157 prodomain scaffolds was generated by combining small domains curated from the PDB and from a large set of previously reported de novo designed "miniproteins" (23) (also see *Methods*). Candidate prodomains feature a wide range of folds and sequences, including alpha-helical bundles and mixed alpha/beta folds. The prodomain library can be downloaded at Zenodo (https:// zenodo.org/record/5553581).

To minimally perturb the activity of the postcleavage form of the enzyme, we used a one-sided design approach in which only the prodomain sequence could be modified while the sequence of the CPG2 scaffold region remained fixed. The interaction surface for the prodomain is not a typical natural protein–protein interaction site, as it includes a significant number of polar and charged moieties such as the dinuclear zinc ion-containing active site. To introduce interactions with these polar groups, we designed explicit hydrogen bond networks between the prodomain and the enzyme scaffold.

A total of ~15,000 designs was obtained using a design workflow implemented in Rosetta (Fig. 2*A*; also see *Methods*). We used a number of criteria (detailed description in *SI Appendix*, Supplementary Results S1, Fig. S4, and Table S3) to select



Fig. 2. Massively parallel design of Pro-CPG2. (*A*) The design strategy is depicted as a flowchart. Abbreviations: AAs (amino acids) and BB (backbone). (*B–D*) Rosetta models of the three Pro-CPG2 designs selected for detailed study: (*B*) Pro-CPG2-1, (*C*) Pro-CPG2-2, (*D*) Pro-CPG2-3. The prodomain (green), catalytic domain (blue), and dimerization domain (wheat) are highlighted. (*E*) NGS results from the massively parallel selection assay. The relative enrichment ratio (log scale) is given at increasing MTX concentrations. The helical designs that are known not to inhibit (green) or to mildly inhibit (orange) CPG2_{CP-N89} are shown alongside the 12 designs predicted to be Pro-CPG2s (red) and predicted not to be Pro-CPG2s (blue).

 \sim 7,500 designs for testing using a high-throughput assay (as described in the next section). As controls, we included four helical prodomain designs (one ineffective and three moderately effective autoinhibitors) that allow the comparison of the relative activities determined in our high-throughput assay to those of prodomains with known inhibitory effects.

Massively Parallel Screening of the Pro-CPG2 Design Pool. We previously reported that periplasmically expressed CPG2 is able to protect Escherichia coli CAG12184 cells from methotrexate toxicity (22, 24-26). We employed this screen to discriminate between inhibiting and noninhibiting pro-CPG2 designs from the set of 7,469 computationally generated designs. In summary, cells expressing designs with poorly inhibiting prodomains (i.e., full CPG2 activity) detoxify methotrexate effectively and survive, while cells expressing designs with strongly inhibiting prodomains (i.e., low CPG2 activity) are unable to detoxify methotrexate and do not survive. We generated a relative ranking of prodomain efficacy by comparing the fraction of NGS reads attributable to each design in cultures grown at various methotrexate concentrations (30, 50, and 100 µM) and comparing them to cultures grown in the absence of methotrexate. By including both moderately effective and ineffective control prodomains identified from our original helical pro-CPG2 designs, the NGS results could be analyzed to identify designs that are expected to be at least as strongly inhibited as the helical pro-CPG2s (SI Appendix, Fig. S2 and Table S4). Between 400,000 and 1.3 million NGS reads were obtained for each individual test culture corresponding to methotrexate concentrations of 0 to 100 µM, suggesting adequate coverage of the 7,500 unique sequences contained in the library.

Upon analysis of the NGS data, reliable enrichment values could be obtained for 1,384 out of the tested sequence pool of 7,500 sequences, allowing us to identify 107 inhibited designs and 27 high-probability inhibited designs, with the latter defined as designs whose relative enrichment was lower in the 30 μ M methotrexate culture as compared to the 0 μ M culture and lower in both the 50 µM and 100 µM methotrexate cultures relative to the 30 µM culture (also see SI Appendix, Supplementary Methods). To confirm the ability of this highthroughput assay to effectively discriminate between inhibiting and noninhibiting designs, we selected 12 of these 27 designs, as well as 4 designs identified to be noninhibiting (Fig. 2E and SI Appendix, Fig. S5). The 12 designs selected for further study were chosen to maximize sequence diversity and eliminate designs with very similar predicted folds and binding modes. The diversity within the set of inhibiting designs suggests that our selection is not biased toward a particular fold. Each of these 16 designs was purified as an individual construct alongside the most inhibited helical pro-CPG2 design and the CPG2_{CP-N89}-K177A construct. All 12 of the predicted inhibiting pro-CPG2 designs were more strongly inhibited than the helical pro-CPG2 design, while the 4 predicted noninhibiting pro-CPG2 designs exhibited activities resembling CPG2_{CP-N89}-K177A (SI Appendix, Fig. S5). These results confirm that our screening system accurately discriminates between inhibiting and noninhibiting designs, using the helical pro-CPG2 controls as benchmarks. Of the 12 designs, we identified 3 for further characterization, which we refer to hereafter as Pro-CPG2-1, Pro-CPG2-2, and Pro-CPG2-3 (SI Appendix, Table S1). The models of the Pro-CPG2-1 and -2 prodomains have mixed alpha/beta folds, whereas Pro-CPG2-3 is a helical bundle.

The Michaelis-Menten kinetic parameters of Pro-CPG2-1, -2, and -3 were determined and compared to the fully active

CPG2_{CP-N89}-K177A (Fig. 3 *A*–*C* and *SI Appendix*, Table S5). Significantly, there is an approximate 50% reduction in the catalytic rate constant (k_{cat}) for all three designs, indicating a dramatic decrease in the ability of the enzyme to complete its catalytic cycle. In addition, Pro-CPG2-1 and -2 elicited a twofold to threefold increase in the Michaelis constant (K_M), suggesting that the effective substrate binding affinity is reduced. These three designs exhibit up to an 80% decrease in catalytic efficiency, k_{cat}/K_M .

