HIGH-AFFINITY SMALL PROTEIN INHIBITORS OF HUMAN CHYMOTRYPSIN C (CTRC) SELECTED BY PHAGE DISPLAY REVEAL UNUSUAL PREFERENCE FOR P4' ACIDIC RESIDUES

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Human chymotrypsin C (CTRC) is a pancreatic protease that participates in the regulation of intestinal digestive enzyme activity. Other chymotrypsins and elastases are inactive on the regulatory sites cleaved by CTRC, suggesting that CTRC recognizes unique sequence patterns. To characterize the molecular determinants underlying CTRC specificity, we selected high-affinity substrate-like small protein inhibitors against CTRC from a phage library displaying variants of a natural chymotrypsin inhibitor from Schistocerca gregaria (SGPI-2). On the basis of the sequence pattern selected, we designed eight inhibitor variants in which amino acid residues in the reactive loop at P1 (Met or Leu), P2’ (Leu or Asp) and P4’ (Glu, Asp or Ala) were varied. Binding experiments with CTRC revealed that (i) inhibitors with Leu at P1 bind 10-fold stronger than those with P1 Met; (ii) Asp at P2’ (versus Leu) decreases affinity but increases selectivity and (iii) Glu or Asp at P4’ (versus Ala) increase affinity 10-fold. The highest affinity SGPI-2 variant (K_D 20 pM) bound to CTRC 575-fold tighter than the parent molecule. The most selective inhibitor variant exhibited a K_D of 110 pM and a selectivity ranging from 225-fold to 112,664-fold against other human chymotrypsins and elastases. Homology modeling and mutagenesis identified a cluster of basic amino acid residues (Lys⁵¹, Arg⁵⁶ and Arg⁹⁰) on the surface of human CTRC that interact with the P4’ acidic residue of the inhibitor. The acidic preference of CTRC at P4’ is unique among pancreatic proteases and might contribute to the high specificity of CTRC-mediated digestive enzyme regulation.

Digestion of dietary proteins in the small intestines is catalyzed by proteases secreted from the pancreas. These proteases are produced as inactive proenzymes (zymogens) and their activation is spatially restricted to the duodenum and proceeds in a cascade like manner. The membrane-localized serine protease enteropeptidase (enterokinase) activates trypsinogens to active trypsins, which, in turn, activate all other proteolytic proenzymes; in the human these include chymotrypsinogens B1, B2, C and L1, proelastases 2A, 3A and 3B, and procarboxypeptidases A1, A2 and B1. This classic textbook paradigm of digestive enzyme activation has been recently revised by the discovery of interactions between digestive proteases which can modulate the ultimate levels of active enzymes. In this regard, we demonstrated that human chymotrypsin C (CTRC) regulates activation and degradation of human cationic trypsinogen and trypsin [1, 2]. CTRC facilitates trypsin-mediated activation (autoactivation) of cationic trypsinogen by processing the trypsinogen activation peptide, at the Phe¹⁸-Asp¹⁹ peptide bond, to a shorter form which is more readily cleaved by trypsin [1]. Mutation A16V in the trypsinogen activation peptide stimulates CTRC-mediated processing and subjects with this mutation are at increased risk of developing chronic pancreatitis [1, 3]. CTRC can also promote degradation of human cationic trypsinogen and trypsin by cleaving the conserved Leu⁸¹-Glu⁸² peptide bond in the calcium binding loop, when the calcium concentration is low and the binding loop is unoccupied [2].
combination of this cleavage and an autolytic cleavage at Arg^{122} results in the degradation and complete inactivation of cationic trypsin. Hereditary pancreatitis-associated mutation R122H eliminates the Arg^{122} autolysis site in cationic trypsin(ogen) and prevents CTRC-induced degradation, suggesting that trypsin(ogen) degradation is an important defensive mechanism in the pancreas [2, 4]. Consistent with this notion, genetic association studies identified loss-of-function mutations in CTRC as risk factors for chronic pancreatitis [5, 6].

More recently, we found that human CTRC is a physiological co-activator of human procarboxypeptidases A1 and A2 (proCPA1 and proCPA2) [7]. Digestive carboxypeptidases are secreted by the pancreas as inactive proenzymes containing a 94-96 amino acid long propeptide which acts as a strong inhibitor of the enzymes thereby maintaining their zymogen state. Activation of procarboxypeptidases is initiated by trypsin-mediated proteolytic cleavages at the C-terminal end of the propeptides. However, the trypsin-processed propeptides are still inhibitory and subsequent cleavages by CTRC are required for full activation of proCPA1 and proCPA2. CTRC cleaves the propeptides at multiple sites, but the critical step for activation appears to be cleavage of the conserved Leu^{96}-Leu^{97} peptide bond in both proenzymes [7].

The CTRC-catalyzed regulatory functions are highly specific as other human pancreatic chymotrypsins and elastases exhibit no such activity [1, 2, 7]. This may be partly due to the primary (P1) specificity of CTRC, which is chymotrypsin-like, but CTRC exhibits higher activity on leucyl peptide bonds than chymotrypsin A or B [8-10]. However, other determinants may be also important, as suggested by the preponderance of acidic amino acid residues on the prime side of the CTRC-specific regulatory cleavage sites (Table 1). Despite the similarities in these sequences, the number of CTRC-specific cleavage sites identified hitherto is too few to determine the inherent binding site preference of CTRC. Functional importance of individual amino acid residues within a linear binding epitope of substrates can be assessed through mutational replacements, but a complete analysis would require an impractically large number of mutants. If recognition relies on cooperating substrate positions, the combination of mutants required for full coverage of all possible sequence variations would reach astronomical numbers. Directed evolution combines combinatorial mutagenesis with functional selection and provides a straightforward solution to these problems [11]. Substrate-like canonical inhibitors mimic natural peptide substrates in their interactions with proteases [12, 13]. In the present study we utilized phage-display, possibly the most robust directed evolution approach, to define the substrate specificity of human CTRC. We fully randomized six protease binding loop positions in the chymotrypsin inhibitor SGPI-2, displayed the library on M13 phage, and selected tight binding variants against CTRC. Based on the selected sequence pattern, eight SGPI-2 variants designated C1 through C8 were produced and characterized. The results suggest that a unique electrostatic interaction between a cluster of basic residues on CTRC and the P4' acidic amino acid residue of CTRC substrates is an important determinant of CTRC specificity.

