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An Episulfide Cation (Thiiranium Ring) Trapped in the Active Site of HAV 3C Proteinase Inactivated by Peptide-based Ketone Inhibitors

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We have solved the crystal and molecular structures of hepatitis A viral (HAV) 3C proteinase, a cysteine peptidase having a chymotrypsin-like protein fold, in complex with each of three tetrapeptidyl-based methyl ketone inhibitors to resolutions beyond 1.4 Å, the highest resolution to date for a 3C or a 3C-Like (e.g. SARS viral main proteinase) peptidase. The residues of the β-hairpin motif (residues 138–158), an extension of two β-strands of the C-terminal β-barrel of HAV 3C are critical for the interactions between the enzyme and the tetrapeptide portion of these inhibitors that are analogous to the residues at the P4 to P1 positions in the natural substrates of picornaviral 3C proteinases. Unexpectedly, the Sγ of Cys172 forms two covalent bonds with each inhibitor, yielding an unusual episulfide cation (thiiranium ring) stabilized by a nearby oxyanion. This result suggests a mechanism of inactivation of 3C peptidases by methyl ketone inhibitors that is distinct from that occurring in the structurally related serine proteinases or in the papain-like cysteine peptidases. It also provides insight into the mechanisms underlying both the inactivation of HAV 3C by these inhibitors and on the proteolysis of natural substrates by this viral cysteine peptidase.

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Introduction

Hepatitis A virus belongs to the Picornaviridae, a large family of positive single-stranded RNA viruses that also includes poliovirus (PV), foot-and-mouth disease virus (FMDV) and human rhinoviruses (HRV).¹ Based on the extensive similarity in their genomic organization, it is likely that the basic features of their viral life cycles in the cell are also similar. Upon entry into the cytoplasm of susceptible cells, the viral RNA genome is translated by host ribosomes to produce a large polyprotein. Subsequent proteolytic cleavage of this polyprotein by virally encoded peptidase(s) (and in some cases by an unknown host peptidase) leads to mature viral proteins that perform key functions such as replication of the viral RNA genome, assembly of progeny virions and virus–host antagonism. The latter include inhibition of cellular mRNA translation and transcription, inhibition/induction of apoptosis, and membrane vesiculization.

In picornaviruses, most of the polyprotein is processed by the virus’s 3C peptidase, although a 2A peptidase (entero- and rhinoviruses) and a small leader proteinase (aphthoviruses) participate to a minor extent.² 3C and 3C-like (3CL) peptidases also occur in caliciviruses and coronaviruses; they play

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Abbreviations used: HAV, hepatitis A virus; FMDV, foot-and-mouth disease virus; TGEV, transmissible gastroenteritis coronavirus; SARS, severe acute respiratory syndrome; BBL, carboxylbenzyloxyl-L-serine-β-lactone; CMK, chloromethylketone; FMK, fluoromethylketone; Ac, acetyl; Qmm, N,N-dimethyl glutamine.

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major roles in polyprotein processing in these viruses, although the coronavirus polyprotein is considerably larger. These enzymes are cysteine peptidases with folds also similar to that of chymotrypsin rather than that of papain. 3CL peptidase has an additional domain that mediates homodimerization, which is thought to be critical for the enzymatic activity of 3CL. The catalytic mechanisms of 3C and 3CL are thought to proceed via a tetrahedral intermediate resulting from the nucleophilic attack on the carbonyl carbon of the scissile bond by the $\delta^\gamma$ of the catalytic cysteine residue (Cys172 in HAV 3C). However, many mechanistic details remain unknown. Detailed understanding at atomic resolution of how 3C enzymes cleave their cognate substrates would greatly facilitate the rational design of antiviral drugs. Such knowledge will help to control these dangerous pathogens in the event of a pandemic outbreak, such as the recent FMDV epidemic in England that caused significant loss of animals and the accompanying financial woes.