Analysis of Inhibited vs. Not Inhibited Designs. Based on the high predictive power of our screening approach, we sought to compare properties of designs identified by our assay to be inhibitors to those identified not to be inhibitors. We performed two-sided t tests comparing the two groups of designs for all computed statistics (selected metrics shown in SI Appendix, Table S6). Of note, the number of carbon-carbon contacts between the prodomain and two regions of interest were found to be significantly greater in the inhibited designs, namely, the base of the dimerization domain beta-sheet (residues 215 to 219 and 320 to 324) and a catalytic domain loop that contains the K177A residue previously highlighted in our helical designs and spatially proximal to the circular permutant's termini (residues 173 to 182) (SI Appendix, Fig. S6). These observations suggest that forming contacts with these regions may be important for achieving sufficiently tight binding and inhibition. Upweighting designs with contacts in these regions may yield higher success rates or stronger inhibition in future studies.

Recovery of Activity using Specific Proteases. In order to function as switches, zymogens must be reactivated by proteolytic processing using a specific protease. To probe the activatability of the pro-CPG2 designs, we generated interdomain linker variants containing the canonical cleavage sequences of two dissimilar proteases, namely, the 3C-type TEV cysteine protease (27) and zinc-dependent matrix metalloproteinase MMP-2 (28). The activity of all Pro-CPG2 designs is recovered following incubation with the appropriate protease (Fig. 3 D and E). As demonstrated by mass spectroscopy, both the residual uncleaved proenzyme and retention of the bound but cleaved prodomain (when free prodomain is removed through dialysis) may be responsible for the incomplete recovery of CPG2 activity (SI Appendix, Fig. S7). As two unrelated proteases are able to cleave and reactivate the corresponding CPG2 proenzyme, it is likely that a broad range of other site-selective proteases could be used to reactivate pro-CPG2 by including the appropriate cleavage site in the interdomain linker.

Characterization of the Designed Prodomains. As an initial computational characterization of the designed proenzymes, we performed a series of simulations to assess their quality (*SI Appendix*, Supplementary Results S2 and Fig. S8). Ab initio forward folding successfully refolds the designs both as isolated proteins and in the context of CPG2, suggesting that the appended domains are well-folded in the designed conformations.

To further investigate the prodomains as independent proteins, the prodomains of Pro-CPG2-1, -2, and -3 were expressed individually and their secondary structure features analyzed using circular dichroism (CD) spectroscopy. All three prodomains exhibit CD spectra characteristic of well-folded proteins that are stable at physiologically relevant temperatures. Consistent with the Rosetta models, Pro-CPG2-1 and -2 have characteristic mixed alpha/beta spectra, while the spectrum of Pro-CPG2-3 appears primarily alpha-helical in character.



Fig. 3. Activity and thermodynamic data for the Pro-CPG2 designs and their disulfide variants. (*A*–*C*) The Michaelis–Menten kinetics (*A*, k_{cat} ; *B*, K_M ; *C*, k_{cat}/K_M) for the three base Pro-CPG2 designs are shown. In yellow, the disulfide variants are grouped with the base design they are derived from. See also *SI Appendix*, Table S5. (*D*–*E*) The recovery of molar specific activity following incubation with MMP-2 (*D*) and TEV (*E*) protease is shown. Differences in absolute molar specific activity between *D* and *E* are due to different buffer conditions required for MMP-2 and TEV activity. (*F*) Temperature-dependent thermodynamic binding signatures characterizing the association of CPG2_{CP-N89}-K177A and Pro-Domain-1 deduced from ITC measurements. See also *SI Appendix*, Table S7. (*G*) Temperature-dependent ITC-derived K_d of CPG2_{CP-N89}-K177A and Pro-Domain-1. Error bars in panels *A*–*C*, *F*, and *G* are standard errors obtained from curve fitting. Error bars in panel *D*–*E* are standard deviations from replicate measurements. See also *SI Appendix*, Table S7.

Inspection of thermal melting profiles reveals relatively broad noncooperative unfolding transitions that do not plateau even when heated to 100 °C, thereby precluding an accurate determination of a discrete melting temperature (*SI Appendix*, Fig. S9). The high stability of these constructs suggests that they are likely to adopt the same, well-structured fold when fused to the terminus of CPG2_{CP-N89}-K177A.

Next, we assessed the binding affinity of Pro-Domain-1 for the CPG2 scaffold in trans. We conducted isothermal titration calorimetry (ITC) experiments in which Pro-Domain-1 is titrated successively into CPG2_{CP-N89}-K177A. The acquisition of a series of binding profiles at varying Pro-Domain (150 to 500 μ M) and enzyme (15 to 50 μ M) concentrations yielded a consistent set of thermodynamic parameters, thereby ensuring that aggregation or precipitation (at higher concentrations) or dissociation (at lower concentrations) do not appreciably impact the binding energetics. Binding isotherms acquired at 5.0 °C reveal that the interaction heats are relatively modest and endothermic. Moreover, an analysis of these association profiles indicates that the formation of the CPG2_{CP-N89}–Pro-Domain-1 complex is characterized by binding

affinities on the order of 105 M-1 (i.e., dissociation constanty $[K_d] \sim 3$ to 6 μ M) over the temperature range studied. Inspection of the thermodynamic binding parameters (Fig. 3 F and G and SI Appendix, Table S7) acquired at 5.0 °C reveals that the Pro-Domain-enzyme association is entropy driven, as manifested by the relative contributions of the binding enthalpy (ΔH , 1.5 kcal·mol⁻¹) and the binding entropy (T Δ S, 8.1 kcal·mol⁻¹) to the binding Gibbs Free Energy ($\Delta G = -6.6 \text{ kcal} \cdot \text{mol}^{-1}$). Significantly, the interaction is accompanied by a slightly negative heat capacity ($\Delta C_p \sim -0.098 \text{ kcal} \cdot \text{mol}^{-1} \cdot \text{deg}^{-1}$) as reflected in the temperature dependence of binding enthalpies measured at 5 and 15 °C. Based on these results, we anticipate that the enthalpic term approaches zero near 25 °C and is presumably slightly exothermic under selection, assay, and physiological conditions. The observed temperature dependence is a characteristic hallmark of hydrophobic interactions (29-31) that presumably reflects the desolvation and burial of hydrophobic surfaces, potentially including the methotrexate binding site, upon prodomain binding.