**EXPERIMENTAL PROCEDURES**

**Nomenclature.** Amino acid residues in human pancreatic proteases were numbered starting with the initiator methionine of the primary translation product; according to the recommendations of the Human Genome Variation Society. Where indicated, the traditional crystallographic numbering system based on the bovine chymotrypsin A sequence was used. Amino acids in SGPI-2 were numbered starting from the first residue in the mature inhibitor (Glu^1).

**SGPI library construction.** Library construction was based on the Tag-wtSGPI-2-pGP8 phagemid vector [14], which monovalently displays SGPI-2 on the p8 coat protein of the M13 phage. The library was produced in two successive Kunkel mutagenesis [15] steps in order to avoid wild-type SGPI-2 contamination. First, the Tag-SGPI-2-pGP8-STOP vector was constructed in which all codons to be randomized were replaced with stop codons (where the stop codons are underlined): 5'-GC GGT AGC GAT GGC AAA AGC GGT ACC GGT GGC AAA AGC GCG TAA TGC TAA TAA TAA TGC TAA CAG GGT ACC GGT GGA GG-3'. In the second step, the resulting Tag-
wtSGPI-2-pGP8-STOP vector was used as template in a combinatorial Kunkel mutagenesis step modified slightly for large scale generation of diverse libraries [16]. Stop codons were replaced with amino acid coding codons randomized using NNK degeneracy, where N denotes nucleotides A, C, G or T, while K stands for G or T. NNK codons represent a set of 32 codons covering all 20 amino acid residues. The following mutagenesis primer was used (where NNK codons are underlined): 5'-GGT AGC GGT GGC AAA AGC GCG NK TGC NNK NNK NK TGC GNK CAG GGT ACC GGA GG -3'. The Tag-SGPI-2-pGP8-Lib library construct is illustrated in Supplementary Figure S1. The phagemid library was electroporated into E. coli SS320 cells to generate phage libraries as described [16].

**Selection of inhibitor-phages on CTRC.** Human CTRC was immobilized in 12 wells of an Immobilizer Amino plate (Nunc International) using 5 µg CTRC per well in 100 µL of 10 mM Hepes buffer (pH 7.8) containing 0.15 M NaCl; for 3 hours. The wells were rinsed with phosphate-buffered saline (PBS, pH 7.2) and blocked with 200 µL of bovine serum albumin (BSA, 5 mg/mL dissolved in PBS) for 1 hour. A control plate was treated with BSA only without adding CTRC. The wells were rinsed four times with PBS containing 0.05% Tween-20 (final concentration). Phages (100 µL, ~5×10¹¹ phage particles per well) were added to the wells in PBS/BSA solution containing 0.05% Tween-20 and incubated for 2 hours. Plates were rinsed 12 times with PBS containing 0.05% Tween-20 and bound phages were eluted at pH 1.0 with 100 mM HCl (100 µL/well) for 1 min. The eluted phage solution was neutralized by adding 15% volume of 1 M Tris base solution and phages were amplified in E. coli XL1-Blue. Three selection and amplification cycles were performed as described [16]. After the second and third cycles, the inhibitor-phage titers eluted from target and control plates were determined and enrichment values were calculated to characterize the efficiency of the selection process. The enrichment was 40-fold and 900-fold after the second and third cycles, respectively.

**Phage-ELISA of selected library members.** Individual clones from the third selection cycle were tested in phage-ELISA performed as described [16]. Clones producing ELISA signals 3-fold higher on CTRC containing plates than on albumin coated control plates were selected for DNA sequencing.

**Sequence analysis.** DNA sequences coding for SGPI-2 variants were PCR-amplified from the selected library members with the following primers annealing to invariant vector sequences. Forward primer, pTacUp35T7: 5'- CGA AAT TAA TAC GAC TCA CTA TAG GGC TAT AGG GTC GTA ATG ATA ATG TTT TTT GCC CC -3' and reverse primer, pVIII-rev: 5'- GTT ATG CTA GTT ATT CCT AGC CCT CTT GTC TTC AAC CCG CC -3'. The forward PCR primer was designed to contain the sequence of the T7 promoter sequencing primer: 5'- CGA AAT TAA TAC GAC TCA CTA TAG GG -3', which was then used for the sequencing reactions. Clones with unique DNA sequences were aligned and amino acid frequencies at the randomized positions were determined. These frequencies were normalized to the expected codon frequencies in the NNK degenerated set, to eliminate the effects of codon bias. For logo representation of the normalized results an input sequence dataset containing 100 sequences was generated representing the normalized amino acid frequencies at each randomized positions. Sequence logos were generated by the web-based application, WebLogo [17] available at http://weblogo.berkeley.edu/logo.cgi.