The use of substrate analogue inhibitors to probe enzymatic mechanisms has a long tradition in research on proteinases. Peptidyl chloromethyl ketones react with papain and serine proteinases in dissimilar ways to yield different types of covalent adducts with the catalytic residues. In chymotrypsin, Ser195 forms a covalent linkage to the carbonyl carbon of chloromethyl ketone inhibitors, resulting in a tetrahedral geometry thought to mimic the transition state that occurs during the hydrolysis of natural substrates. In addition, the $\delta^\gamma$ atom of His57 attacks the halogenated $\alpha$-carbon yielding a second covalent attachment between the inhibitor and the now alkylated enzyme. When the chloromethyl function is replaced with a chloroethyl group in the inhibitor, His57 is alkylated with retention of configuration at the chiral center. Together with kinetic studies, this indicates an internal nucleophilic displacement of chloride by the newly formed oxyanion in the tetrahedral intermediate leading to a Ser195-epoxy ether adduct that is opened by the histidine. In contrast, papain is inactivated by direct covalent linkage between the $\delta^\gamma$ atom of Cys25 and the halogenated $\alpha$-carbon of the same class of inhibitors. The imidazole ring of His159 in papain is implicated in protonating the leaving group due to its shift in orientation from a position in-plane with the $\delta^\gamma$ atom of Cys25 to a position in-plane with the $\alpha$-carbon of the inhibitor, an atom equivalent to the N of a scissile bond. However, it is unclear whether reversible formation of a tetrahedral intermediate occurs prior to alkylation of Cys25.

Sequence analyses indicate that residues His44, Asp84 and Cys172 in HAV 3C should form the canonical catalytic triad. However, this was not confirmed structurally until the hydrogen bond between $\alpha^\gamma$ of Asp84 and $\delta^\gamma$ of His44 was observed in the recently crystal structures of HAV 3C in complex with a $\beta^\gamma$-lactone (BBL, N-benzoylxylocarbonyl (Cbz) L-serine-$\beta^\gamma$-lactone) inhibitor. Fortuitously, in these crystals, the inhibitor BBL binds covalently to His102, a surface residue near the RNA binding motif of 3C, while leaving the proteolytic active site and the catalytic Cys172 residue unobstructed. Here we report the crystal structures of HAV 3C in complex with each of three tetrapeptidyl inhibitors: N-acetyl-leucyl-alanyl-alanyl-(N,N-dimethyl)-glutamine-fluoromethylketone (Ac-LAAQ$^{\gamma^\gamma}$-FMK, inhibitor Ia), N-acetyl-leucyl-phenylalanyl-phenylalanyl-glutamate-fluoromethylketone (Ac-LFFE-FMK, inhibitor II) and N-acetyl-leucyl-alanyl-alanyl-(N,N-dimethyl)-glutamine-(1,4-dioxo-3,4-dihydro-1H-phthalazin-2-yl) methylketone (Ac-LAAQ$^{\gamma^\gamma}$-pMK, inhibitor Ib) (Figure 1). These complexes were obtained by a soaking method using the pre-grown crystals of HAV 3C-BBL.

The peptidyl portion of inhibitors Ia and Ib offers a fairly good resemblance to the most favored sequence (P4 to P1 positions) of the cleavage sites by HAV 3C in the viral polyprotein. Inhibitor II was included in this study as a mimic of a less optimal substrate than the LAAQ sequence in inhibitors Ia and Ib, mainly due to the substitutions at the P1 (Gln->Glu) and P2 (Ala->Phe) positions of its peptidyl portion. Sequence alignment analysis of the naturally occurring cleavage sites for HAV 3C protease in the hepatitis A viral polyprotein indicates a clear preference for small residues such as serine and threonine at the P2 position. Available structural information strongly supports this conclusion. Furthermore, glutamate does occasionally occur as the P1 residue at the natural cleavage sites within picornaviral polyproteins, although at a much lower frequency than glutamine. Here, we are interested in understanding how the interactions between the HAV 3C enzyme and peptidyl inhibitors are influenced by the amino acid make-up of the inhibitors. The substrate recognition sites (S1 to S4) of HAV 3C are identified crystallographically. No covalent linkage occurred between His44 and any of the inhibitors. More interestingly, an unusual episulfide cation (thiiranium ring) is observed at the catalytic cysteine residue.

Results and Discussion

Crystallization and enzyme–inhibitor complex formation

Native and variant forms of HAV 3C protein have been crystallized in several different crystal forms. Until the recently reported crystallization of a Cys24Ser variant of HAV 3C in space group P2$_1$2$_1$2$_1$, there was no structural evidence that the three catalytic residues in the proteolytic active site, His44, Asp84 and Cys172, form the canonical triad assembly as observed in chymotrypsin-like serine proteinases and other picorna-viral 3C enzymes. Furthermore, the earlier crystal structures were of inactivated forms of HAV 3C containing either a substituted catalytic cysteine
residue (Cys172Ala) or an oxidized form of Cys172, thereby complicating the study of enzyme-substrate/inhibitor interactions. Indeed, soaking of substrate-like inhibitors into pre-grown crystals of these active site variants of HAV 3C has met with little success. Here, HAV 3C crystals formed after incubating the Cys24Ser variant with BBL were used for the subsequent binding of three tetrapeptidyl-based inhibitors (Figure 1). Analysis of the soaked crystals by mass spectrometry revealed a difference in mass between 3C-BBL and 3C-BBL-inhibitor Ib, that is within experimental error of the calculated molecular mass of acetyl-LAA(N,N-dimethyl)Q-methylene (data not shown). This suggests that the phthalhydrazide group is lost during the covalent attachment of inhibitor Ib to the HAV 3C enzyme in a fashion similar to the loss of fluoride from the other two inhibitors.  

