Directed evolution of the metalloproteinase inhibitor TIMP-1 reveals that its N- and C-terminal domains cooperate in matrix metalloproteinase recognition

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Tissue inhibitors of metalloproteinases (TIMPs) are natural inhibitors of matrix metalloproteinases (MMPs), enzymes that contribute to cancer and many inflammatory and degenerative diseases. The TIMP N-terminal domain binds and inhibits an MMP catalytic domain, but the role of the TIMP C-terminal domain in MMP inhibition is poorly understood. Here, we employed yeast surface display for directed evolution of full-length human TIMP-1 to develop MMP-3–targeting ultra-binders. By simultaneously incorporating diversity into both domains, we identified TIMP-1 variants that were up to 10-fold improved in binding MMP-3 compared with WT TIMP-1, with inhibition constants ($K_i$) in the low picomolar range. Analysis of individual and paired mutations from the selected TIMP-1 variants revealed cooperative effects between distant residues located on the N- and C-terminal TIMP domains, positioned on opposite sides of the interaction interface with MMP-3. Crystal structures of MMP-3 complexes with TIMP-1 variants revealed conformational changes in TIMP-1 near the cooperative mutation sites. Affinity was strengthened by cinching of a reciprocal “tyrosine clasp” formed between the N-terminal domain of TIMP-1 and proximal MMP-3 interface and by changes in secondary structure within the TIMP-1 C-terminal domain that stabilize interdomain interactions and improve complementarity to MMP-3. Our protein engineering and structural studies provide critical insight into the cooperative function of TIMP domains and the significance of peripheral TIMP epitopes in MMP recognition. Our findings suggest new strategies to engineer TIMP proteins for therapeutic applications, and our directed evolution approach may also enable exploration of functional domain interactions in other protein systems.

The majority of eukaryotic proteins consists of at least two domains, and these multidomain proteins can perform elaborate tasks, often employing a binding or active site located at the domain interface. The separate domains of a multidomain protein fold independently, and yet the interplay between protein domains has an important role in protein structure, stability, and function. Efforts to understand the cooperative impact of individual protein domains on function of multidomain proteins can provide deeper insight into natural protein function and regulation and can also inform new strategies for rational design and engineering of multidomain proteins.

Tissue inhibitors of metalloproteinases (TIMPs), a family of four proteins in vertebrates, are endogenous inhibitors of matrix metalloproteinases (MMPs) that regulate MMP function and activity. TIMPs comprise two domains that pack side by side. The N-terminal domain is widely recognized as the primary inhibitory domain that blocks MMP enzymatic activity by binding at the active site and interacting with the catalytic zinc. The TIMP C-terminal domain in some cases regulates MMP activation through interactions with noncatalytic MMP hemopexin domains or stimulates MMP-independent cell signaling by binding to membrane receptors. Although the TIMP C-terminal domain may also contribute to the binding interface with an MMP catalytic domain, conferring a degree of binding affinity, the significance of this interaction and mechanism by which it contributes to MMP recognition have been largely overlooked. Here, we sought to probe the role of the TIMP C-terminal domain and of cooperativity between TIMP domains in MMP recognition.

Directed evolution of proteins is commonly used to improve or alter protein function toward desired properties but, importantly, can also be used to uncover insights relating sequence to function. Yeast surface display (YSD) is a directed evolution platform for engineering protein binders, including anti-

The abbreviations used are: TIMP, tissue inhibitor of metalloproteinases; MMP, matrix metalloproteinase; YSD, yeast surface display; MMP-3cd, MMP-3 catalytic domain; MTL, multiple-turn loop; FN3, fibronectin type III domain; TLS, translation/libration/screw; PDB, Protein Data Bank; Bis-Tris, 2-[bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol.
bodies (16, 17) or other scaffolds (18), and benefits from eukaryotic secretion machinery that provides quality control for displaying full-length, stable, and properly folded proteins (17, 19), making it a suitable platform for engineering multidomain proteins. YSD has typically been applied to evolve single protein domains for altered binding characteristics, as in the recent engineering of the TIMP-2 N-terminal domain for selectivity toward MMP-9 and -14 (20–22). We hypothesized that by simultaneously applying directed evolution to both domains of a full-length TIMP on the yeast surface, we might probe the roles of each individual domain as well as any cooperative dynamic in MMP recognition; by evolving the intact multidomain protein holistically, we could optimize functional domain interplay.

Here, by creating a YSD library incorporating diversity into both domains of TIMP-1 and analyzing mutations of variants selected for improved binding to an already extremely high-affinity natural target, the MMP-3 catalytic domain (MMP-3cd), we aimed to evaluate the potential contributions of each TIMP domain to MMP affinity. The results of our analyses identify an unsuspected role for the TIMP-1 C-terminal domain in evolving MMP-3 affinity, mediated through synergistic effects between distant sites on the opposing TIMP domains. Protein structures of TIMP-1 mutants cocrystallized with MMP-3cd provide insights into the domain cooperativity of TIMP-1 variants with improved MMP-3cd binding. Our results have important implications for development of therapeutic TIMPs with optimized MMP-binding characteristics, while our conceptual approach may be more widely applicable to the application of directed evolution as a tool to probe domain interplay in other multidomain proteins.