**Expression and purification of SGPI-2 variants.** Recombinant SGPI-2 variants were expressed into the periplasm of E. coli as MBP fusion proteins [18]. PCR amplified genes of SGPI-2 variants were subcloned into the pMal-p2G vector (New England Biolabs) using the EcoRI and HindIII restriction sites. The following universal 5’ primer was used, which contained an EcoRI site (underlined), a TEV protease cleavage site coding segment (bold), a SerGlySer linker coding segment (italic) and the first six codons of the SGPI-2 gene (italic and underlined): 5'- A CTG GAA TTC GAA AAC CTG TAT TTT CAG GGA TCC GCC GAG GTG ACC TGC GAA CCG -3'. Each 3’ primer contained a variant-specific segment and a common segment with two consecutive stop codons and a HindIII restriction site. The expression plasmid is illustrated in
Supplementary Figure S1. *E. coli* BL21 Star™ (Invitrogen) cells transformed with the appropriate expression vector were grown in 1 liter LB/ampicillin medium at 37 °C until the optical density of the culture measured at 600 nm reached 0.5 and expression was induced overnight with 0.3 mM IPTG (final concentration). Cells were harvested by centrifugation (10 min; 6000g; 4°C), resuspended in 80 mL ice cold 1 mM MgCl₂ solution and kept frozen at -20°C overnight. Next morning the suspension was thawed and cells were removed by centrifugation (10 min; 15,000g; 4°C). The supernatant containing the periplasmic fraction was treated with 20 units/mL concentration Benzonase™ (Novagen) overnight at room temperature to eliminate nucleic acid contamination. Ammonium sulfate was added to 90% saturation and precipitated proteins were recovered by centrifugation (10 min; 15,000g; 4°C). Proteins were suspended in 70 mL 2.5 mM HCl and dialyzed against the same solution twice for 2 hours each. Precipitated proteins were removed by centrifugation (10 min; 15,000g; 4°C). The pH of the supernatant was adjusted to 8.0 by adding Tris-HCl buffer to 50 mM final concentration and the MBP-SGPI-2 fusion protein was purified on a 5 mL HiTrap Q HP (GE Healthcare Life Sciences) ion-exchange column using the same buffer and a linear 0-0.5 M NaCl gradient on a Pharmacia FPLC system. MBP-SGPI-2 fusion protein was purified on a 5 mL HiTrap Q HP (GE Healthcare Life Sciences) ion-exchange column using the same buffer and a linear 0-0.5 M NaCl gradient on a Pharmacia FPLC system. The fusion protein eluted at 0.25 M NaCl buffer to 50 mM final concentration and the MBP-SGPI-2 fusion protein was purified on a 5 mL HiTrap Q HP (GE Healthcare Life Sciences) ion-exchange column using the same buffer and a linear 0-0.5 M NaCl gradient on a Pharmacia FPLC system. The fusion protein eluted at 0.25 M NaCl concentration. Two or three peak fractions (1 mL each) were pooled; and AcTEV™ protease (10,000 units/mL containing 5 mM DTT; Invitrogen) was added to attain 100 units/mL protease concentration and 50 µM final DTT concentration. The sample was incubated overnight at room temperature and completeness of the cleavage reaction was verified by SDS-PAGE and Coomassie Blue staining. The sample was then loaded on a column containing immobilized amylase (New England Biolabs) to capture the released MBP tag. The flow-through containing the SGPI-2 variant was filtered through a 22 µm membrane and was loaded onto a Jupiter C4 300A (250x10mm) RP HPLC column (Phenomenex). Solvent A was 0.1% trifluoroacetic acid (TFA) and solvent B was acetonitrile with 0.08% TFA. SGPI-2 variants eluted as symmetrical peaks and were found to be homogeneous and of correct size by mass spectrometry on an HP1100 HPLC-ESI-MS system (Agilent Technologies). Purified SGPI-2 variants were lyophilized and dissolved in 10 mM Tris-HCl (pH 8.0) before use.

Expression and purification of human pancreatic proteases. Human cationic trypsin (PRSS1), human anionic trypsin (PRSS2), chymotrypsins CTRB1, CTRB2, CTRC, CTRL1, and elastases ELA2A, ELA3A and ELA3B were expressed as inactive precursors (zymogens) either in *E. coli* BL21 (DE3) or in HEK 293T cells and were purified as previously described [7 and references therein], CTRB1, CTRB2, CTRC, CTRL1 and ELA3B zymogens were activated with immobilized bovine trypsin (Pierce; Thermo Fisher Scientific, Rockford IL) and the trypsin-containing beads were removed by centrifugation. ELA3A and ELA2A precursors were activated with human cationic trypsin and anionic trypsin, respectively, and trypsin was removed by passing the activation mixtures through a ~1 mL benzamidine column (Pharmacia). Active proteases were stored in 0.1 M Tris-HCl (pH 8.0) and 0.05% Tween-20 (final concentrations) at 4 °C. Protease concentrations were verified before each experiment either by active site titration or activity assay.

Concentration determinations. Bovine trypsin (Worthington Biochemicals) was dissolved in 0.1 M Tris-HCl (pH 8.0), 1 mM CaCl₂ at an approximately 50 µM concentration and active-site titrated using p-nitrophenyl p'-guanidinobenzoate, as described [19]. The panprotease inhibitor ecotin [20-21] was overexpressed in *E. coli* BL21 (DE3) and purified from the periplasm as described previously [22]. The concentration of ecotin was determined by titration against the active-site titrated bovine trypsin. This ecotin batch served then as a universal titrant for all human pancreatic proteases studied here. We have determined the equilibrium dissociation constants (*K*<sub>D</sub>) of ecotin with human pancreatic proteases (Supplementary Table S1) and titrations were performed using protease concentrations at least two orders of magnitude above the *K*<sub>D</sub> values, with the exception of ELA3A which was titrated at a concentration ~5-times above *K*<sub>D</sub>. The concentration of SGPI-2 and its variants was determined by titration against CTRC. Immediately before the binding
experiments, all tight-binding protease-inhibitor pairs were re-titrated against each other to ensure "self-consistent" concentrations.

**Equilibrium binding assays.** Binding of SGPI-2 variants to human chymotrypsins and elastases was characterized by determining the \( K_D \) value of the reaction in equilibrium, according to the method of the Laskowski laboratory [23]. A fixed protease concentration (between 0.5 nM and 10 nM) was reacted with increasing concentrations of the inhibitor and the free (unbound) protease concentration was determined by enzymatic assays using appropriate fluorogenic or chromogenic peptide substrates (listed in Supplementary Table S2). Inhibitors and proteases were incubated in 0.1 M Tris-HCl (pH 8.0), 1 mM CaCl\(_2\), and 0.05% Tween-20 (final concentrations) for 1 hour at 22 °C in microtiter plates in 100 µL final volume and substrates were given in 5 µL volume so as not to perturb the equilibrium. The free protease concentration was plotted as a function of the total inhibitor concentration and the experimental points were fitted with the following equation. \( y = E-(E+x+K-sqrt((E+x+K)^2-4Ex))/2 \), where the independent variable \( x \) represents the total inhibitor concentration, the dependent variable \( y \) is the free protease concentration in equilibrium; \( K \) is \( K_D \) and \( E \) designates the total protease concentration.

**Structural modeling of the CTRC:C6 complex.** Several homology models of human CTRC were constructed using the SWISS-MODEL Workspace [24] using as templates porcine elastase bound to Ascaris chymotrypsin/elastase inhibitor (PDB 1EAI chain A; 52% identity) [25], bovine chymotrypsin bound to pacifastin inhibitor PMP-C (PDB 1GL1 chain A, 36% identity) [26], and bovine chymotrypsinogen C (PDB 1PYT chain D; 80% identity) [27]. Since substantial conformational changes throughout the substrate binding cleft take place upon serine protease activation, we chose the model based on porcine elastase as the best approximation of the active CTRC structure. The three homology models were quite similar overall but differed considerably over a 5-residue loop (residues 238-242 in the human CTRC sequence) that in the elastase-based model clashed with the anticipated orientation of the C6 inhibitor; these residues were replaced with the corresponding residues from the homology model based on bovine chymotrypsinogen C. A homology model of the C6 inhibitor was constructed in the SWISS-MODEL Workspace using as a template PMP-C (PDB 1GL1 chain I; 76% identity) [26]. The homology models of human CTRC and inhibitor C6 were first docked manually based on the orientation of PMP-C and chymotrypsin in 1GL1, and then a local docking search with energy minimization of side chain rotamers at the intermolecular interface was performed using the RosettaDock server [28].