Examination of the binding interface in our model reveals several nonpolar contact regions; 766 Å² of the 1,117 Å² buried solvent accessible surface area calculated using Rosetta is hydrophobic. The discrete surface burial, particularly in terms of hydrophobic residues, is generally consistent with a modest heat capacity change estimated for the interaction and may provide further support for our design model. Given that the binding interface includes a metal coordinating glutamate from the prodomain, it is entirely plausible that water molecules may be dislodged upon association. The zinc-glutamate coordination might therefore contribute additional unfavorable enthalpic contributions as previously reported for protein-metal interactions (32), resulting in an overall net entropy-driven association. An evaluation of the interdependence of the CPG2 prodomain binding sites is included in the SI Appendix, Supplementary Results S3.

An Open-Closed Equilibrium Associated with CPG2 Activity Revealed by SAXS. In the absence of a crystal structure where the binding mode of our designed prodomains could be identified, we sought to use SAXS experiments to understand the structural states occupied by our designs as compared to the uninhibited enzyme (Fig. 4). We hypothesized that, similar to other members of the M20 family of metallodipeptidases, CPG2 exists in the following two distinct conformations: an open conformation responsible for substrate binding and a closed conformation optimized for catalytic turnover (Fig. 4E). The former conformation is observed in the crystal structure of CPG2 reported here (SI Appendix, Fig. S3), as well as a previously published structure of CPG2 (21). The latter conformation is observed in many closely related homologs such as mouse carnosinase 2 (mCN2) (33), Sapep (34), PepV (35), peptidase T (36), aminoacylase-1 (37), and N-succinyl-L,L-diaminopimelic acid desuccinylase (DapE) (38). Anticipating that the addition of excess product (which is converted entirely from the substrate by the enzyme prior to data collection) would push the conformational equilibrium to the closed conformation, we acquired SAXS data in the presence and absence of hydrolyzed methotrexate to monitor changes in the accessibility of these two states. A model of the closed state was generated by individually superimposing the CPG2 catalytic and dimerization domains onto the structure of mCN2 and using generalized kinematic loop closure (39) to rebuild the interdomain linkers.

An analysis of the SAXS data (Fig. 4*A*, *SI Appendix*, Table S8) for CPG2_{CP-N89}-K177A in the absence of the products resulted in a particle with a radius of gyration (R_p) of 38.9 Å.

While neither the open crystal structure nor the closed model produced a reasonable fit of the data alone ($\chi^2 > 25$), we observed a good fit ($\chi^2 = 2.2$) for a two-state model using FoXS (40). The two states (open and closed) each contribute roughly 50% of the total scattering (Fig. 4B). In contrast, when hydrolyzed methotrexate is added to the sample, one obtains a smaller $R_{\rm g}$ of 36.0 Å which is consistent with the more compact closed model (Fig. 4A). The closed model produced a reasonable fit to the data using CRYSOL (41) ($\chi^2 = 4.9$), although a moderately better fit is obtained with the two-state FoXS model ($\chi^2 = 2.3$), with 75% and 25% contributions from the closed and open conformations, respectively (Fig. 4B). These data support our hypothesis and are consistent with the mechanisms of homologous enzymes (33-38), namely, that the addition of the product triggers a shift in the conformational equilibrium toward the closed conformation of CPG2.

We conducted a similar analysis on Pro-CPG2-1, -2, and -3 (Fig. 4 and SI Appendix, Table S8), using Rosetta-generated models of open and closed states of these proteins. In all three cases, the R_gs of samples containing substrate are reduced when compared to samples in the absence of product; however, the magnitudes of these changes are smaller than those for CPG2_{CP-} N89-K177A, primarily owing to a smaller observed Rg for samples without product (Fig. 4A). When two-state models of the Pro-CPG2s are used to fit the scattering data, the conformational equilibrium is $\sim 50\%/50\%$ open/closed, as observed for CPG2_{CP-} _{N89}-K177A, ranging between 48 to 59% in the open conformation. In contrast, the conformational equilibrium does not shift with the addition of product, remaining at roughly 50%/50% levels for the proenzymes (Fig. 4B). It should be noted that while the precise percentages of populations of the two states are highly dependent on the quality of the models used for fitting the data and should not be interpreted in absolute terms, the lack of a shift toward the closed conformation in relative terms is consistent, regardless of the models used. Thus, the SAXS-based characterization of all Pro-CPG2 variants is consistent with a scenario in which the open state is preferentially stabilized by the designed inhibitory domain.

To further corroborate our findings and investigate the impact of bias introduced by the quality of both the CPG2_{CP-N89}-K177A and proenzyme models, we utilized model-free analyses of the SAXS data, including dimensionless Kratky (42, 43) and Porod (42) exponent analyses (*SI Appendix*, Supplementary Results S4 and Fig. 4). These analyses confirm that in the absence of product, CPG2_{CP-N89}-K177A behaves as a well-folded but flexible particle. While the addition of the product renders this particle less flexible, the proenzymes' flexibility is even further reduced, with the concentration of mass occurring closer to the particle's center of mass. These results are therefore consistent with our initial findings that the proenzymes exhibit reduced conformational flexibility as compared to the circular permutant, with a greater population adopting the inactive, open state.