Results

Screening of full-length TIMP-1 surface-displayed library identifies variants with improved binding to MMP-3

To explore the roles of each TIMP-1 domain in modulating MMP binding and inhibition, we first developed a platform for YSD of full-length human TIMP-1. The free, exposed N terminus of TIMP-1 is required for MMP binding and inhibition (6, 23); accordingly, we designed a fusion construct connecting the C terminus of TIMP-1 to the N terminus of yeast cell wall protein Aga2p, with secretion directed by the yeast α-factor signal sequence (Fig. 1A). Display of TIMP-1 bearing the correctly processed, mature N terminus was optimized by deletion of Glu-Ala residues from the signal sequence, which otherwise were inefficiently removed by yeast STE13 protease (Fig. 1). The optimized construct was N-terminally processed by yeast protease Kex2, resulting in efficient display of a functional TIMP-1 fusion that demonstrated robust MMP binding, as detected using biotinylated MMP-3cd (Figs. 1C and 2, A and D). MMP-3cd was selected as a model for the TIMP-1/MMP interaction as it offers a well-studied, well-behaved minimal MMP catalytic domain, and its interaction with TIMP-1 has been structurally characterized (6). We also evaluated the potential benefit of codon optimization of the human TIMP-1 gene for yeast but found minimal impact on efficiency of display (Fig. 1D).

We next generated a targeted library of TIMP-1 mutants, incorporating diversity into MMP contact zones located on both domains of TIMP-1, including 8 residues within the N-terminal domain and 9 residues within the C-terminal domain (Fig. 2, B and C). The TIMP-1 mutant library was constructed aiming to achieve on average three to four mutations per variant through incorporation of degenerate codons and yielded a library size of ~5 × 10^6 independent TIMP-1 variants
in yeast upon transformation. The YSD TIMP-1 library was screened using fluorescence-activated cell sorting (FACS) by incrementally decreasing the concentration of biotinylated MMP-3cd in successive rounds of sorting, selecting for clones with enhanced affinity toward MMP-3cd (Fig. 2A and D). After several rounds of FACS to enrich the pool for variants with highest MMP-3 binding relative to TIMP-1 expression, the enriched pool showed significant enhancement in MMP-3 binding compared with both the naïve library and WT TIMP-1 (Fig. 2D). The further enriched TIMP-1 variants isolated after six rounds of screening, representing a total of four unique sequences, showed up to 10-fold binding improvement compared with WT TIMP-1 (Fig. 3, A–C).

**Analysis of TIMP-1 single and double mutants reveals cooperative impact of N- and C-terminal domain mutations on MMP-3 binding**

The high-affinity MMP-3-binding TIMP-1 variants obtained from library screening contained four to five mutations per gene; notably, all sequences contained mutations in both the N-terminal and C-terminal domains (Fig. 3C). To identify the mutations directly responsible for affinity improvements, we...
next generated single-mutant constructs representing the most prevalent mutations and assessed MMP-3cd binding on the yeast surface. Mutations L34G located in the AB-loop and T98D located in the EF-loop of the TIMP-1 N-terminal domain each conferred significant improvement (Fig. 4A), and yet the binding enhancements of these single mutations were modest by comparison with the composite mutants selected from the library (Fig. 3B). Interestingly, the L34G mutation co-occurred repeatedly in combination with Gly-154 mutations in the multiple-turn loop (MTL) of the C-terminal domain located ~30 Å away. To evaluate the potential functional interaction of these distant residues, we next generated and evaluated several TIMP-1 double-mutant constructs. We found that although TIMP-1 single mutations affecting Gly-154 in isolation had no evident cooperative effect when combined with the L34G mutation (Fig. 4, B–D). In each case examined, the L34G/G154A double mutants showed binding improvements indistinguishable from those of the corresponding library clones, suggesting that mutations at these two sites were sufficient to confer the full functional improvements of selected clones C1, C9, and C14. By contrast, TIMP-1 double mutants combining G154A/H with T98D did not demonstrate similar additive augmentation of MMP-3 binding affinity above that of the T98D single mutant (Fig. 4, E and F), suggesting that Gly-154 mutations modify MMP affinity in a context-dependent fashion rather than conferring an independent and additive effect.

To extend our findings and quantitatively evaluate protease-inhibitory activity of TIMP-1 variants in solution, we next cloned, expressed, and purified soluble forms of several of the mutant TIMP-1 proteins using a human cell expression system. We then assessed inhibition of MMP-3cd activity by soluble WT TIMP-1 and TIMP-1 variants L34G, L34G/G154A, and C1 using an enzyme inhibition assay that measured the reduced rate of cleavage of a fluorometric substrate in the presence of increasing concentrations of the inhibitor. Data were fitted to Morrison's tight binding equation (Equation 1) to determine equilibrium inhibition constants ($K_i$) (Fig. 5 and Table 1). Importantly, these solution studies corroborated the YSD findings; variant C1 showed ~5-fold improvement compared with WT TIMP-1, an effect fully recapitulated by the TIMP-1-L34G/G154A variant (Table 1). Thus, we find consistent evidence from both solution studies and YSD demonstrating that TIMP-1 C-terminal domain mutations can act cooperatively with N-terminal domain mutations to modulate MMP binding and inhibition.