**RESULTS**

**Analysis of CTRC specificity through phage-display selection of substrate-like SGPI-2 inhibitor variants.** To identify determinants of CTRC substrate specificity, we utilized a phage-display approach, in which a library of inhibitor variants was constructed and phages carrying tight-binding inhibitors were selected. We used a chymotrypsin inhibitor from the desert locust *Schistocerca gregaria* (SGPI-2) as our scaffold for inhibitor selection (Fig 1A) [29-32]. The 35 amino acid SGPI-2, which belongs to the pacifastin inhibitor family, had already been efficiently displayed on the surface of M13 phage [14]. Wild-type SGPI-2 inhibited human chymotrypsins and elastases with \( K_D \) values ranging from 0.2 nM to 2 µM, and generally stronger inhibition was apparent for chymotrypsins (Table 2). To display SGPI-2 on the surface of the M13 phage, it was fused to the N terminus of the p8 coat protein. Six positions in the reactive loop of SGPI-2 were fully randomized, which included P4, P2, P1, P1', P2' and P4' (Fig 1A, Supplementary Fig S1A). The P3 and P3' Cys residues are structurally indispensable and these were therefore conserved. Selection of tight-binding clones was performed against immobilized human CTRC. Three selection cycles were performed and tight-binding inhibitor containing phages were verified by ELISA and subjected to DNA sequencing.

**Selected tight-binding SGPI-2 variants reveal unexpected acidic residues at P4' position of the reactive loop.** From the sequence analysis of 32 inhibitor-phage clones, 22 were unique at the DNA level, indicating that these were the results of independent selection events. The reactive loop
sequence of the individual clones is listed in Supplementary Table S3. To define the characteristic sequence pattern for CTRC inhibition, we have created a codon-normalized sequence logo from the individual sequences, in which symbol height was used to represent sequence conservation at positions P4 through P4’ (Fig 1B). The normalized sequence pattern indicated selection of Met and Leu amino acid residues at the P1 position, which corresponded to the expected P1 specificity of CTRC. The P2 position showed a strong consensus for Thr; however, this most likely reflects stabilizing interactions between the reactive loop and the inhibitor core (see Discussion). A slight predominance of Ala and Gly were detected at the P4 position, which showed the lowest degree of sequence conservation of all positions randomized. On the prime side of the reactive loop, both the P1’ and P2’ positions showed a clear preference for hydrophobic side chains, Met and Leu, respectively. Interestingly, acidic amino acid residues Asp and Glu were also selected at position P2’, although with a lower frequency than Leu. Finally, the most striking feature of the sequence logo was the almost exclusive presence of acidic Glu and Asp amino acid residues at position P4’. Taken together, the phage-display results suggest that the sequence pattern of the reactive loop of SGPI-2 variants that most likely bind to CTRC with high affinity is Xaa-Cys-Thr-Met/Leu-Met-Leu-Cys-Glu/Asp.

Phage-display evolved CTRC inhibitors exhibit improved binding affinity relative to parent SGPI-2. On the basis of the reactive loop sequence pattern derived from tight-binding inhibitor-phage clones, we have designed six SGPI-2 inhibitor variants, in which positions P1 (Met or Leu), P2’ (Leu or Asp) and P4’ (Glu or Asp) were varied, while positions P4 (Ala), P2 (Thr) and P1’ (Met) were fixed (see Fig 1C). In addition, to test the contribution of the P4’ acidic side chain to binding affinity, we designed two inhibitors with P4’ Ala (Fig 1C). The eight inhibitors were expressed in E. coli, purified to homogeneity and their binding to CTRC was characterized by determining the equilibrium dissociation constants ($K_D$). Representative binding experiments are shown in Fig 2A and the $K_D$ values for CTRC are demonstrated by Fig 2B. $K_D$ values for the eight inhibitors were also measured with the full complement of human pancreatic chymotrypsins and elastases (Table 2). To better illustrate the selectivity of the inhibitors, $K_D$ values were also expressed after normalization to the CTRC $K_D$ which was set to equal one (Table 2). Full characterization of wild-type SGPI-2 and phage-display selected variants required 88 independent $K_D$ determinations in the present work.

As expected, compared to the inhibition of CTRC by the parent molecule SGPI-2 ($K_D$ 11.5 nM, see Table 2), all eight SGPI-2 variants exhibited significantly higher binding affinities, with $K_D$ values ranging from 0.02 nM to 0.9 nM, which corresponded to 13-fold to 575-fold improvement. Interestingly, relative to inhibition by the parent molecule SGPI-2, chymotrypsins CTRB1, CTRB2 and CTRL1 were inhibited by the CTRC-selected inhibitors less strongly, whereas inhibition of elastases ELA2A, ELA3A and ELA3B showed a tendency for improvement. With respect to binding affinity, the best CTRC inhibitor was C4 ($K_D$ 20 pM), which also exhibited good selectivity (range 46-12,650-fold) against other pancreatic chymotrypsins and elastases. With respect to selectivity, the best inhibitor was C6, with a $K_D$ value of 110 pM and a selectivity that ranged from 225-fold to 112,664-fold (Table 2).

Leu at P1 affords tighter binding than Met. According to the sequence logo, Met or Leu at P1 should result in comparable inhibition of CTRC. Surprisingly, however, when purified inhibitors with P1 Leu or Met were compared, inhibitors with Leu consistently showed stronger inhibition of CTRC by about an order of magnitude (average 12-fold; range 8-16-fold) (cf. C1 with C4, C2 with C5 and C3 with C6 in Fig 2B, Fig 3 and Table 2). CTRB2 exhibited a similarly strong P1 preference for Leu and the same general trend was apparent for other chymotrypsins and elastases, although the magnitude of the difference between inhibition by Leu and Met P1 was typically smaller than seen with CTRC (Fig 3, Table 2). Consequently, relative to Met, the Leu P1 not only increased affinity for CTRC but also increased the selectivity of the CTRC inhibitors against other chymotrypsins and elastases by an average of 4-fold (range 1-12-fold) (Table 2). Interestingly, the
presence of Asp (versus Leu) at the P2' position mitigated the Leu P1 preference of CTRC and CTRB2 by 2-fold and 5-fold, respectively (cf. Fig 3C with Fig 3A and B).