Effects of Disulfide Stapling on Pro-CPG2. To validate our structural models of Pro-CPG2 and to establish an "upper bound" of enzyme inhibition using our current designs, we generated disulfide-stapled variants of Pro-CPG2-1 and -2 with the goal of restricting the prodomain to its designed open-state conformation. For each proenzyme, we used a Rosetta-based approach to identify locations where a disulfide bond spanning the CPG2-prodomain interface is feasible (Fig. 1*E*, *SI Appendix*, Fig. S10). We identified one disulfide variant of Pro-CPG2-1 and two variants of Pro-CPG2-2 that exhibited substantially reduced activity (Fig. 3 *A*–*C*) relative to the CPG2 cysteine



Fig. 4. SAXS analysis for CPG2_{CP-N89}-K177A and the Pro-CPG2 designs. (*A*) The R_g of the enzymes in the absence (red) and presence (green) of hydrolyzed MTX (*Left* axis) and difference in R_g (blue, *Right* axis) are shown. Error bars are the standard error obtained from curve fitting. (*B*) Results from fitting the two-state model (open and closed) to the SAXS data using FoXS are shown, with -MTX and +MTX indicating the absence or presence of hydrolyzed MTX, respectively. The quality of the fits is shown in *SI Appendix*, Fig. S12. (*C*-*D*) The R_g-based (*C*) and volume of correlation (V_c)-based (*D*) dimensionless Kratky plots are shown. Samples with solid lines have no hydrolyzed MTX present, while samples with dashed lines have hydrolyzed MTX present. The horizontal and vertical dotted lines indicate a cross at the peak position of a compact, spherical particle. In the R_g-based plot, if the peak is shifted up and to the right, this indicates that the particle is more elongated, and if the curve becomes a hyperbolic plateau, this is indicative of a disordered particle. In the V_c-based plot, a downward shift indicates an increase in surface-to-volume ratio relative to that of a sphere. (*E*) Rosetta models of the conformational equilibrium required for CPG2 activity, highlighting the role of the prodomain (green).

variant (SI Appendix, Table S1). Notably, when the other three CPG2 cysteine variants (without the prodomain) generated by Rosetta (of a total of six) were tested, they resulted in enzymes that were catalytically inactive. In other words, all disulfide proenzyme variants designed by Rosetta that were compatible with the structure and function of CPG2 resulted in reduced activity. SAXS experiments on the Pro-CPG2-1-Disulfide variant reveal a further moderate decrease in flexibility as deduced from the dimensionless Kratky plots (SI Appendix, Supplementary Results S4 and Fig. 4 C and D). We conclude that a disulfide bridge is effectively created between the prodomain and scaffold, thereby validating our structural models. Moreover, pinning down the prodomain using a secondary covalent linkage assists in strengthening the effectiveness of the prodomain, presumably by decreasing its conformational freedom, and establishes a reasonable maximum inhibition that could be obtained following optimization of the Pro-CPG2-1 design.

Pro-CPG2-1-Disulfide Has a Reduced Ability to Detoxify Methotrexate in Cancer Cell Culture. As we plan to use the designs described here as a starting point for the development of therapeutics or diagnostics, we sought to ascertain the level of inhibition in a more complex biological context prior to initiating the extensive optimization required for specific applications. To that end, we treated an aggressive prostate cancer cell line, PC-3, with Pro-CPG2-1-Disulfide and CPG2_{CP-N89}-K177A. The PC-3 cell line was derived from a bone metastasis of a grade IV prostate cancer patient and is commonly used to assess treatment responses, including chemotherapies, making it ideal for evaluating the capability of designed proenzymes to degrade the chemotherapeutic methotrexate in a cell survival assay (44). As methotrexate is toxic to PC-3 cells, CPG2 activity in the cell culture medium is expected to increase cell survival in the presence of methotrexate. Designed Pro-CPG2 enzymes have significantly lower methotrexate hydrolysis



Fig. 5. Pro-CPG2-1-Disulfide has a reduced ability to cleave and detoxify MTX compared to CPG2_{CP-N89}-K177A. PC-3 cells were treated with MTX at 50 nM and then treated with either CPG2_{CP-N89}-K177A or Pro-CPG2-1-Disulfide. Enzyme concentrations between 250 and 5,000 pM showed 100% viability. Enzyme concentrations between 15 and 62 pM showed a significant difference (unpaired *t* test, *P* < 0.05) of viability in PC-3 cells between the two enzyme treatment groups, with cells treated with Pro-CPG2-1-Disulfide exhibiting a lower survival rate than cells treated with active CPG2. The dashed line indicates the cell viability in the presence of MTX alone (no enzyme treatment). Error bars indicate the SD. *, *P* < 0.05; **, *P* < 0.01; ****, *P* < 0.0001. Two-factor analysis of variance *P* < 10⁻⁸. See also *SI Appendix*, Fig. S11.

activity and are, therefore, expected to lead to decreased cell survival compared to unmodified CPG2. PC-3 cells were incubated with Pro-CPG2-1-Disulfide or CPG2_{CP-N89}-K177A and then subsequently treated with 50 nM methotrexate over 6 days. Cells treated with the Pro-CPG2-1-Disulfide variant indeed displayed significantly lower dose-dependent cell viability at enzyme concentrations of 15 to 62 pM when compared to CPG2_{CP-N89}-K177A (Fig. 5). These data indicate that the designed disulfide-containing proenzyme remains effectively autoinhibited in a cell culture-based assay. When a similar experiment was performed with Pro-CPG2-1, which is less strongly inhibited but could allow proteolytic removal of the prodomain, a smaller, qualitative reduction in PC-3 cell viability was also observed as compared to CPG2_{CP-N89}-K177A (SI Appendix, Fig. S11). Nonetheless, the increased cell survival of the disulfide variant suggest that further optimization of the Pro-CPG2-1 design is possible and may yield effective switches in selected therapeutic contexts.