A previous study of the in vitro inhibition of HAV 3C by inhibitor Ia showed an IC₅₀ value of 41 μM using a partial viral polyprotein (ΔP1-2A-2B) as substrate. Moreover, when added to HAV-infected cells at a concentration of 5 μM, inhibitor Ia reduced the viral yield by 25-fold, indicative of a much lower IC₅₀ value in vivo. The efficacy of the inhibitor in cells is probably due to a combination of effects, including the irreversibility of inhibition and the fact that polyprotein processing is upstream of many key residues.
replicative events during HAV infection. Therefore, even partially blocking the 3C proteinase function with inhibitor Ia would have a profound impact on progeny virion production.

The inhibition of HAV 3C by inhibitor Ib had been studied initially using a short time course (15 min) and low concentrations of a fluorogenic substrate to minimize the latter’s cleavage and self-quenching. Under these conditions, potent, competitive inhibition was observed with an IC₅₀ value of 13 μM.²⁰ In the current study, a more extensive time course study demonstrated that the initial competitive inhibition (Figure 2(a)) is followed by slow inactivation of HAV 3C by inhibitor Ib in an irreversible and time-dependent fashion (Figure 2(b)). This is in agreement with the structural data presented below that HAV 3C forms essentially the same complexes with both inhibitors Ia and Ib. The IC₅₀ value of inhibitor Ib was estimated to be 15 μM, close to a previously reported value of 13 μM (Figure 2(a)).²⁰ No enzymatic activity was detected after HAV 3C was incubated with inhibitor Ib for 3 h prior to the assay did not lose activity detectably.

**Structural overview**

Structural analysis revealed few differences in the overall protein fold or in the 3C–BBL interactions among the three new complexes and 3C–BBL (Figure 3). In the three new structures, BBL was bound to the Nε² atom of His102 in the same manner as in the 3C–BBL structure.¹⁵ The average r.m.s.d. values between the three new structures reported here and 3C–BBL are in the range of 0.23 Å to 0.30 Å over the Cα positions of all residues (Table 1), suggesting that little structural rearrangement is required for the binding of substrate-analogue inhibitors. This confirms our previous claim that the crystallized enzyme is in its catalytically
Comparison of the r.m.s. d. values at each of the Cα positions reveals that the three new complexes differ significantly from 3C-BBL in only three regions (Figure 4): those comprising residues 49–51, 142–154 and 194–196. These differences suggest that enzyme–substrate complex formation involves an induced-fit mechanism, whereby subtle structural adjustments optimize enzyme–substrate interactions. Residues 142–154 form a major part of the β-hairpin motif that sits atop the catalytic residues in HAV 3C. Some of the residues in this β-hairpin help form the S sub-sites for substrate binding (vide infra). The complexes with inhibitors Ia and Ib share greater structural similarity with each other than with the inhibitor II complex (Table 1). This supports our hypothesis that inhibitors Ia and Ib react similarly in the active site of 3C despite the different substituents on the
terminal α-carbon of these two inhibitors. Additionally, since both inhibitors Ia and II are fluoromethyl ketones, the differences in their tetrapeptidyl sequence must be the key cause for their structural divergences. This suggests that HAV 3C uses subtle structural adjustments to “measure up” the binding fitness of each inhibitor/substrate. The structural differences between the inhibitor Ia and II complexes are consistent with HAV 3C’s preferences for (1) small residues at the P2 position of the cleavage site and (2) glutamine at the P1 position.

3C proteinase–inhibitor interactions and the specificity pockets of HAV 3C

The peptidyl portions of the three ketone inhibitors form canonical β-sheet interactions with the 3C enzyme via four hydrogen bonds: those between the main-chain N of Val144 and O of P4-Leu, the main-chain O of Val144 and N of P2-Ala (or P2-Phe) on one side, and those between the main-chain O of G194 and N of P3-Ala (or P3-Phe), the main-chain N of G194 and O of P3-Ala (or P3-Phe) on the other (Figure 5 and Table 2). This β-ladder is slightly twisted as shown in the relative angles of the above hydrogen bonds to allow for optimal interactions between the side-chains of the inhibitor and the corresponding substrate binding sites (S sites) of the enzyme. Additionally, the main-chain N atoms of the P1 residues in inhibitors Ia and Ib (P1-Gln) form hydrogen bonds (3.2 Å) to the main-chain O atom of Val192. This interaction is not observed in the inhibitor II complex structure that has a longer corresponding distance of 3.9 Å, which likely results from the non-optimal binding of the P2-Phe residue of inhibitor II at the S2 site (see below).