Crystal structures of TIMP-1 variants in complex with MMP-3 show conformational changes that stabilize interdomain interactions and MMP-3 binding

To gain structural insights into the effect of amino acid substitutions on the binding mechanism of TIMP-1 with MMP-3cd, we cocryrstallized the TIMP-1-L34G and TIMP-1-C1 (L34G/L151C/L133P/G154A) mutants in complex with MMP-3cd and solved the crystal structures of the complexes. The TIMP-1-L34G/MMP-3cd and TIMP-1-C1/MMP-3cd structures were solved by molecular replacement and refined against diffraction data extending to resolutions of 2.37 and 2.67 Å, respectively. Data collection and refinement statistics are summarized in Table 2. Each complex shows the expected protein architecture of TIMP-1 and MMP-3cd as described previously for the complex with WT TIMP-1 (PDB code 1UEA) (6); however, notable deviations are seen in the vicinity of the mutated residues that can account for the observed improvements in binding and inhibition.

A striking feature observed in both mutant structures, proximal to the L34G mutation at the enzyme/inhibitor interface, is the observed displacement of the corresponding C-terminal domain residues and the concomitant conformational changes of the TIMP-1 C-terminal domain that stabilize interdomain interactions and MMP-3 binding.
is a reciprocal “clasp” formed by Tyr-35 of TIMP-1 and Tyr-153 of MMP-3 (Fig. 6, A and B). The two Tyr rings, one from each protein, stretch across the interface to form H-bonds with the partner protein, TIMP-1 Tyr-35 with the MMP-3 Phe-154 backbone carboxyl and MMP-3 Tyr-155 with the TIMP-1 Gly-34 backbone carboxyl. Additionally, the two Tyr aromatic rings form hydrophobic and edge-to-face π interactions bridging the interface. Formation of this reciprocal tyrosine clasp is facilitated by substantial back-bone displacements in the TIMP-1-L34G AB-loop and the MMP-3 S-loop compared with the WT TIMP-1/MMP-3cd complex (Fig. 6C). In the WT TIMP-1 structure, bulkier Leu-34 pushes these two loops apart, preventing Tyr-35 of TIMP-1 from H-bonding to the Phe-154 carboxyl, whereas in the L34G mutant structure the distance between oxygens is shortened from 3.5 to 3 Å. The H-bond between MMP-3 Tyr-155 and TIMP-1 Gly-34 is already present in the WT TIMP-1 complex with a distance of 2.6 Å between oxygens; however, the shift of the MMP-3 S-loop in the mutant complexes enables this short H-bond to be cinched even tighter in the TIMP-1 mutant complexes (Fig. 6).

The TIMP-1-C1/MMP-3cd crystal structure reflects near-identical conformational changes in the TIMP-1 AB-loop and MMP-3cd S-loop, resulting from the L34G mutation, with consequent cinching of the reciprocal tyrosine clasp. In addition, mutations in the C-terminal domain of TIMP-1 produce significant conformational changes in this domain, the effect of which is to strengthen interactions both with the TIMP-1 N-terminal domain and with MMP-3cd. TIMP-1-C1 mutation G154A results in adoption of an α-helical conformation by residues 154–157 of the multiple-turn loop at the interface between the TIMP-1 N- and C-terminal domains (Fig. 7, A, D, and E). Glycine, the native TIMP-1 residue at this position, is known to destabilize α-helical secondary structures (24), and each of the mutations of this residue identified in our library screen, to Ala, His, and Lys, are predicted to confer greater propensity for α-helix formation (25, 26), explaining the surprising array of varied substitutions selected at this position. This alteration
in secondary structure allows the two domains of TIMP-1 to pack together more tightly and introduces a new H-bond stabilizing the interdomain interaction (Fig. 7D). Additional conformational alterations in the TIMP-1-C1 structure result in closer contact with MMP-3cd, including hydrophobic contacts of TIMP-1-C1 Leu-152 with MMP-3 Thr-215, Tyr-220, and Leu-222 side chains (Fig. 7B) and a new interdomain H-bond between TIMP-1 residues Gln-150 and Pro-133 (Fig. 7C). In sum, the stabilizing effects of the mutations on TIMP-1 C-terminal domain secondary structure and on interdomain interactions serve to integrate the domains of this multidomain protein, producing functional cooperativity, which in turn improves MMP-3cd binding.