Discussion

We have developed a massively parallel enzyme design and screening approach to engineer a proenzyme form of CPG2, an FDA-approved drug and one of the best-studied DEPT enzymes. This approach has allowed us to achieve a reversible decrease in activity of up to 80% relative to the CPG2_{CP-N89}-K177A variant without further design optimization. Inhibition can be reversed by the activity of two structurally and function-ally distinct proteases, including MMP-2 that is overexpressed by many cancer cell lines. Introduction of a disulfide bond between the prodomain and the enzyme, which allows us to

simultaneously validate our structural model and establish an upper bound for inhibition for our design, resulted in 98% inhibition compared to the base $CPG2_{CP-N89}$ -K177A variant, although the reactivation of disulfide-containing protein is less efficient.

The design challenge for developing a pro-CPG2 enzyme is substantial. Unlike natural and previously engineered zymogens which act upon macromolecular substrates, CPG2 substrates are relatively small molecules, offering less opportunity to directly sterically impede substrate binding by design. The prodomain binding site is not known to naturally form protein-protein interactions whose binding modes can serve as templates for computational design. Moreover, considering its proximity to the active site, any substitutions within the catalytic or dimerization domains could decrease the activity of the postcleavage form of the designed proenzyme. Therefore, we employed a one-sided design strategy in which only the prodomain is designed while the rest of the protein sequence is held fixed, further increasing the challenge as compared to design strategies in which both partners in a protein-protein interface are allowed to be optimized simultaneously.

Computational approaches using de novo-designed miniproteins have shown great promise in the development of protein binders against a variety of therapeutically relevant targets (23, 45), but these methods have not been used thus far for the design of proenzymes. The ability to synthesize genes corresponding to tens of thousands of designed proteins as oligonucleotide pools coupled with a high-throughput screening assay and NGS allow one to test a broad swathe of sequence and structure space for the desired functionality, allowing for successful design campaigns despite low success rates (<10%) (45) in the computational design step. We used these developments to design miniprotein-based prodomains. Although the individual candidate miniprotein scaffolds are predicted to be well folded in isolation, success in proenzyme design requires that they fold properly and remain soluble and stable when fused to the CPG2 circular permutant. To serve as an effective prodomain, they should feature sufficiently robust inhibitory interactions while still allowing dissociation upon protease cleavage. Furthermore, as is common for enzymes, multiple states (open and closed) are likely involved in the catalytic cycle, but highresolution structures are available for only one (open) state, highlighting the additional challenges unique to proenzyme design compared to the protein-protein interface design. Given these considerations, we reasoned that a massively parallel design and screening approach with a diversity of prodomain structures would maximize opportunities to identify strongly inhibiting prodomains that can also be effectively removed by proteolysis. It is noteworthy that the prodomain structures and sequences that we identified are highly varied, suggesting that the design approach developed here is able to identify multiple, distinct solutions to the problem, rather than converging on a single solution, as is often the case with winner-takes-all directed evolution strategies.

Compared to natural zymogens in which activity level differences between inhibited and activated states are ~100-fold, the best (non-disulfide-containing) designed CPG2 zymogens show ~fivefold increases upon activation, indicating that these diverse solutions identified by computation require further improvements, possibly using directed evolution, in future efforts aimed at improving the dynamic range of the designed proenzymes for therapeutic applications. One avenue for improvement would be to allow further substitutions in the catalytic domain that may improve prodomain binding. A comparative analysis of computational designs that are and are not effective inhibitors also provides insights for further improvement (*SI Appendix*, Fig. S6 and Table S6). Contacts between the prodomain and specific regions in the catalytic and dimerization domains appeared to be significantly greater in designs identified as inhibitors in the massively parallel screening assay, suggesting that increasing the focus on these regions may yield stronger inhibitors during design optimization.

There are at least two putative mechanisms for the inhibition of activity observed in Pro-CPG2 designs. In one case, if the prodomain blocks the active site or binds nearby, it may impede binding of the substrate, either directly or through subtle distortions of the active site. Alternatively, as CPG2 is expected to undergo a closing conformational change during catalysis, the prodomain may act as a door stop, preventing the enzyme from closing after substrate binding, thereby impeding catalysis. Because we designed the prodomains to be in close proximity to the catalytic zinc ions while also having them located at the hinge region of the expected closure site, our design strategy could impact either or both of these mechanisms of inhibition. Our structural and kinetic characterization provide insight into the mechanisms of inhibition. If the prodomain is acting to block the active site, rather than directly interfering with the catalytic machinery of the enzyme, one might expect that this would manifest as a decrease in affinity with the substrate, as evidenced by an increase in K_M. As the catalytic residues and conformational changes are unimpeded in this model, the k_{cat} should be minimally affected. Conversely, if the substrate is free to bind the enzyme, but the enzyme's ability to pass through its catalytic cycle is impeded, the k_{cat} is expected to be reduced, while the K_M should be unaffected. In two of the three most inhibited designs, we observe an increase in K_M and a decrease in k_{cat}, suggesting that both mechanisms are at play. The third design is characterized solely by a decrease in k_{cat}, suggesting that the primary mechanism of inhibition occurs via disruption of catalytic turnover.

The SAXS data provide further insight into the molecular mechanism of inhibition. We present direct evidence that CPG2 undergoes open-closed conformational changes, as observed for homologous enzymes (33-38). Our data highlight the conformational selection underlying CPG2 activity, namely, that CPG2_{CP-} _{N89}-K177A, upon an addition of product, shifts toward the more compact closed conformation. In the Pro-CPG2 designs, this conformational shift does not seem to occur, suggesting that this change is impeded by the presence of the prodomain. This trend is further validated by an analysis of the SAXS flexibility data; while various SAXS-derived metrics suggest that CPG2_{CP-N89}-K177A is more compact and less flexible in the presence of hydrolyzed methotrexate than in its absence, the Pro-CPG2 samples display minimal corresponding variation. Furthermore, these data are generally consistent with a more compact Pro-CPG2 structure.