Our structural observations accord well with previous biochemical data that indicate that the P1 and P4 residues primarily determine the specificity in enzyme–substrate interactions. The S1, S2 and S4 sites of HAV 3C are quite well defined, whereas the S3 site is not discernible, and the side-chain of the P3 residue of each of the inhibitors simply protrudes into the solvent. The S2 site, formed by the side-chain atoms of His44, Phe48, Tyr143, His145 and Leu155 and the main-chain atoms of His44, Tyr143, Val144 and His145, is neither wide nor deep enough to accommodate a large hydrophobic residue such as Phe. Consequently, the P2-Phe residue of inhibitor II does not enter the S2 pocket. With a χ1 angle of 79°, its benzene moiety bends toward solvent at the entrance of S2 pocket, stacking against the imidazole ring of His145 (Table 2). The S4 site is a shallow depression on the surface of HAV 3C clustered with the side-chains of hydrophobic residues, namely, Tyr143, Ile198 and Val200, all forming hydrophobic interactions with the side-chain atoms of P4-Leu of the inhibitors.

Finally, the S1 site is of paramount importance in substrate recognition. It is formed by residues 192–195 as one wall and residues 167–169 as the other. His191 sits at the bottom of the S1 pocket and Leu199 at the back end opposite to Cys172. The imidazole side-chain of His191 is locked in a particular tautomeric conformation by the hydrogen bond between its Nδ1 atom and one of the two buried water molecules that are also hydrogen bonded to the carboxylate of Glu132. His191 is poised to interact with the P1 residue of the incoming peptidyl substrate: the Oε1 atom of a P1 glutamine residue would form an optimal hydrogen bond to the Nε2 atom of His191. Indeed, the Oε1 atoms of P1-Glnmm in either inhibitor Ia and Ib are at a distance of 3.0 Å and 2.9 Å to the Nε2 atom of His191. It is interesting to note that neither carboxyl oxygen atoms of P1-Glu in inhibitor II forms a direct

| Complex | r.m.s.d. (Å) |
|---------|-------------|
| 3C–BBL  | 0.30        |
| 3C–inhibitor Ia | 0.20 (0.55) |
| 3C–inhibitor II | 0.19 (0.52) |

*a r.m.s.d. values (Cα positions) over all protein residues and BBL (the carboxyl carbon was used in lieu of Cα) in Å. **Parentheses indicate r.m.s.d. values for inhibitors alone.
Inhibition of HAV 3C by Substrate Analogue Inhibitors

Figure 5. Interactions between the tetrapeptidyl inhibitors and HAV 3C in the active site. The protein residues are distinguished by cyan carbon bonds, the inhibitors by grey carbon bonds/spheres. The catalytic triad, His44, Asp84 and Cys172 are identified by green carbon bonds. Atoms of the 3C proteinase within van der Waals radius to the inhibitors are represented as spheres. The yellow broken lines mainly show the hydrogen bonds formed between the inhibitors and the 3C enzyme. For the sake of clarity, not all protein residues are labeled and solvent atoms are omitted with the exception of a water-bridged interaction between the $O^{\epsilon1}$ atom of the P1 glutamate residue of inhibitor II and the $N^{\epsilon2}$ atom of His191 in B (inhibitor II complex).
hydrogen bond to His191. Instead, the O$_{\varepsilon 1}$ atom of P1-Glu is hydrogen bonded to the N$_{\varepsilon 2}$ atom of His191 via a bridging water molecule, which is at distances of 2.7 Å and 2.6 Å to the former and latter atoms, respectively. In the crystal structure of 3C–BBL complex, a similar water molecule forms a hydrogen bond to the N$_{\varepsilon 2}$ atom of His191 with a comparable distance of 2.8 Å. Similarly, a solvent molecule also forms a hydrogen bond to the equivalent P1 specificity-determining residues in the FMDV 3C (His181) and the TGEV 3CL (His162) crystal structures.\textsuperscript{22,23} In most serine proteinases, the water molecules interacting with the residues in the S1 specificity pockets are displaced upon binding substrate-analogue inhibitors, probably exiting the S1 site via a network of hydrogen bonding water molecules formed at the “back” end of the S1 specificity pocket.\textsuperscript{24} Similar scenes might have happened in our HAV 3C complexes with the ketone inhibitors Ia and Ib.