**Acidic amino acid residues at P4' markedly increase inhibitor affinity to CTRC.** The essentially exclusive selection of Glu and Asp residues at the P4' position in tight-binding inhibitor-phage clones suggested that acidic residues are important determinants of inhibitor and substrate binding to CTRC. In agreement with these expectations, the equilibrium binding experiments demonstrated that CTRC binds inhibitors with P4' Glu circa an order of magnitude stronger (average 13-fold, range 8-18-fold) than inhibitors with P4' Ala (cf. C4 with C7 and C6 with C8 in Fig 4 and Table 2). In contrast, other chymotrypsins generally bound inhibitors with P4' Glu more weakly than those with P4' Ala, whereas in the case of elastases the Glu versus Ala exchange at P4' had no effect on inhibitor binding (Fig 4). Binding affinities of inhibitors with P4' Glu or Asp were comparable (cf. inhibitors C1 with C2 and C4 with C5 in Table 2), although inhibitors with P4' Asp showed a tendency toward slightly weaker binding, but the difference was within experimental error. Interestingly, the P2' position had a small, but measurable influence on the effect of the P4' position on chymotrypsins (cf. C4 with C6 and C7 with C8 in Fig 4). The P2' Asp (versus Leu) reduced the affinity-increasing effect of the P4' Glu (versus Ala) toward CTRC by about 2-fold, whereas it enhanced the unfavorable effect of the P4' Glu on binding affinity toward CTRB1 and CTRB2 by about 3-fold.

Because the positive effect of the P4' Glu/Asp on inhibitor binding is highly specific for CTRC, this residue is also the most important determinant of inhibitor selectivity against other pancreatic chymotrypsins and elastases. CTRC inhibitors with P4' Glu (versus Ala) are on average 10-fold (range 6-13-fold) more selective against elastases and CTRB2 than inhibitors with P4' Ala (cf. C1 with C3, C4 with C6 and C7 with C8 in Fig 5 and Table 2). The P2' Asp decreases binding affinity but increases selectivity of SGPI-2 variants. The phage-selected sequence pattern suggests that the optimal residue at P2' for CTRC binding is Leu. However, contrary to our expectations, acidic residues (Asp or Glu) were also selected at this position. This observation suggested that acidic residues at P2' offer a non-homologous alternative solution for binding to the S2' site of CTRC without significant loss of affinity. Binding experiments with purified inhibitors confirmed that, relative to a P2' Leu, the presence of P2' Asp caused only an average of 3.6-fold (range 2.5-5.5-fold) decrease in binding affinity toward CTRC (cf. C1 with C3, C4 with C6 and C7 with C8 in Fig 5 and Table 2). Binding to CTRB1 and CTRB2 was similarly affected. In contrast, the P2' Asp caused a much more significant (on average ~25-fold, range 5-49-fold) decrease in the inhibition of elastases and CTRL1 (Fig 5). Owing to this difference, the P2' Asp appreciably (on average ~7-fold, range 2-17-fold) increased the selectivity of the CTRC inhibitors against elastases and CTRL1 (Table 2).

**Modeling suggests that a cluster of positively charged CTRC residues interact with the acidic P4' residue of the inhibitor.** The most intriguing finding from the phage display selection was the preference of CTRC for acidic residues at the P4' position; this preference is unique among pancreatic proteases. To gain insight into potential structural determinants of this aspect of CTRC specificity, we generated homology models of CTRC (modeled on porcine elastase and bovine CTRC zymogen) and the C6 inhibitor (modeled on PMP-C, a homologous inhibitor from *Locusta migratoria*). We docked the CTRC:C6 model complex using a local docking search with energy minimization of side chain rotamers at the intermolecular interface. The resultant C6 inhibitor orientation and placement of residues occupying the P4-P4' positions closely mimic those observed in the crystal structure of the complex formed by bovine chymotrypsin with PMP-C (PDB 1GL1) [26], lending support to the validity of our model. In examining the environment surrounding the P4' Glu residue in the model, we observed a cluster of basic residues that apparently interact with and stabilize the negative charge borne by this residue (Fig 6). In the model, the side chain carboxylate oxygens of the P4' Glu form salt bridges with Lys51 and Arg80, with distances of about 2.6 and 3.4 angstroms, respectively, and the backbone carbonyl oxygen of the P4' Glu accepts an H-bond...
from nearby Arg\textsuperscript{56}. We hypothesize that these interactions contribute to the preference of CTRC for acidic residues at the P4' position.

Beyond the P4' position, we examined the CTRC:C6 model for insights into binding specificity at additional prime-side subsites of CTRC which differ from those of typical chymotrypsins and elastases. At the P1' position, mammalian chymotrypsins typically show a preference for basic residues Arg and Lys, which has been attributed to interactions with conserved acidic residues Asp\textsuperscript{35} and Asp\textsuperscript{64} (chymotrypsin numbering) \[33\]. In human CTRC, each of these Asp residues is substituted with the bulkier, uncharged Tyr, potentially explaining a preference for the shorter, hydrophobic side chain of Met at the P1' position. At the P2' position, human CTRC proved to be unexpectedly permissive of acidic side chains Asp and Glu; this may be due to the positioning of the bound P2' residue between the basic side chains of Arg\textsuperscript{56} (distance ~5 Å) and Arg\textsuperscript{52} (distance ~6 Å). Although these distances may not suggest particularly close ion pair contacts, it has been well documented that electrostatic interactions in the range of 4-7 Å can contribute substantially to the energetics of protein-protein association, depending upon the local environment \[34\].

### Mutational analysis of CTRC residues Lys\textsuperscript{51}, Arg\textsuperscript{56} and Arg\textsuperscript{80} supports their interaction with the acidic P4' residue of the inhibitor.

To test the predictions of our CTRC model with respect to the binding of the P4' Glu amino acid residue of inhibitors, we mutated Lys\textsuperscript{51} and Arg\textsuperscript{56} to Ala, individually (single mutants K51A and R56A) and in combination (double mutant K51A,R56A). We also mutated Arg\textsuperscript{80} to Leu (mutant R80L). Leu was used at this position because in most mammalian CTRC molecules Arg\textsuperscript{80} is not conserved and Leu is found in its place. Finally, to test the effect of an unfavorable negative charge within this basic amino acid cluster, Arg\textsuperscript{56} was changed to Glu (mutant R56E). The CTRC mutants were expressed and purified and kinetic parameters were determined on the peptide substrate Suc-Ala-Ala-Pro-Phe-p-nitroanilide. As shown in Supplementary Table S4, the \(k_{cat}, K_M\) and \(k_{cat} / K_M\) values for the six mutants were comparable to those of wild-type CTRC, indicating that the mutations did not perturb the catalytic machinery of CTRC to a significant extent. Binding of the wild-type SGPI-2 inhibitor by the CTRC mutants was slightly improved (mutants K51A, R56A, and K51A,R56A) or unchanged (mutant R56E), with the exception of the R80L mutant, which bound SGPI-2 almost 10-fold stronger than wild-type CTRC (Supplementary Table S5). This latter effect is probably due to the elimination of a repulsive interaction between Arg\textsuperscript{80} on CTRC and the P1' Lys residue in SGPI-2 (not shown).