Discussion

Despite the common belief that the N-terminal domain of a TIMP is solely responsible for MMP binding and inhibition, here we have found that the C-terminal domain of TIMP-1 and the interplay between the two domains play a significant role in MMP-3 binding. We further have shown that not only the MMP-contacting interface but also the interactions between TIMP domains can be optimized in an integral fashion to achieve desired binding characteristics. Prior studies seeking to define and reshape the protease-inhibitory function of TIMPs have focused on the N-terminal domain due to the early finding that the isolated N-TIMP domain is independently capable of MMP inhibition with only modest loss of affinity compared with full-length TIMPs (7, 12). Site-directed mutagenesis experiments identified several key amino acid residues, mutation of which can shift the inhibition specificity of N-TIMPs (27–35). More recently, directed evolution efforts have employed phage display or YSD screening of N-TIMP-2 libraries to more broadly scan the sequence determinants of binding specificity at the N-TIMP/MMP interface (20–22, 36). These studies have shown TIMPs to be malleable scaffolds, tolerant of mutation and relatively easy to reengineer for an altered spectrum of protease-inhibitory activity. However, prior work has overlooked the potential functional role of the TIMP C-terminal domain, which can contribute up to a third of the contact surface area at the TIMP/MMP interface (11). Furthermore, the more complex tertiary structures of multidomain proteins are often accompanied by more complex functional capabilities (1) where cooperation between domains renders an intact protein as more than the sum of its parts. Here, by applying evolutionary pressure to a library of full-length TIMP-1 variants, diversified in both domains, we have uncovered such functional cooperativity between TIMP domains.

Our structural analyses of TIMP-1 variants lend insight into how TIMP mutations at the periphery of the interface with an MMP, in both the N- and C-terminal domains, can stabilize binding interactions. Within the N-terminal domain AB-loop, we identified the L34G mutation, which facilitates tighter cinching of a reciprocal tyrosine clasp involving Tyr-35 of TIMP-1 and Tyr-153 of MMP-3. Our structures demonstrate shortening of one intermolecular H-bond and formation of a new additional H-bond. H-bonds have long been appreciated as major contributors to protein–protein binding affinity and specificity (37, 38), where the strength of H-bonds generally correlates well with interatomic distance between the proton donor and acceptor (39). Tyrosine residues in particular are highly represented in protein/protein interaction hot spots (40, 41), likely due to the ability to form aromatic π interactions as well as H-bonds, both of which are evidenced in our structures.

### Table 2

X-ray crystallographic data collection and refinement statistics

| Crystal data                  | TIMP-1-L34G/MMP-3cd | TIMP-1-C1/MMP-3cd |
|-------------------------------|-------------------|-------------------|
| Structure name                | TIMP-1             | TIMP-1            |
| PDB code                      | 6MAV              | 6N9D             |
| Data collection               |                   |                   |
| Resolution range (Å)          | 52.59–2.37 (2.46–2.37) | 43.99–2.67 (2.77–2.67) |
| Space group                   | P 6, 2 2          | P 6, 2 2         |
| a, b, c (Å)                   | 90, 90, 120       | 90, 90, 120      |
| R<sub>merge</sub>              | 0.066 (0.791)     | 0.247 (1.838)    |
| R<sub>free</sub>               | 0.069 (1.180)     | 0.253 (1.882)    |
| Ramachandran outliers (%)     | 12.1 (12.7)       | 20.4 (21.1)      |
| Multiplicity                  | 99.9 (100.0)      | 99.5 (89.3)      |
| Refinement                    |                   |                   |
| Unique reflections used in refinement | 19,911 | 14,162 |
| R<sub>work</sub>/R<sub>free</sub> | 0.217/0.298       | 0.184/0.258      |
| Number of non-hydrogen atoms  | 2,681             | 2,675            |
| Macromolecules                | 2,643             | 2,624            |
| Ligands                       | 5                 | 5                |
| Solvent                       | 33                | 46               |
| Protein residues              | 335               | 336              |
| r.m.s.d. bonds (Å)            | 0.010             | 0.009            |
| r.m.s.d. angles (°)           | 1.35              | 1.30             |
| Ramachandran favored (%)      | 91                | 94               |
| Ramachandran outliers (%)     | 1.2               | 0.3              |
| Average B-factor              | 77.60             | 55.10            |
| Macromolecules                | 77.70             | 55.20            |
| Ligands                       | 69.80             | 51.60            |
| Solvent                       | 71.30             | 49.20            |

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Within the C-terminal domain, we identified the G154A mutation, responsible for conformational changes in the multiple-turn loop that impact both the binding interface with MMP-3cd and the interface between the TIMP domains. The interface with MMP-3cd gains new hydrophobic stabilizing contacts, while the interdomain interface is stabilized by the defined /H9251-helical secondary structure adopted by TIMP residues 154–157 and by a new interdomain H-bond. Interestingly, TIMP-1 Gly-154, a residue with low helical propensity, was found mutated to Ala, Lys, and His, residues with higher helical propensities (25, 26), in three of four isolated clones with enhanced MMP-3cd–binding affinity. Given that we observed a function-enhancing role for these substitutions only in combination with the L34G mutation, it may be that a G154X mutation alone is insufficient to fully stabilize the helical turn but requires additional structural stability conferred by the reciprocal tyrosine clasp on the opposite side of the intermolecular interface.