It is apparent that the binding of inhibitor II to HAV 3C was not sufficiently optimal to expel the solvent molecule interacting with His191 from the S1 site. Consequently, P1-Glu of inhibitor II does not bind as deeply in the S1 pocket as does the P1-Gln$^\text{mm}$ residues of inhibitors Ia and Ib. This provides strong evidence that the imidazole ring of His191 is neutral in these complexes, as otherwise a direct hydrogen bonded ion-pair interaction between P1-Glu and His191 would favor the opposite scenario. With the side-chain of His191 uncharged, the negatively charged carboxylate side-chain of P1-Glu naturally favors a more solvent-exposed location. Additionally, the non-optimal binding of P2-Phe residue of inhibitor II to the S2 site also contributes to the shallow penetration of P1-Glu in the S1 pocket. Compared to their equivalent atoms in inhibitors Ia and Ib, the C$^\beta$ and C$^\alpha$ atoms of P2-Phe in inhibitor II were pushed outwards solvent by 0.97 Å and 0.73 Å, respectively, to avoid clashes between the benzene

### Table 2

Interactions between the tetrapeptidyl inhibitors and HAV 3C protease (episulfide mode) and B. same

| Position$^a$ | Inhibitor Ia | Inhibitor II | Inhibitor Ib |
|--------------|--------------|--------------|--------------|
| **A. A tabulation of interactions by the components in the inhibitors** |
| P4           | 32/9$^b$     | 34/9         | 31/8         |
| P3           | 16/2         | 28/9         | 17/2         |
| P2           | 15/4         | 36/7         | 15/4         |
| P1           | 35/3         | 30/11        | 35/3         |
| C$^\alpha$   | 8/2          | 7/3          | 8/2          |
| O(P1)$^e$    | 8/1          | 12           | 8/1          |
| H-bonds      |              |              |              |
| P4O:Val144N (2.94)$^e$ | P4O:Val144N (3.00) | P4O:Val144N (2.93) |
| P3N:Gly194O (3.04) | P3N:Gly194O (2.86) | P3N:Gly194O (3.04) |
| P3O:Gly194N (2.94) | P3O:Gly194N (2.80) | P3O:Gly194N (2.89) |
| P2N:Val144O (2.72) | P2N:Val144O (2.98) | P2N:Val144O (2.76) |
| P1N:Val192O (3.21) | Water bridged P1O$^\varepsilon 1$:HOH:His191N$^{22}$ | P1N:Val192O (3.19) |
| P1O$^\varepsilon 1$:His191N$^\varepsilon 2$ (2.97) | O(P1):Gly170N (2.77) | O(P1):Gly170N (2.87) |
| O(P1):Cys172N (2.92) | O(P1):Cys172N (2.69) | O(P1):Cys172N (2.84) |
| **B. Statistics assorted by protein residues involved in the interactions (interactions with solvent omitted)** |
| Met29        | 1            |              | 1            |
| His44        | 5            | 1            | 5            |
| Thr142       | 1            | 1            | 1            |
| Tyr143       | 5            | 4            | 4            |
| Val144       | 14(2)$^f$    | 14(2)        | 14(2)        |
| His145       | 3            | 12           | 3            |
| Lys146       | 0            | 14           | 0            |
| Arg162       | 0            | 1            | 0            |
| Gly167       | 5            | 3            | 5            |
| Leu168       | 3            | 3            | 3            |
| Pro169       | 0            | 2            | 0            |
| Gly170       | 3(1)         | 3(1)         | 3(1)         |
| Met171       | 2            | 3            | 2            |
| Cys172       | 12(1)        | 12(1)        | 12(1)        |
| His191       | 3(1)         | 0            | 4(1)         |
| Val192       | 4(1)         | 2            | 4(1)         |
| Ala193       | 6            | 4            | 5            |
| Gly194       | 20(2)        | 22(2)        | 21(2)        |
| Gly195       | 5            | 3            | 5            |
| Asn196       | 2            | 3            | 2            |
| Ile198       | 3            | 3            | 3            |
| Val200       | 2            | 2            | 2            |

$^a$ Residue positions with respect to scissile bond as defined by Schechter and Berger.\textsuperscript{38}

$^b$ Total number of van der Waals interactions/van der Waals interactions with solvent (less than or equal to 4 Å).

$^c$ The terminal $\alpha$-carbon atom next to the carbonyl carbon (C) of the P1 residue.