To determine whether the three mutated basic residues contribute to binding the P4' Glu of CTRC inhibitors, we compared binding of two inhibitor pairs (C4-C7 and C6-C8) differing only in the P4' position (Glu versus Ala). Table 3 lists the \(K_D\) values measured. The single neutral mutations (K51A, R56A, R80L) in CTRC decreased binding affinity to inhibitors with P4' Glu by an average of 6-fold (range 4-9-fold), and this effect was additive in the double mutant K51A,R56A, which caused an average of 31-fold (range 21-40-fold) decrease. Binding of these CTRC mutants to inhibitors carrying a P4' Ala was only slightly weakened (on average 1.7-fold, range 0.4-4-fold), and in some cases the mutations actually improved affinity (R80L with C7 and C8, and K51A,R56A with C7, see Table 3). Similarly, the negative charge of the R65E mutant more significantly compromised binding to inhibitors with a P4' Glu (average 33-fold, range 21-46-fold) than those with Ala (average 6-fold, range 4-7-fold). To visualize the effects of the mutations on the P4' Glu specificity, the ratio of the \(K_D\) values for inhibitors containing Ala versus Glu at P4' were graphed. Fig 7 demonstrates that individual mutations of Lys\textsuperscript{51} and Arg\textsuperscript{56} decreased, while double-mutation of these residues (K51A,R56A) or mutation of Arg\textsuperscript{80} alone (R80L) abolished the P4' Glu specificity. Introduction of a negative charge in place of Arg\textsuperscript{56} (mutation R56E) also reduced the P4' Glu preference of CTRC. Taken together, the mutagenesis results are consistent with the predictions of the modeling and confirm that Lys\textsuperscript{51}, Arg\textsuperscript{56} and Arg\textsuperscript{80} participate in stabilizing interactions with the P4' Glu of phage-selected CTRC inhibitors.

### DISCUSSION

Recent studies highlighted human CTRC as a unique digestive enzyme with the ability to...
regulate the activity levels of other digestive proteases, trypsins and carboxypeptidases in particular, through promotion of their activation or degradation [1, 2, 7]. These regulatory functions, reviewed in detail in the introduction to this paper, are highly specific to CTRC as other human pancreatic chymotrypsins and elastases exhibit no such activity. The CTRC specificity is partly explained by the known affinity of CTRC for leucyl peptide bonds [8-10]. However, taking into account that Leu is the most abundant amino acid in proteins, other recognition determinants in the regulatory CTRC cleavage sites should be also relevant. In this respect, alignment of sequences flanking the regulatory cleavage sites in human cationic trypsinogen (Phe\textsuperscript{8}-Asp\textsuperscript{19} and Leu\textsuperscript{81}-Glu\textsuperscript{97}) and in human proCPA1/proCPA2 (Leu\textsuperscript{96}-Leu\textsuperscript{97}) reveals a clustering of acidic amino acid residues on the prime side of the scissile peptide bonds, from P1' through P4' (Table 1). To define the cleavage specificity of CTRC and thereby gain further insight into its physiological function, in the present study we utilized a phage-display approach to select tight-binding substrate-like chymotrypsin inhibitor variants. To borrow the title of a recent comprehensive review by Zani and Moreau (2010), "phage display is a powerful tool to engineer protease inhibitors"; either to gather information on protease specificity or to generate specific reagents for inhibition studies [35]. Here, we used SGPI-2, an arthropod-derived small chymotrypsin inhibitor from the pacifastin family [29-32] as a scaffold and fully randomized six positions in the reactive loop, spanning from P4 through P4', with the exception of the structurally essential P3 and P3' Cys residues (see Fig 1).

The results of the phage-display selection against CTRC yielded SGPI-2 variants with a reactive loop sequence pattern that identified the amino acid preference of CTRC at five of the six positions randomized, i.e. at P4, P1, P1', P2' and P4'. The sole exception was position P2, where a Thr residue was selected predominantly. The P2 Thr is a characteristic feature of many canonical protease inhibitors and it is also strictly conserved in the pacifastin family [32, 33]. Both NMR and crystal structures demonstrated that in SGPI homologues the P2 Thr in the reactive loop and Asn\textsuperscript{15} in a supporting loop participate in a stabilizing H-bond network [26, 37-45]. Consequently, conservation of the P2 Thr upon phage display evolution simply recapitulates natural evolution and does not represent the selection of a true protease specificity determinant. From the five positions allowing insight into CTRC specificity, four (P1, P1', P2' and P4') demonstrated a clear predilection toward certain amino acid residues, whereas selection at position P4 was diverse, with only a small preference for small amino acid residues Ala and Gly. On the basis of the sequence pattern selected we have designed and expressed eight SGPI-2 variants and studied their binding to CTRC and a panel of human chymotrypsins and elastases. All eight variants showed significantly improved binding to CTRC relative to the parent SGPI-2 molecule. The C4 variant with the highest affinity ($K_D$ 20 pM) represented a 575-fold improvement over wild-type SGPI-2.

With respect to the primary (P1) specificity determinant, the large majority of selected phage clones carried SGPI-2 variants with Met or Leu residues at this position. These two residues are characteristically found in the reactive site of naturally occurring chymotrypsin inhibitors. In a now classic project, the Laskowski laboratory purified and sequenced ovomucoid third domains from 153 bird species, and in the P1 position of the reactive loop found Met and Leu 60 and 66 times, respectively [46]. Nevertheless, when we compared purified SGPI-2 variants in direct binding experiments, the inhibitors containing a P1 Leu bound to human CTRC with an order of magnitude higher affinity than those with P1 Met (see Fig 3). Thus, Leu is the preferred amino acid residue at the P1 position for human CTRC, which is consistent with the known activity of porcine and bovine CTRC on leucyl peptide bonds [8-10]. The comparable selection of phage clones with Met and Leu at P1 suggests that a binding equilibrium has not been reached in our selection experiments. Several phage display studies established that ratios of normalized amino acid frequencies correlate with affinity ratios of the corresponding variants [47-50]. However, this correlation applies only when the selection process reaches binding equilibrium, which for high affinity variants with slow off rates may take longer than an experimentally practical time frame.