An analysis of the calorimetric profiles characterizing in trans binding of the prodomain of Pro-CPG2-1 revealed dissociation constants in the low micromolar range. In a therapeutic context, this affinity is sufficient to inhibit the enzyme, while likely being weak enough to be readily released from the binding site following proteolytic processing. In order to evaluate whether proenzymes with similar inhibitory characteristics have potential utility for in vivo or clinical applications, we examined whether the proenzyme form of CPG2 has a diminished ability to protect PC-3 cancer cells from methotrexate toxicity. Our results demonstrate that the decreased activity of the Pro-CPG2-1-Disulfide variant results in a statistically significant reduction in the enzyme's ability to detoxify the drug over a multiday experiment in tumor cells, while the base Pro-CPG2-1 variant shows a smaller, qualitative difference in cell survival. This activity differential suggests that further investigation of these proenzymes on additional cell lines and animal models as well as in clinically valuable contexts is warranted. For example, the use of a tumor-activated proenzyme able to activate a chemotherapeutic prodrug in antibody DEPT (ADEPT) would further mitigate the serious side effects associated with conventional chemotherapy beyond what is already achieved using ADEPT. ADEPT enzymes such as CPG2 are directed to the tumor using a targeting molecule, such as a tumor-directed antibody, where they can activate the drug, thereby localizing the cytotoxic effects to the tumor site. Clearance times for the circulating enzyme vary widely from patient to patient (46, 47), requiring the development of a variety of complex methods for assaying serum enzyme levels (46) and ultimately precluding their clinical use to date. While we previously designed a light-activated, prodrug-activating enzyme (48), challenges associated with the low tissue penetrance of light may limit the applicability of this approach in the clinic as compared to protease-activated enzymes. As the Pro-CPG2 enzymes that we have developed exhibit reduced activity compared to their activated form in tumor cell culture, selection of an appropriate protease that is upregulated in the tumor microenvironment could facilitate synergistic improvement in the pharmacological safety profile of ADEPT therapy. The ability to easily switch the activating protease offers a distinct advantage, as the approach can be readily adapted to tumors with varied protease expression profiles.

Conclusion

The computational design of zymogen forms for any chosen enzyme is a challenging problem, particularly for enzymes acting on small molecule substrates. This study presents a massively parallel computational design and screening approach to develop diverse proenzymes of CPG2, a clinically useful enzyme acting on small molecule substrates, whose inhibition is readily reversed through proteolytic cleavage of the enzymeprodomain linker. A key aspect of our computational method is the use of a structurally diverse set of prodomains to create protein-protein interfaces, and our screening approach exploits advances in synthetic DNA technology and NGS to build and screen a library of thousands of prodomains without requiring specialized assays or equipment. The characterization of our best proenzyme design reveals that the following two mechanisms of inhibition are at play: binding of the substrate to the enzyme is impeded, and domain movements required for catalytic activity are also disrupted. Finally, we demonstrate that the activity differential between the disulfide-linked, inhibited form and activated form is sufficient to significantly impact cell survival under methotrexate treatment, while the design without the disulfide link showed a smaller impact on cell survival. This design and screening strategy has potential implications for the development of a broad range of enzyme inhibitors and proenzymes, potentially yielding therapeutics and diagnostic tools.

Materials and Methods

Complete experimental methods are described in SI Appendix.

Hotspot Placement and Prodomain Design. Rosetta's PlaceStub mover (49) was employed to place small domains interacting with the catalytic zinc ion. An interaction between one of the active site zincs and a glutamate introduced in

the prodomain was defined as the hotspot. A library of 466 inverse rotamers of glutamate interacting with the active site zinc was built using the TryRotamers mover. Hotspot-directed prodomain placement was performed using script 3 (*SI Appendix*, Table S9). This protocol produced 4,444 designs, which were further filtered to 1,668 designs after analyzing the scores and filter values.

Increasing Prodomain Sequence Diversity. To generate a larger set of designs, further rounds of fixed backbone design were applied to the set of 4,444 designs produced from the Hotspot protocol. Either the standard fixed-backbone design protocol or a design protocol (HBNet) aimed at obtaining hydrogen bond networks at the interface (50, 51) was used as described in scripts 4 and 5 (*SI Appendix*, Table S9). These protocols produced 8,757 (fixed backbone) and 4,993 (HBNet) additional designs, for a total of 15,418 designs (*SI Appendix*, Table S4). As described in *SI Appendix*, 7,500 designs and controls were ordered and screened using a high-throughput cell survival assay.

Disulfide Variant Design. Disulfide-bridge variants were designed starting from the base prodomain models using the Disulfidize mover (39) (script 8, *SI Appendix*, Table S9) and evaluated for quality manually.

Determination of CPG2 Molar-Specific Activity. CPG2 was first diluted to 100 μ g/mL in 50 mM Tris and 100 mM NaCl (pH 7.4) and then to 1 to 10 μ g/mL in the same buffer, after which 10 μ L was mixed with 90 μ L of 50 mM Tris, 50 mM NaCl, and 0.1 mM ZnSO₄ (pH 7.4) containing 100 μ M of methotrexate preheated to 37 °C (final protein concentration of 100 to 1,000 ng/mL). The absorbance at 320 nm was measured over time in a 96-well plate (Greiner Bio-One UV-Star Half Area 96-well Microplate) using a Tecan Infinite M200 Pro. Absorbance units were converted to units of molar concentration of methotrexate, and initial rates (molar concentration of methotrexate consumed per second) were determined using linear regression. Activity is reported as the initial rate divided by the molar-specific activity is divided by the molar-specific activity of the reference enzyme and converted to a percentage.

Michaelis–Menten Kinetics. CPG2 was first diluted to 100 µg/mL in 50 mM Tris and 100 mM NaCl (pH 7.4) and then to 50 ng/mL, or 500 ng/mL for the disulfide variants, in the same buffer. A total of 10 µL of this sample was mixed with 90 µL of 50 mM Tris, 50 mM NaCl, and 0.1 mM ZnSO₄ (pH 7.4) containing between 1 and 300 µM methotrexate, preheated to 37 °C. The final protein concentration was ~1 nM (or 10 nM for the disulfide variants). The absorbance at 320 nm was measured over time as described above. Absorbance values were converted to methotrexate concentrations, and initial rates were determined using linear regression. Michaelis–Menten parameters were determined using nonlinear regression.