$^d$ The O atom of the P1 residue.

$^e$ Parentheses indicate distance in Å.

$^f$ Total number of interactions (number of hydrogen bonds).
The inhibitors form an unusual episulfide linkage with the catalytic cysteine residue

In fitting the initial model of the inhibitor into the experimental electron densities near the catalytic cysteine, both the keto carbonyl carbon (CC) of the P1 residues and the adjacent α-methylene carbon (C') of the inhibitors had to be placed within 2.2 Å of the Sγ atom of Cys172. Thus, an episulfide cation fits the electron density far better than a single Sγ-C' bond to either the CC or the C' atom. Although related thiranium cations bearing only carbon substituents are well-recognized intermediates in chemical syntheses,25,26 including a few especially stable, structurally characterized examples,27 to the best of our knowledge there is only one report of a thiranium (episulfide) cation having an oxanion substituent detected as an intermediate in gas phase studies.28 In the final model of HAV 3C with inhibitor Ia, the distances between the Sγ atom of Cys172 and C' or CC (P1-Gln) are 1.83 Å and 1.82 Å, respectively (Figure 6(a)). The imidazole ring of His44 is coplanar with Sγ and C' and its Nε atom is at distances of 4.1 Å and 3.3 Å from Sγ and C', respectively. The distance between C' and CC is 1.56 Å, corresponding to that of a C-C single bond and indicating that the physical strain is well distributed around the entire three-membered ring. The C'-Sγ-CC angle is ∼51°, which is feasible due to the fact that sulfur has a larger atomic radius than either oxygen or carbon. The latter elements would not be able to form a three-membered ring structure with similar stability, which is perhaps why the epoxy ether intermediate proposed to develop during the inactivation of the serine proteinases by chloromethyl ketone inhibitors has not been observed in previous crystal structures.5,29 The episulfide ring structures in the inhibitor Ia and inhibitor Ib complexes are essentially identical (Table 1 and Figures 5 and 6).

Structural refinement of the 3C–BBL–inhibitor II complex structure using the coordinates of the episulfide ring in the 3C–BBL–inhibitor I complex revealed an additional positive electron density peak near the Sγ atom of Cys172 (Figure 6(c)). Further fitting indicated that this represents an alternative mode of bonding of the inhibitor characterized by a single covalent linkage between Cys172 Sγ and the C' of inhibitor II. Interestingly, this mode of bonding corresponds to that observed in the crystal structure of an inactivated papain in which Cys25 is covalently attached to a chloromethyl ketone compound.12 Furthermore, the relative orientations of the methylketone moiety of the inhibitor, the catalytic cysteine and histidine residues are also shared between these two cysteine proteinases of different folds. Nevertheless, the two enzymes made different structural adjustments in arriving at these conformations. In the case of pain, the imidazole ring of His159, in-plane with the Sγ atom of Cys25 in the apoenzyme, rotated about the Cδ–Cγ bond to become coplanar with the atom equivalent to the N atom of the scissile peptide bond. In the 3C–BBL–inhibitor II complex structure containing an episulfide ring, the Sγ atom of Cys172 is coplanar with the C' atom of the inhibitor and the imidazole ring of His44. In the alternative conformation lacking this ring, the atomic positions of His44 are unchanged, whereas the Sγ–Cδ bond of Cys172 is rotated ∼91° out toward the solvent, causing the Sγ atom to roll out of the common plane containing His44 imidazole ring and C'. The new position of the Sγ atom is 2.8 Å from CC atom of P1-Glu of inhibitor II, which is comparable to the average distance (2.8 Å) between Sγ and Cδ in the eight methionine residues in the HAV 3C molecules.

Possible mechanisms for inactivation of HAV 3C by substituted methyl ketone compounds

A comparison of the three structures of HAV 3C–BBL bound with tetrapeptidyl ketone inhibitors with the 3C-iodo-Val-Phe (iVF) complex structure (PDB code 1QA7) suggests that the episulfide ring structure may be an intermediate state on the iVF inactivation pathway. In the latter structure, the imidazole ring of His44 is rotated about its Cδ–Cγ bond by roughly 10° and becomes more in-plane with the carbon at the α position of the ketomethylene function of the iVF inhibitor than with the Sγ atom of Cys172. Judging from the two alternate conformations of the 3C–inhibitor II complex, it seems possible that an internal displacement reaction would resolve the episulfide ring and eventually form an alkylated Cys172 in the active site. However, it is unclear how an episulfide cation is produced in the first place. There are at least three potential routes leading to the formation of an episulfide ring (Figure 7). In scheme I, the Sγ atom of Cys172 attacks C' directly in a nucleophilic substitution reaction to replace the fluoride. This would then be followed by the attack of a lone electron pair of the Sγ atom on the carbonyl carbon to form the oxanion and the episulfide cation. In schemes II and III, the Sγ atom of Cys172 first attacks the carbonyl carbon to form a tetrahedral intermediate and the oxanion. In scheme II, a lone electron pair of Sγ then displaces the fluoride. In scheme III, the oxanion displaces the fluoride to form a transient epoxy ether, which then rearranges to the more stable episulfide ring. As X-ray crystallography shows only snapshots of structures of interest, unless one of these proposed intermediate species can be uniformly formed and stably preserved in crystals, other experimental tools will be needed to differentiate among these mechanistic pathways.