On the prime side of the reactive loop of selected SGPI-2 variants hydrophobic amino acid
residues Met and Leu dominated at the P1' and P2' positions, respectively. The P1' preference of human CTRC resembles that of trypsin rather than chymotrypsin. Thus, bovine cationic and rat anionic trypsin prefer Met side-chains at P1', whereas bovine and rat chymotrypsin prefer positively charged Arg or Lys side chains at this position [30, 34, 51-54]. The characteristic Arg/Lys P1' preference of chymotrypsin is attributed to electrostatic interactions with two negatively charged residues, Asp \(^{35}\) and Asp \(^{64}\) (chymotrypsin numbering), which are not conserved in human CTRC [34]. The selection of Leu at P2' matches the published P2' preference of rat chymotrypsin determined from measurements of acyl-transfer to pentapeptide nucleophiles [51]. In addition to the preferred Leu, some of the phage clones selected contained Asp or Glu at the P2' position of the SGPI-2 reactive loop (see Fig 1B and Supplementary Table S3). This finding was unexpected, as the aforementioned acyl-transfer experiments indicated that acidic side-chains at P2' are poorly recognized by rat chymotrypsin or rat anionic trypsin [52], but was consistent with potentially stabilizing electrostatic interactions with Arg\(^{56}\) and Arg\(^{162}\) identified in our structural model. Binding experiments with SGPI-2 variants carrying a P2' Asp revealed that binding affinity to CTRC was only moderately compromised (on average ~3.6-fold versus Leu P2') but the P2' Asp increased the selectivity of the inhibitors more significantly, particularly against pancreatic elastases (see discussion on selectivity below).

Arguably, the most exciting observation from the present study was the highly restricted selection of acidic Asp and Glu amino acid residues at the P4' position of the reactive loop of SGPI-2 variants. Binding experiments comparing inhibitors with P4' Glu versus Ala demonstrated that the acidic side chain at P4' increases binding affinity by an order of magnitude on average. Furthermore, inspection of known regulatory cleavage sites for human CTRC in digestive proenzymes reveals that an acidic residue at P4' is present in all four sequences (Table 1). Taken together, the observations strongly suggest that an interaction between CTRC and the P4' position of substrates and inhibitors is a major determinant of CTRC recognition and specificity. This property of CTRC appears unique among the pancreatic digestive proteases, as no P4' preference has been described for trypsin, chymotrypsin or pancreatic elastase so far. Interestingly, a similar but less pronounced acidic P4' preference was reported for human granzyme B, a serine protease found in the granules of cytotoxic lymphocytes [55, 56].

In the reactive loop of natural canonical inhibitors that inhibit pancreatic proteases the presence of an acidic Glu or Asp at P4' is unusual, and Pro is often found here. Among the 153 ovomucoid third domains isolated by Laskowski the Pro 4' is strictly conserved [46]. Within the 20 known members of the pacifastin family Pro occupies the P4' position in eight inhibitors (including SGPI-2), other hydrophobic residues (Leu, Ile, Val or Ala) are present in seven cases, while Arg is found in four members and Gln in one [32]. Examination of crystal structures of pacifastin inhibitors bound to serine proteases reveals that in the complex between bovine chymotrypsin and PMP-C a P4' Pro stacks against Phe\(^{39}\) (chymotrypsin numbering) of the enzyme [26]. The same packing interaction was found between P4' Pro of SGTL and Phe\(^{39}\) (chymotrypsin numbering) of crayfish trypsin [42, 44]. In the complex between Fusarium oxysporum trypsin and LMPI-3 the P4' Arg participates in an H-bond with Asn\(^{37}\) (chymotrypsin numbering) of the enzyme [45]. On the other hand, in the complex of bovine chymotrypsin and a P1'-P1' mutant of PMP-D2, in which the P4' residue is Gln, there is no detectable P4'-S4' interaction [26]. These structural examples indicate that the P4' position can contribute to inhibitor binding, although functional studies confirming the significance of the P4'-S4' interactions in these complexes are lacking. We also note that the interacting loop of SGPI-2 is flexible [41], and the P4' position, being the penultimate residue in SGPI-2, may be conformationally less constrained than on other canonical inhibitors.

To identify the amino acid residues in human CTRC that interact with a P4' Glu/Asp side chain, we created a model of the CTRC:C6 inhibitor complex using homology modeling. On the basis of this model, we then subjected three positively charged residues (Lys\(^{31}\), Arg\(^{56}\) and Arg\(^{80}\)) to mutagenesis and confirmed that this cluster of basic residues on the surface of CTRC is responsible for the observed acidic preference at P4'. Interestingly, Arg\(^{80}\) is not conserved in the bovine, rat, mouse, horse and dog CTRC, whereas
pig CTRC lacks Lys51, suggesting that the strong P4' Glu/Asp reference may be a distinctive property of the human enzyme.

The SGPI-2 variants selected against CTRC exhibited not only high binding affinity but also good selectivity against other pancreatic chymotrypsins and elastases. In this respect, the best inhibitor was C6, which was from 225-fold to 112,664-fold more selective toward CTRC than other proteases. The highest selectivity was observed against ELA3A, whereas the lowest was against CTRL1 and ELA2A. Selectivity was determined by the amino acids at the P1, P2' and P4' positions of the reactive loop. The P1 Leu (versus Met) increased selectivity by an average of 4-fold. The P2' Asp (versus Leu) increased selectivity on average 7-fold against elastases and CTRL1 but it had only a small impact on selectivity against CTRB1 and CTRB2. Finally, the most important determinant of selectivity was the P4' Glu amino acid residue, which increased selectivity (over P4' Ala) on average by 10-fold against elastases and 55-fold against chymotrypsins.

In summary, we selected novel small protein inhibitors against human CTRC using a phage-display approach and identified a functionally important interaction between the acidic P4' amino acid residue of inhibitors and a basic amino acid cluster in CTRC. The high affinity and strong selectivity of the inhibitors developed in the present study opens up new avenues for experiments on the physiological function of CTRC in digestion and on the pathological role of CTRC mutants in the development of chronic pancreatitis.