Because MMPs represent high-value but challenging therapeutic targets, our results suggest new avenues for engineering MMP-targeted therapeutic proteins. Many human MMPs contribute to pathology of cancer, fibrosis, arthritis, cardiovascular, pulmonary, and other diseases when improperly expressed (23, 42, 43), but clinical trials of broad-spectrum MMP inhibitors in cancer and arthritis proved disappointing (44, 45). Engineered designer TIMPs have been viewed as a possible pathway to achieve greater selectivity and efficacy (23, 46, 47), and recent campaigns to develop such selectivity have employed YSD platforms for directed evolution of N-TIMP-2 aided by computational library design (20–22). Our present findings suggest that future TIMP engineering efforts may benefit from employing a full-length TIMP scaffold and, further, that maximum advantage of the multidomain scaffold may be best achieved through simultaneous optimization to capture cooperative mutations across domains.
Finally, the utility of our approach for identifying and engineering domain cooperativity may extend beyond the TIMP family and have relevance for other systems. Prior studies have demonstrated how connectivity and cooperativity of multiple domains contribute to protein folding, stability, and function (2, 48, 49). Dynamic interactions between individual domains orchestrate folding pathways, determining kinetic and equilibrium properties of cooperative folding (49). Favorable interactions between domains can dictate the stability of individual domain structures (48), and thus interdomain interactions have been critical in the natural evolution of multidomain proteins. Directed evolution can recapitulate this natural evolution of multidomain functionality, e.g. when a low-affinity peptide-binding PDZ domain was linked to an unrelated, functionally inert fibronectin type III domain (FN3), optimization of FN3 led to a highly selective domain-interdependent “affinity clamp” (1). To date, however, the ability to pinpoint specific sequence and structural determinants of domain cooperativity has been limited. Here, directed evolution simultaneously targeting residues within both domains of TIMP-1 enabled us to search for and identify pairs of residues responsible for interdomain cooperativity in TIMP-1 function. This strategy may be more broadly useful to detect domain cooperativity, probing the structure–function paradigm, and to engineer enhanced synergistic function into other multidomain proteins for varied practical applications.

Figure 7. Crystal structure of TIMP-1-C1 variant bound to MMP3cd. A, cartoon representation of the TIMP-1-C1/MMP-3cd complex crystal structure is shown using the same color scheme described for Fig. 6. Altered inter- and intramolecular interactions attributable to TIMP-1-C1 mutations are highlighted in B–D. B, a hydrophobic cluster is formed at the intermolecular interface between TIMP-1 Leu-152 and MMP-3cd residues Thr-215, Tyr-220, and Leu-222. C, a new H-bond within the TIMP-1-C1 C-terminal domain is formed between the carbonyl of Pro-133 and the side chain of Gln-150. D, as a consequence of the G154A mutation, residues 154–157 adopt an α-helical conformation, and a new interdomain H-bond is formed between the side chain of Ser-155 in the C-terminal domain and the carbonyl of Pro-6 in the N-terminal domain. E, superposition of the TIMP-1-C1/MMP-3cd crystal structure (colored as above) with the WT TIMP-1/MMP-3cd structure (PDB code 1UEA; shown in pale blue/pale orange) highlights conformational changes in the multiple-turn loop of TIMP-1-C1, which facilitate stabilizing interdomain interactions within TIMP-1 as well as favorable hydrophobic interactions at the interface with MMP-3cd. F, stick representation of the protein environment surrounding the hydrophobic cluster at the intermolecular interface with 2Fo-Fc map contoured at 1.0σ.
Engineering TIMP-1 domain cooperation

Experimental procedures

Strains and plasmids

The yeast Saccharomyces cerevisiae strain EBY100 (MATa AGA1::GAL1-AGA1::URA3 ura3–52 trp1 leu2-D200 his3-D200 pep4::HIS3 prb11.6R can1 GAL) was used for yeast surface display of human TIMP-1 protein and its variants. Yeast display plasmids for displaying WT TIMP-1 and TIMP-1 mutants at the N terminus of Aga2p protein on the yeast surface were derived from the pCHA backbone, pCHA-VRC01 vector (50) purchased from Addgene.

Yeast surface display of TIMP-1

pCHA-TIMP-1 plasmids (the yeast display vectors) were transformed into the yeast strain EBY100 by electroporation using a Bio-Rad Gene Pulser and 2-mm electroporation cuvettes. The yeast cells were grown in minimal SD-CAA medium, pH 6 (20 g/liter dextrose, 6.7 g/liter yeast nitrogen base lacking amino acids, 5.4 g/liter Na₂HPO₄, 8.6 g/liter NaH₂PO₄·H₂O, and 5 g/liter Bacto casamino acids). After 16 h growth at 30 °C, cells were pelleted at 3000 × g and resuspended in SGR-CAA medium (20 g/liter galactose, 6.7 g/liter yeast nitrogen base lacking amino acids, 5.4 g/liter Na₂HPO₄, 8.6 g/liter NaH₂PO₄·H₂O, 5 g/liter Bacto casamino acids, and 20 g/liter raffinose) for induction. Cells were grown in SGR-CAA medium at 30 °C for 16 h before harvesting. The yeast cells displaying TIMP-1 variants were collected at an OD₆₀₀ of 0.2 and washed with cold PBSA buffer (8 g/liter NaCl, 0.2 g/liter KCl, 1.44 g/liter Na₂HPO₄, 0.24 g/liter KH₂PO₄, pH 7.4, and 1% BSA). The cells were first incubated with 100–250 nM biotinylated MMP-3cd and mouse anti-c-myc 9e10 (Sigma) for 1 h on ice. The cells were then washed and resuspended in 100 μl of cold PBSA buffer containing fluorescein-conjugated goat anti-mouse antibody (Thermo Fisher) and streptavidin-Alexa Fluor 647 (each at 1:100 dilution) and incubated on ice for 30 min. After harvesting and washing with cold PBSA buffer, the cells were centrifuged and resuspended in 750 μl of PBSA buffer. Flow cytometry data were then collected using an Attune NxT flow cytometer from at least 10,000 cell events per sample and analyzed using FlowJo software (FlowJo, LLC).