CPG2 Cell Viability Assay. PC-3 cells were seeded at 1,400 cells/100 μ L in 96-well plates in RPMI media (Sigma-Aldrich) with 10% fetal bovine serum (FBS), 1% penicillin-streptomycin (Pen/Strep), and 1% GlutaGro (Corning). After an

- A. R. Khan, M. N. G. James, Molecular mechanisms for the conversion of zymogens to active proteolytic enzymes. *Protein Sci.* 7, 815–836 (1998).
- J. Chang *et al.*, Enzyme-instructed activation of pro-protein therapeutics in vivo. J. Am. Chem. Soc. 141, 18136–18141 (2019).
- L. Boon, E. Ugarte-Berzal, J. Vandooren, G. Opdenakker, Protease propeptide structures, mechanisms of activation, and functions. *Crit. Rev. Biochem. Mol. Biol.* 55, 111–165 (2020).
- J. L. Arolas, T. Goulas, A. Cuppari, F. X. Gomis-Rüth, Multiple architectures and mechanisms of latency in metallopeptidase zymogens. *Chem. Rev.* 118, 5581–5597 (2018).
- R. J. Johnson, S. R. Lin, R. T. Raines, A ribonuclease zymogen activated by the NS3 protease of the hepatitis C virus. FEBS J. 273, 5457–5465 (2006).
- R. F. Turcotte, R. T. Raines, Design and characterization of an HIV-specific ribonuclease zymogen. AIDS Res. Hum. Retroviruses 24, 1357–1363 (2008).
- D. M. Mitrea, L. S. Parsons, S. N. Loh, Engineering an artificial zymogen by alternate frame protein folding. Proc. Natl. Acad. Sci. U.S.A. 107, 2824–2829 (2010).
- P. Plainkum, S. M. Fuchs, S. Wiyakrutta, R. T. Raines, Creation of a zymogen. Nat. Struct. Biol. 10, 115–119 (2003).
- J. Puskas *et al.*, Development of an attenuated interleukin-2 fusion protein that can be activated by tumour-expressed proteases. *Immunology* 133, 206–220 (2011).
- J. Gerspach et al., Restoration of membrane TNF-like activity by cell surface targeting and matrix metalloproteinase-mediated processing of a TNF prodrug. Cell Death Differ. 13, 273–284 (2006).
- T. Yogo, K. Umezawa, M. Kamiya, R. Hino, Y. Urano, Development of an activatable fluorescent probe for prostate cancer imaging. *Bioconjug. Chem.* 28, 2069–2076 (2017).
- L. M. DeAngelis, W. P. Tong, S. Lin, M. Fleisher, J. R. Bertino, Carboxypeptidase G2 rescue after high-dose methotrexate. J. Clin. Oncol. 14, 2145–2149 (1996).

overnight incubation at 37°C and 5% CO₂, media in the wells were replaced with fresh RPMI media with 5% charcoal-stripped serum (CSS)-FBS, 1% Pen/Strep, and 1% GlutaGro (Corning). Cells were then grown for 3 days. We then added one of the following three drug groups: 50 nM methotrexate alone, 50 nM methotrexate + CPG2_{CP-N89}-K177A (100 to 5,000 pM), or 50 nM methotrexate + Pro-CPG2-1-Disulfide (100 to 5,000 pM). Treatment lasted for 6 days with replenishment of the media, drug, and enzymes after 3 days. Cell viability was measured using WST-1 at 1:10 dilution with CSS-FBS media at absorbance of 450 nm (Tecan 1100 Plate Reader).

Data Availability. Structure data have been deposited in the Protein Data Bank (Accession code: 7M6U). All study data are included in the article and/or *SI Appendix*.

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- 13. J. M. Green, Glucarpidase to combat toxic levels of methotrexate in patients. *Ther. Clin. Risk Manag.* **8**, 403–413 (2012).
- K. D. Wyatt et al., Delayed methotrexate clearance despite carboxypeptidase-G2 (glucarpidase) administration in 2 patients with toxic methotrexate levels. J. Pediatr. Hematol. Oncol. 40, 152–155 (2018).
- A. M. Christensen *et al.*, Resumption of high-dose methotrexate after acute kidney injury and glucarpidase use in pediatric oncology patients. *Cancer* **118**, 4321–4330 (2012).
- R. J. Francis *et al.*, A phase I trial of antibody directed enzyme prodrug therapy (ADEPT) in patients with advanced colorectal carcinoma or other CEA producing tumours. *Br. J. Cancer* 87, 600–607 (2002).
- 17. M. P. Napier *et al.*, Antibody-directed enzyme prodrug therapy: Efficacy and mechanism of action in colorectal carcinoma. *Clin. Cancer Res.* **6**, 765–772 (2000).
- K. Y. Choi, M. Swierczewska, S. Lee, X. Chen, Protease-activated drug development. *Theranostics* 2, 156–178 (2012).
- M. Tholen et al., Design of optical-imaging probes by screening of diverse substrate libraries directly in disease-tissue extracts. Angew. Chem. Int. Ed. Engl. 59, 19143–19152 (2020).
- Y. Jamin, T. R. Eykyn, E. Poon, C. J. Springer, S. P. Robinson, Detection of the prodrug-activating enzyme carboxypeptidase G2 activity with chemical exchange saturation transfer magnetic resonance. *Mol. Imaging Biol.* 16, 152–157 (2014).
- S. Rowsell et al., Crystal structure of carboxypeptidase G2, a bacterial enzyme with applications in cancer therapy. Structure 5, 337–347 (1997).
- B. J. Yachnin, S. D. Khare, Engineering carboxypeptidase G2 circular permutations for the design of an autoinhibited enzyme. *Protein Eng. Des. Sel.* **30**, 321–331 (2017).
- G. J. Rocklin *et al.*, Global analysis of protein folding using massively parallel design, synthesis, and testing. *Science* **357**, 168–175 (2017).