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REFERENCES

LEGENDS TO FIGURES

Figure 1. Phage-display selected *Schistocerca gregaria* chymotrypsin inhibitor (SGPI-2) variants. A. Primary structure of wild-type SGPI-2. The amino acid positions within the reactive loop that were randomized in the phage display library are highlighted as white letters on a black background. The disulfide bonds are indicated. B. Sequence logo representation of the reactive loop of phage-selected CTRC inhibitors. The logo was generated with the Weblogo program (http://weblogo.berkeley.edu/). The overall height of the stack of symbols indicates the sequence conservation at that position, while the height of symbols within the stack indicates the relative frequency of each amino acid. The colors indicate the chemical properties of the amino acid side-chains; aliphatic is green, aromatic is orange, acidic is red, basic is dark blue, and polar with no charge is light blue. Note that the Cys residues shown in gray at the P3 and P3’ positions were not randomized. Their symbol height represents the maximal height corresponding to a completely conserved residue. C. Reactive loop sequences of eight SGPI-2 variants designed to test the significance of the P1 (Met or Leu), P2’ (Leu or Asp) and P4’ (Glu, Asp or Ala) positions in binding affinity. The corresponding P4-P4’ segment in wild-type SGPI-2 is indicated in panel A. These eight inhibitors as well as wild-type SGPI-2 were expressed, purified and their inhibitory activity was tested on human pancreatic chymotrypsins and elastases. See text for further details.

Figure 2. Inhibition of human CTRC by phage-display selected SGPI-2 variants. A. Representative inhibitor-binding experiments with the C4 and C6 SGPI-2 variants using 0.5 nM CTRC concentration. B. Equilibrium dissociation constant (K_D) values determined for eight CTRC-specific inhibitors. Measurements were carried out as described in Experimental Procedures.

Figure 3. The effect of the P1 residue (Leu or Met) on the inhibition of human pancreatic chymotrypsins and elastases by phage-display selected SGPI-2 variants. Pairwise comparison of relative K_D values (P1 Leu set to equal one) is shown for inhibitor pairs differing only at the P1 position. The amino acid residues found at P2' and P4' are also indicated.

Figure 4. The effect of the P4’ residue (Glu or Ala) on the inhibition of human pancreatic chymotrypsins and elastases by SGPI-2 variants. Pairwise comparison of relative K_D values (P4’ Glu set to equal one) is shown for inhibitor pairs differing only at the P4’ position. The amino acid residues found at P1 and P2’ are also indicated.

Figure 5. The effect of the P2’ residue (Leu or Asp) on the inhibition of human pancreatic chymotrypsins and elastases by phage-display selected SGPI-2 variants. Pairwise comparison of relative K_D values (P2’ Leu set to equal one) is shown for inhibitor pairs differing only at the P2’ position. The amino acid residues found at P1 and P4’ are also indicated.

Figure 6. Structural modeling predicts specificity determinants. A. Overview of homology model of the CTRC:C6 complex. CTRC is surface-rendered in gray, with a cluster of three basic residues surrounding the P4’ binding site highlighted in blue (nitrogens) and yellow (carbons); the inhibitor is shown in orange. B. Enlarged view of the P4’ binding site, showing interactions of Lys^{51}, Arg^{56}, and Arg^{80} with the P4’ Glu residue of the C6 inhibitor.

Figure 7. Effect of CTRC mutations on binding to SGPI-2 variants differing in the P4’ position only (Ala versus Glu). Equilibrium dissociation constants (K_D) were determined for the C7/C4 and C8/C6 inhibitor pairs, as described in Experimental Procedures. The ratio of the K_D values (Ala over Glu ) was plotted. A high ratio signifies preference for Glu-containing inhibitors, whereas a low ratio indicates loss of P4’ specificity. The dotted line indicates a ratio of one, with no discrimination between inhibitors containing Ala or Glu at P4’ in their reactive loop.
Table 1. Sequence context of CTRC-specific regulatory cleavage sites in human cationic trypsinogen and procarboxypeptidases A1 and A2. The P1 residues are highlighted in bold.

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<th>CTRC cleavage sites</th>
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<td>-Gln^{94}-Val^{95}-<strong>Leu^{96}</strong>-Leu^{97}-Asp^{98}-Lys^{99}-Glu^{100}--</td>
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Table 2. Inhibition of human pancreatic chymotrypsins and elastases by *Schistocerca gregaria* chymotrypsin inhibitor (SGPI-2) and phage-display selected SGPI-2 variants. Equilibrium dissociation constants ($K_D$) shown in bold on gray background were determined as described in *Experimental Procedures* and expressed in nM units of concentration. To quantify the selectivity of the inhibitors, $K_D$ values were normalized to the CTRC $K_D$ which was set to equal one. These selectivity values are shown in italics underneath their corresponding $K_D$. Values above one were rounded to the nearest whole number. See Fig 1 for the primary structure of the inhibitors used.

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Table 3. Effect of mutations of Lys$^{51}$, Arg$^{56}$ and Arg$^{80}$ in CTRC on the binding of SGPI-2 variants. Equilibrium dissociation constants ($K_D$) were determined as described in Experimental Procedures and expressed in nM units of concentration. See Fig 1 for the primary structure of the inhibitors used.

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Figure 1
Figure 2
Figure 3

A

- C4 inhibitor, **P1 Leu**, P2’ Leu, P4’ Glu
- C1 inhibitor, **P1 Met**, P2’ Leu, P4’ Glu

B

- C5 inhibitor, **P1 Leu**, P2’ Leu, P4’ Asp
- C2 inhibitor, **P1 Met**, P2’ Leu, P4’ Asp

C

- C6 inhibitor, **P1 Leu**, P2’ Asp, P4’ Glu
- C3 inhibitor, **P1 Met**, P2’ Asp, P4’ Glu
Figure 4
Figure 5

A. C1 inhibitor, P1 Met, P2' Leu, P4' Glu
C3 inhibitor, P1 Met, P2' Asp, P4' Glu

B. C4 inhibitor, P1 Leu, P2' Leu, P4' Glu
C6 inhibitor, P1 Leu, P2' Asp, P4' Glu

C. C7 inhibitor, P1 Leu, P2' Leu, P4' Ala
C8 inhibitor, P1 Leu, P2' Asp, P4' Ala
Figure 6
Figure 7

- C7/C4 ratio, P2' Leu, P4' Ala versus Glu
- C8/C6 ratio, P2' Asp, P4' Ala versus Glu