Generating the targeted library of TIMP-1 mutants

A library of human TIMP-1 gene variants, targeting for mutation 17 residues in five loops shown to interact with MMP catalytic domains in crystal structures (Fig. 2C), was purchased from Invitrogen GeneArt Gene Synthesis. The library of mutants was constructed by assembling gene blocks of synthetic oligonucleotides with 13% chance of NNS degenerate codon incorporation at each position targeted for mutation (where N = any nucleotide and S = G or C), to achieve on average three to four mutations per variant. The yeast display TIMP-1 mutant library was generated by PCR amplification of the TIMP-1 gene from the targeted TIMP-1 library using primers with 50-bp overhangs containing homology to sequence upstream and downstream of the TIMP-1 gene in the pCHA-TIMP-1 yeast display vector. PCR products of TIMP-1 gene variants were gel-purified and concentrated using Pellet Paint® Co-Precipitant (EMD Millipore) following the manufacturer’s protocol. The pCHA-TIMP-1 vector was double-digested using BsrGI and BamHI restriction enzymes, and the pCHA digested vector (1 μg) and purified PCR product (5 μg) of the TIMP-1 mutant library were mixed and electrotransformed into yeast cells as described previously (51). Five separate electrotransformations were performed and combined. The yeast cells were then resuspended in cold YPD medium (20 g/liter glucose, 20 g/liter peptone, and 10 g/liter yeast extract) and grown at 30 °C with shaking for 1 h. The library size was estimated by plating yeast transfectants in serial dilution and found to contain about 5 × 10⁷ independent variants. The yeast cells containing the library of TIMP-1 mutants were pelleted and resuspended in 50 ml of SD-CAA medium, pH 4.5 (same as SD-CAA medium described above, but phosphate components were substituted with 13.7 g/liter sodium citrate dehydrate and 8.4 g/liter citric acid anhydrous) and grown for 16 h at 30 °C with shaking.

Screening the yeast-displayed TIMP-1 library using FACS

Prior to each round of sorting, yeast cells were grown and induced; counted (OD₆₀₀ of 1 × 10⁷ cells/ml); incubated with biotinylated MMP-3cd followed by anti-c-myc, streptavidin, and secondary antibody; washed; and suspended in PBSA buffer as described above under “Yeast surface display of TIMP-1.” Samples were maintained on ice and covered from light until loading onto a flow cytometer for analysis (BD Accuri) or library sorting (BD Aria II). Sorted cells were recovered in SD-CAA, pH 4.5, containing 1% penicillin-streptavidin and incubated at 30 °C overnight. Yeast surface protein expression was then induced by culturing cells in SGR-CAA medium at 30 °C overnight. The initial library (~5 × 10⁷ yeast) was subjected to two rounds of screening (staining, sorting, recovery, and regrowth) using a concentration of 100 nM biotinylated MMP-3cd. Screening was continued through four additional rounds, for a total of six rounds, under equilibrium sorting conditions employing incrementally decreasing MMP-3cd concentration from 50 to 5 nM.

DNA sequencing

Following rounds of sorting, recovered yeast were cultured on SD-CAA plates, and plasmid DNA was extracted from the isolated individual yeast clones using the Zymoprep™ yeast plasmid miniprep II kit (ZymoResearch). TIMP-1 mutant plasmids extracted from yeast were transformed into Escherichia coli cells and plated on LB-Amp (100 μg/ml ampicillin). The pCHA-TIMP-1 variant plasmid was then extracted and purified from the bacteria using the Qiagen Miniprep kit according to the manufacturer’s protocol (Qiagen) and sequenced (Eurofins Scientific) with primers up- and downstream of the TIMP-1 gene. Sequences were analyzed using SnapGene software (SnapGene, GSL Biotech, LLC).