- S. J. Kopytek, J. C. Dyer, G. S. Knapp, J. C. Hu, Resistance to methotrexate due to AcrAB-dependent export from *Escherichia coli*. Antimicrob. Agents Chemother. 44, 3210–3212 (2000).
- J. P. Volpato, E. Fossati, J. N. Pelletier, Increasing methotrexate resistance by combination of active-site mutations in human dihydrofolate reductase. J. Mol. Biol. 373, 599–611 (2007).
- J. P. Volpato et al., Multiple conformers in active site of human dihydrofolate reductase F31R/Q35E double mutant suggest structural basis for methotrexate resistance. J. Biol. Chem. 284, 20079–20089 (2009).
- J. Phan et al., Structural basis for the substrate specificity of tobacco etch virus protease. J. Biol. Chem. 277, 50564-50572 (2002).
- E. Morgunova et al., Structure of human pro-matrix metalloproteinase-2: Activation mechanism revealed. Science 284, 1667–1670 (1999).
- C. A. S. A. Minetti *et al.*, Energetics of lesion recognition by a DNA repair protein: Thermodynamic characterization of formamidopyrimidine-glycosylase (Fpg) interactions with damaged DNA duplexes. J. Mol. Biol. **328**, 1047–1060 (2003).
- R. S. Spolar, J. H. Ha, M. T. Record, Jr., Hydrophobic effect in protein folding and other noncovalent processes involving proteins. *Proc. Natl. Acad. Sci. U.S.A.* 86, 8382–8385 (1989).
- J. H. Ha, R. S. Spolar, M. T. Record, Jr., Role of the hydrophobic effect in stability of site-specific protein-DNA complexes. J. Mol. Biol. 209, 801–816 (1989).
- A. M. Nunes, C. A. S. A. Minetti, D. P. Remeta, J. Baum, Magnesium activates microsecond dynamics to regulate integrin-collagen recognition. *Structure* 26, 1080–1090.e5 (2018).
- H. Unno et al., Structural basis for substrate recognition and hydrolysis by mouse carnosinase CN2. J. Biol. Chem. 283, 27289–27299 (2008).
- T. S. Girish, B. Gopal, Crystal structure of *Staphylococcus aureus* metallopeptidase (Sapep) reveals large domain motions between the manganese-bound and apo-states. *J. Biol. Chem.* 285, 29406–29415 (2010).
- D. Jozic et al., Crystal structure of the dinuclear zinc aminopeptidase PepV from Lactobacillus delbrueckii unravels its preference for dipeptides. Structure 10, 1097–1106 (2002).
- K. Håkansson, C. G. Miller, Structure of peptidase T from Salmonella typhimurium. Eur. J. Biochem. 269, 443-450 (2002).
- H. A. Lindner *et al.*, Essential roles of zinc ligation and enzyme dimerization for catalysis in the aminoacylase 1/M20 family. *J. Biol. Chem.* **278**, 44496–44504 (2003).

- B. Nocek *et al.*, The dimerization domain in DapE enzymes is required for catalysis. *PLoS One* 9, e93593 (2014).
- G. Bhardwaj *et al.*, Accurate de novo design of hyperstable constrained peptides. *Nature* 538, 329-335 (2016).
- D. Schneidman-Duhovny, M. Hammel, J. A. Tainer, A. Sali, FoXS, FoXSDock and MultiFoXS: Singlestate and multi-state structural modeling of proteins and their complexes based on SAXS profiles. *Nucleic Acids Res.* 44 (W1), W424–W429 (2016).
- D. Svergun, C. Barberato, M. H. J. Koch, CRYSOL–A program to evaluate X-ray solution scattering of biological macromolecules from atomic coordinates. J. Appl. Cryst. 28, 768–773 (1995).
- F. E. Reyes, C. R. Schwartz, J. A. Tainer, R. P. Rambo, Methods for using new conceptual tools and parameters to assess RNA structure by small-angle X-ray scattering. *Methods Enzymol.* 549, 235–263 (2014).
- R. P. Rambo, J. A. Tainer, Characterizing flexible and intrinsically unstructured biological macromolecules by SAS using the Porod-Debye law. *Biopolymers* 95, 559–571 (2011).
- S. Tai et al., PC3 is a cell line characteristic of prostatic small cell carcinoma. Prostate 71, 1668–1679 (2011).
- L. Cao et al., De novo design of picomolar SARS-CoV-2 miniprotein inhibitors. Science 370, 426–431 (2020).
- C. J. Springer, G. K. Poon, S. K. Sharma, K. D. Bagshawe, Analysis of antibody-enzyme conjugate clearance by investigation of prodrug and active drug in an ADEPT clinical study. *Cell Biophys.* 24-25, 193-207 (1994).
- 47. K. D. Bagshawe, Targeting: The ADEPT story so far. Curr. Drug Targets 10, 152-157 (2009).
- K. M. Blacklock, B. J. Yachnin, G. A. Woolley, S. D. Khare, Computational design of a photocontrolled cytosine deaminase. J. Am. Chem. Soc. 140, 14–17 (2018).
- S. J. Fleishman et al., Hotspot-centric de novo design of protein binders. J. Mol. Biol. 413, 1047-1062 (2011).
- S. E. Boyken et al., De novo design of protein homo-oligomers with modular hydrogen-bond network-mediated specificity. *Science* 352, 680–687 (2016).
- J. B. Maguire, S. E. Boyken, D. Baker, B. Kuhlman, Rapid sampling of hydrogen bond networks for computational protein design. J. Chem. Theory Comput. 14, 2751-2760 (2018).