Mechanism of hydrolysis of peptide bonds by the 3C enzyme

Although none of the reactive groups in these inhibitors represents a true peptide bond substrate
Figure 6. Electron densities (from $|2| F_o | - | F_c | \alpha_{calc}$ map contoured at 1 sigma) surrounding inhibitor Ia (a), Ib (b) and II (c). (a) and (b) Residues in 3C are distinguished by green carbon bonds/spheres, whereas the carbon atoms in the inhibitors are colored grey. The electron density contours surrounding the S$^\gamma$ and C$^\beta$ atoms are colored magenta, those around the inhibitors are in cyan. Hydrogen bonds between His44 and Asp84, His191 and P1Gln, as well as those involving the oxyanion are shown as yellow broken lines. (c) Electron densities showing the two possible binding modes of inhibitor II. The color scheme is similar to that in Figure 5, with the ringless alternate conformation of the inhibitor distinguished by slate-colored carbon atoms. Hydrogen bonds similar to those in Figure 4 are shown with the exception that the His191–P1Gln direct interaction is now replaced with two hydrogen bonds bridged through a solvent molecule (black sphere).
for the 3C proteinase, the changes in the relative orientations between the \( S^\gamma \) atom of Cys172, the imidazole ring of His44 and the substituted methyl carbonyl group in the inhibitor nevertheless agree with the hypothesized roles of these catalytic residues in hydrolyzing natural peptidyl substrates. His44, with the assistance of Asp84, acts as a general base catalyst to receive a proton from Cys172, thereby producing a relatively strong nucleophilic thiolate ion. This thiolate subsequently attacks the carbonyl-carbon atom of the scissile bond, forming the tetrahedral intermediate. In the next step, His44 donates a proton to the nitrogen atom of the P1' residue making it a better leaving group. This is followed by the cleavage of the peptide bond and the formation of a thiolacyl enzyme. To fulfill its various roles in this mechanism, His44 must adjust its orientation toward Cys172 accordingly. Interestingly, the two predicted conformations of the His44-Cys172 pair correspond well to those observed in the two 3C-BBL-inhibitor II complex structures.

The only other reported structure of a 3C or 3C-like enzyme inactivated by a methyl ketone inhibitor is that of SARS 3CL-proteinase modified by a peptidyl-based epoxide inhibitor have been obtained in two different crystal forms (PDB codes 2A5I and 2A5K). Recently, crystal structures of the SARS 3CL-proteinase modified by a peptidyl-based epoxide inhibitor have been obtained in two different crystal forms (PDB codes 2A5I and 2A5K). Remarkably, the ring-opening reaction between the inhibitor and the 3CL enzyme covalently linked the catalytic sulfur atom to the C2 atom of the inhibitor that is in a similar position as the C' atom of the halomethyl ketone inhibitors. Compared to the apoenzyme structure (PDB code 2A5A), a drastic rotation of \( \sim 95^\circ \) (change in the \( \chi_1 \) angle) of the \( C^\delta--S^\gamma \) bond in the catalytic cysteine residue is reproducibly observed in two different crystal forms (PDB codes 2A5I and 2A5K). This indicates that 3C or 3CL proteases, as suggested for papain, may undergo local but significant conformational changes in the active site as the hydrolysis of substrates progresses. Inhibitors aimed at structurally “trapping” these viral enzymes at one of the intermediate steps may prove extremely effective in neutralizing these essential viral components.

Materials and Methods

Production and assay of HAV 3C proteinase C24S, synthesis and testing of inhibitor Ib

Experimental procedures for enzyme production and assay, as well as the syntheses of inhibitors Ia, Ib and II (Figure 1) have been published\cite[25] and were followed in
the present study. Enzyme inhibition and HMQC NMR experiments were also done according to the procedures described in the same reports.