Soluble TIMP expression and purification

The pTT-TIMP-1 vector, a generous gift from Dr. B. Tousaint (Grenoble, France), was used for expression of full-length human TIMP-1 (52). TIMP-1 mutants were amplified from the
corresponding yeast display vector (pCHA-TIMP-1) using PCR and inserted between HindIII and BamHI restriction sites of the pTT-TIMP-1 vector. WT TIMP-1 and TIMP-1 variants were expressed in HEK293-FreeStyle cells (Thermo Fisher Scientific) in Gibco® FreeStyle™ 293 Expression Medium (Thermo Fisher Scientific) on a shaker incubator at 37 °C and 8% CO₂. The HEK293-FreeStyle cells were grown to a cell density of 1–1.5 × 10⁶ in 300-ml culture, transfected with 300 μg of pTT-TIMP-1 plasmid using 0.9 ml of 1 mg/ml polyethyleneimine (PEI) transfection reagent, and incubated on the shaker incubator for 72 h before harvesting the cells and collecting the supernatant media. For purification, the clarified medium was concentrated, dialyzed into buffer A (50 mM HEPES, pH 6.8, and 25 mM NaCl), and purified over an SP-Sepharose column using a linear gradient of buffer B (50 mM HEPES, pH 6.8, and 0.5 m NaCl). Fractions containing the WT or mutant TIMP-1 were assessed by silver-stained SDS-PAGE. To prepare homogenous purified proteins for crystallization, TIMP-1 variants were then enzymatically deglycosylated with peptide N-glycosidase F (New England Biolabs). Deglycosylation of TIMP-1 protein was confirmed by SDS-PAGE followed by purification by size-exclusion chromatography using a Superdex-75 column (GE Healthcare) that was equilibrated and eluted with 50 mM HEPES, pH 6.8, containing 150 mM NaCl. The highly pure deglycosylated TIMP-1 fractions identified by silver-stained SDS-PAGE were combined and concentrated using Amicon Ultra-15 centrifugal filter units with a molecular weight cutoff of 10,000.

**MMP expression, purification, and biotinylation**

ProMMP-3cd was expressed from *E. coli* using a pET3a construct featuring human proMMP-3 lacking the C-terminal hemopexin domain, a generous gift of H. Nagase (53). ProMMP-3cd was refolded, purified, and activated as described previously (53). Briefly, recombinant protein was extracted from inclusion bodies in a solution containing 8 M urea, 20 mM Tris-HCl, pH 8.6, 20 mM DTT, and 50 μM ZnCl₂. The protein was purified on Q-Sepharose equilibrated with 8 M urea, 20 mM Tris-HCl, pH 8.6, and 50 μM ZnCl₂, and eluted using a linear gradient of NaCl to 0.5 M. Fractions containing proMMP-3cd were combined, diluted to an A₂₈₀ of 0.3, and refolded by stepwise dialysis with 50 mM Tris-HCl, pH 7.5, 10 mM CaCl₂, and 150 mM NaCl. Subsequently, purified proMMP-3cd was activated overnight in the presence of the organomercurial compound 4-aminophenyl mercuric acetate (1 mM at 37 °C). Concentrations of active MMP-3cd were determined by titration against a TIMP-1 reference stock as described previously (54). MMP-3cd was biotinylated using the EZ-Link NHS-PEG4 biotinylation kit (Thermo Scientific) according to the manual with addition of biotin in a 1:10 molar ratio (protein:biotin) and incubated at room temperature for 30 min. The biotinylated MMP-3cd was purified using Zeba spin desalting columns (Thermo Scientific) and tested for degree of biotinylation using the 4′-hydroxyazobenzene-2-carboxylic acid assay as described in the kit protocol.

**Active TIMP-1 concentration determination by titration**

Concentrations of purified TIMP-1 variants were determined by titration against a reference stock of MMP-3cd previously titrated against WT TIMP-1, essentially as described previously (54). Briefly, MMP-3cd (200 nM) was preincubated for 1 h at 25 °C with a range of substoichiometric concentrations of the TIMP-1 variant. MMP/TIMP mixtures were then assayed for residual proteolytic activity using colorimetric thiopeptolide substrate Ac-Pro-Leu-Gly-[2-mercapto-4-methylpentanoyl]-Leu-Gly-OC₂H₅ (Enzo Life Sciences). MMP/TIMP mixtures were diluted 40-fold into a reaction cuvette containing the thiopeptolide substrate at a final concentration 100 μM in 50 mM HEPES, pH 6.0, 10 mM CaCl₂, 0.05% Brij-35, and 1 mM 5,5′-dithiobis(2-nitrobenzoic acid). Reactions were followed for 10 min at 37 °C on a Varian spectrophotometer (Thermo Scientific), and linear initial rates were measured as an increase in the absorbance at 410 nm (ε₄₁₀ = 13,600 M⁻¹ cm⁻¹). Data were fitted using linear regression analyses and extrapolated to the stoichiometric equivalence point in Prism 7 (GraphPad Software, San Diego, CA).

**TIMP-1/MMP-3 inhibition studies**

Equilibrium inhibition constants (Kᵢ) of WT TIMP-1 and TIMP-1 variants toward MMP-3 were measured using a method appropriate for tight binding inhibition, similar to a method described previously (21). MMP-3cd (0.24 nM) was incubated with 0.04-2.5 nM WT TIMP-1 or TIMP-1 variant in TCNB buffer (50 mM Tris, pH 7.5, 100 mM NaCl, 10 mM CaCl₂, and 0.05% Brij) for 1 h at 37 °C. Thereafter, the fluorogenic substrate Mca-Pro-Leu-Gly-[2-mercapto-4-methylpentanoyl]-Leu-Gly-OC₂H₅ (Enzo Life Sciences) was added to the reaction at a final concentration of 10 μM, and fluorescence was monitored with 340/30 excitation and 400/30 emission filters using a Synergy HT plate reader (BioTek, Winoski, VT) at 37 °C. Fluorescence readings were recorded every minute for 120 min, and enzymatic rates were determined from the slope of the linear portion of the fluorescence signal. To determine Kᵢ, data were plotted as initial velocities versus TIMP-1 or variant concentration and fitted by multiple regression to Morrison’s tight binding inhibition equation (55) (Equation 1) where Vₑ is enzyme velocity in the presence of inhibitor, V₀ is enzyme velocity in the absence of inhibitor, [E] is enzyme concentration, [I] is inhibitor concentration, [S] is substrate concentration, Kᵢ is the Michaelis–Menten constant, and Kᵢₚᵣ is an apparent inhibition constant given by Equation 2. Data were fitted using Prism 7 (GraphPad Software) (Fig. 5). Reported inhibition constants are average values obtained from two independent experiments, each with duplicate samples. Calculations were performed using a Kᵢ value of 11.23 μM for MMP-3cd as determined from three Michaelis–Menten kinetic experiments in our laboratory.

\[
Vₑ = \frac{1 - ([E] + [I] + Kᵢₚᵣ) - \sqrt{([E] + [I] + Kᵢₚᵣ)^2 - 4[E][I]}}{2[E]}
\]

(Eq. 1)
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\[ K_{\text{on}}^{\text{prop}} = K_0 + \frac{[S]}{K_m} \quad \text{(Eq. 2)} \]

Crystallization and X-ray diffraction

TIMP-1 variants and MMP-3cd protein were mixed at a 1:1 molar ratio, and TIMP-1 variant/MMP-3cd protein complexes were concentrated to 2.8–4.8 mg/ml prior to crystallization screening. Crystallization was conducted using the hanging-drop method with the protein solution mixed 1:1 (v/v) with a reservoir solution. TIMP-1-L34G/MMP-3cd protein crystals were grown in condition 92 of the Anotract Top96 crystallization kit (Anotract) (0.1 M ammonium acetate, 0.1 M Bis-Tris HCl, pH 5.5, and 17% (w/v) PEG 10,000). TIMP-1-C1/MMP-3cd protein crystals were grown in condition 41 of the Anotract Top96 crystallization kit (Anotract) (0.2 M sodium acetate, 0.1 M sodium cacodylate HCl, pH 6.5, and 18% (w/v) PEG 8000). TIMP-1 variant/MMP-3cd protein crystals appeared over 48 h and were grown for over a few weeks. Crystals were flash-cooled in liquid nitrogen using cryoprotectant buffer matched to the crystallization reservoir solution and additionally containing 30% dextrose. Single-wavelength (1.0 Å) native X-ray diffraction data were collected at 100 K at Advanced Light Source beamline 8.2.2, Lawrence Berkeley National Laboratory. TIMP-1 variant complex structures were each solved from single crystals that diffracted to 2.37 Å (for TIMP-1-L34G) and 2.67 Å (for TIMP-1-C1). The X-ray data were processed with xia2 (56) using DIALS (57) for indexing, refinement, and integration with POINTLESS (58) and AIMLESS (59) for scaling and merging. \( R_{\text{free}} \) flags were assigned to a random 5% of reflections, and this test set was maintained throughout all subsequent stages of structure solution and refinement.

Structure determination and refinement

X-ray crystal structures of the protein complexes of MMP-3cd/TIMP-1 variants were solved using CCP4 software by molecular replacement using the program Molrep (60). The previously solved structure of human MMP-3cd/WT TIMP-1 (PDB code 1UEA) without the corresponding Zn and Ca ions was used as a search model. Following molecular replacement, Phenix.refine was used for sequential automated refinements (61) alternating with manual adjustments in Coot (62). Because of the flexibility of MMP-3cd and TIMP-1 protein segments and the possibility that they might demonstrate local trajectories of motion partially decoupled from the larger protein domains, we employed TLS refinement using the automatic TLS group finding tool in the Phenix software for refinement of both structures. Several residues in the TIMP-1 variant chains showed only weak and indistinct density and were left unmodelled in the final models, including residues Gly-52, Asp-53, Ala-54, and Ala-55 in both structures and additionally Leu-51 in the TIMP1-L34G/MMP-3cd structure. The final stage of restrained refinement included water molecules with peaks greater than 1σ and within acceptable H-bonding distances from neighboring protein atoms. The coordinates and structure factors for the TIMP-1-L34G/MMP-3cd and TIMP-1-C1/MMP-3cd structures have been submitted to the Worldwide Protein Data Bank under the accession codes 6MAV and 6N9D, respectively. Structure figures were generated using PyMOL (Schrödinger, LLC).

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