Preparation and crystallization of the complex of HAV 3C with inhibitors

Crystals of HAV 3C proteinase variant C24S were grown and harvested as described.15 To make the 3C–BBL–ketone inhibitor triple complex, pre-grown 3C–BBL crystals were soaked in solutions containing ∼5 mM of the substituted methyketone inhibitor for at least 1 h before they were cryo-protected and flash-cooled for data collection. To prepare samples for mass spectrometric analysis, HAV 3C crystals before and after soaking were collected and then rinsed in large excesses of mother liquor to remove any residual or peripheral presence of the inhibitor(s).

Data collection and processing

X-ray diffraction data were collected at Beamline 8.3.1 of the Advanced Light Source (ALS) at Berkeley Lawrence National Laboratory. The programs Denzo, Scalepack and the CCP4 suite were used to process and scale the datasets32,33 (Table 3).

Structure determination and refinement

All structures were solved by molecular replacement using program Molrep in the CCP4 program suite32 and the published model of the HAV 3C-BBL complex15 (PDB code 2CXV) as a search probe. The program XFit34 was used for visualization, fine-tuned model building and adjustment. Program Refmac5 was used for the refinement of the structures.35 Crystallographic and refinement statistics are listed in Table 3.

Structure analysis and generation of Figures

The quality of the structures was assessed using the program PROCHECK.36 Intra and intermolecular contacts in various crystal structures were analyzed using the CCP4 suite of program CONTACT.32 Figures were prepared using the program Pymol†.

| PDB code | 2H9H | 2HAL | 2H6M |
|----------|------|------|------|
| 3C variant | C24S | C24S | C24S |
| 3C–inhibitor complex | 3C–BBL–Ia | 3C–BBL–II | 3C–BBL–Ib |
| Space group | P2₁,2₁,2₁ | P2₁,2₁,2₁ | P2₁,2₁,2₁ |
| a (Å) | 44.67 | 44.58 | 44.77 |
| b (Å) | 56.06 | 56.24 | 56.09 |
| c (Å) | 80.97 | 81.05 | 80.91 |
| α (°), β (°), γ (°) | 90, 90, 90 | 90, 90, 90 | 90, 90, 90 |
| No. molecule/asymmetric unit | 1 | 1 | 1 |
| Vm (Matthews’ coefficient)/% solvent content | 2.1/41.7 | 2.1/41.7 | 2.1/41.7 |

Data collection

| Resolution range (Å) | ALS Beamline 8.3.1 |
|----------------------|---------------------|
| ALS Beamline 8.3.1 |
| Resolution range (Å) | 26.5–1.35 (1.40–1.35) |
| Wavelength (Å) | 1.115879 |
| Observations | 98,591 |
| Unique reflections | 36,919 (1013) |
| Data completeness (%) | 81.2 (22.6) |
| Rmergeb | 0.056 (0.307) |
| No. reflections used | 34,342 (1263) |
| Resolution range (Å) | 26.49–1.39 (1.43–1.39) |
| Rworkingc (%) | 18.1 (32.9) |
| Mean B valued | 20.97/32.28/37.22 |
| Bond angle (°) | 1.200 |
| Chirality | 0.081 |
| Ramachandran plot outliers (phi, psi) | Asp36 (51.2, –129.2) |
| Bond length (Å) | 0.009 |
| No. atoms | 1946 |
| No. waters | 263 |
| Rfree (%) | 17.7 (30.0) |

| r.m.s.d. from ideal geometry |
|-----------------------------|
| Bond angle (°) | 1.200 |
| Chirality | 0.081 |
| Ramachandran plot outliers (phi, psi) | Asp36 (51.2, –129.2) |
| Bond length (Å) | 0.009 |
| No. atoms | 1946 |
| No. waters | 263 |
| Rfree (%) | 17.7 (30.0) |

* Parentheses indicate values for the highest resolution shell.

b Rmerge = Σ |Ihj – <Ih>| / Σ |Ih|, where <Ih> is the weighted mean intensity of the symmetry-related reflections Ih.

c Rworking = Σ |Fobs| – |Fc| / |Fobs|, where |Fobs| and |Fc| represent the observed and calculated structure factor amplitudes, respectively. Rfree is Rworking calculated with the reference set.

d Average B factors of the complex/tetrapeptidyl inhibitor/solvent molecules.
Protein Data Bank accession codes

The coordinates and associated structure factors have been deposited into the RCSB Protein Data Bank (PDB codes 2H91, 2H6M and 2HAL for the HAV 3C complexes with inhibitors Ia, Ib and II complexes, respectively).

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2006.06.047

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