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Identification of Essential Residues in Potyvirus Proteinase HC-Pro by Site-Directed Mutagenesis

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Two virus-encoded proteinases are responsible for proteolysis of potyvirus polyproteins. One of these, HC-Pro, is a multifunctional protein that autolytically cleaves at its carboxyl-terminus (J. C. Carrington et al., 1989, EMBO J. 8, 365–370). To identify the class of proteinase to which HC-Pro belongs, tobacco etch virus (TEV) HC-Pro mutants containing single amino acid substitutions at serine, cysteine, aspartic acid, and histidine positions were synthesized by in vitro transcription and translation and were tested for autoproteolytic activity. Combinations of these residues are constituents of the active sites of diverse groups of cellular and viral proteinases. Only those positions that were strictly conserved among four potyvirus HC-Pro proteolytic domains (for which sequences have been deduced) were mutagenized. Of the 19 mutant proteinases synthesized and tested, only those with alterations at Cys-649 and His-722 were defective for HC-Pro autolytic activity. Most of the other mutant proteinases exhibited no impairments in processing experiments. The spectrum of essential residues, as defined by this genetic analysis, supports the hypothesis that HC-Pro most closely resembles members of the cysteine-type family of proteinases.

INTRODUCTION

Tobacco etch virus (TEV), a member of the potyvirus group, possesses a single-stranded RNA genome (10 kb) that encodes a large polyprotein (Allison et al., 1986). All TEV-derived proteins arise by proteolytic cleavage of the polyprotein precursor (Allison et al., 1985, 1986; Dougherty and Carrington, 1989). The activities of two TEV-encoded proteins, HC-Pro and 49-kDa (or Nla) proteinase, are responsible for most or all proteolytic events associated with polyprotein processing (Carrington and Dougherty, 1987a,b; Carrington et al., 1988, 1989a). The TEV 49-kDa proteinase catalyzes cleavage at five positions, all of which reside within the carboxyl-terminal two-thirds of the polyprotein. The proteolytic domain of the 49-kDa proteinase is highly conserved in structure relative to the analogous Nla proteinases from other potyviruses, to the 24-kDa proteinase of cowpea mosaic virus (CpMV), as well as to the 3C proteinase of picornaviruses (Allison et al., 1986; Domier et al., 1987; Krausslich and Wimmer, 1988). Compelling evidence has been presented that shows the Nla, CpMV 24-kDa, and picornaviral 3C proteinases originated from the trypsin superfamily of serine proteinases by an evolutionary pathway that included conversion of the active-site serine to cysteine (Bazan and Fletterick, 1988; Gorbunova et al., 1989b).

The second potyvirus proteinase, HC-Pro, appears to catalyze only one proteolytic reaction, that being an autolytic cleavage at its carboxyl-terminus (Carrington et al., 1989a,b). This protein resides near the amino-terminus of the TEV polyprotein, and cleaves between a Gly-Gly dipeptide located at amino acid positions 763–764 in the TEV polyprotein. Processing of the TEV polyprotein by HC-Pro liberates an 87-kDa precursor that is further cleaved in vivo to products of 31- and 56-kDa (the latter being HC-Pro) (Thornbury et al., 1985; de Mejia et al. 1985). The identity of the proteinase that processes the amino-terminus of HC-Pro has not been identified. In addition to its role as a proteolytic enzyme, HC-Pro functions as a helper component (HC) for virus transmission from plant-to-plant in nature (Thornbury et al., 1985). A limited degree of amino acid sequence similarity has been noted between the amino-terminal region of HC-Pro of potyviruses and the helper component encoded by the cauliflower mosaic virus genome (Domier et al., 1987). Since only the carboxyl-terminal region of HC-Pro is required for proteolytic activity (Carrington et al., 1989a), HC-Pro appears to possess a two-domain structure.

Most proteinases can be grouped into one of four classes according to the chemically reactive groups at their active site: Ser, Cys, Asp, or Zn$^{2+}$ (Neurath, 1984; Barrett, 1986). In most cases, amino acid sequences surrounding the active-site residues, as well as the spatial relationship of active-site component residues to one another, are highly conserved among members of a given class. In theory, therefore, if a proteinase belongs to one of the four classes, its class assignment may be made by generating mutants with substitutions.
of amino acid residues known to function in proteinases of that class. Serine proteinases would require at least one Ser, His, and Asp residue. Typical cysteine proteinases would show a requirement for at least one Cys and one His residue (Polgar and Halasz, 1982). Aspartic acid proteinases would exhibit a requirement for at least two Asp residues (Tang et al., 1973), while the zinc-dependent endoproteinases would require at least two His residues (Kester and Mathews, 1977; Jongeneel et al., 1989). Additionally, the viral cysteine proteinases should exhibit a requirement for at least one Cys, His, and Asp residue (Bazan and Fletterick, 1988).

In this article, we have sought to identify the nature of the proteolytic activity of HC-Pro by selective site-directed mutagenesis. We rationalized that the active-site residues within HC-Pro from several potyviruses would be invariant and spatially conserved. Therefore, the amino acid sequences of the proteolytic domain of HC-Pro encoded by four potyviruses were aligned to identify regions that contained strictly conserved Ser, Cys, His, or Asp residues (Fig. 1). These positions were mutagenized in a TEV cDNA construct, and the modified proteinases were expressed and tested for activity using cell-free transcription and translation systems. On the basis of this genetic analysis, we propose that HC-Pro exhibits greatest similarity to well-established members of the cysteine class of proteolytic enzymes.

MATERIALS AND METHODS

Recombinant plasmids and strains

Complementary DNA representing the HC-Pro coding region (from the highly aphid-transmissible strain of TEV) was inserted into the cell-free expression vectors pTL7SN and pTL7S. These vectors resemble pTL-8 (Carrington and Dougherty, 1987b) except that the M13 intergenic region has been added upstream of the bacteriophage SP6 promoter to facilitate production of single-stranded DNA with the aid of helper phage M13K07. A key feature of these vectors is the presence of TEV 5'-nontranslated region and translational start sequence. DNA sequences lacking their own start codon can be inserted and expressed in vitro using cell-free transcription and translation systems. On the basis of this genetic analysis, we propose that HC-Pro exhibits greatest similarity to well-established members of the cysteine class of proteolytic enzymes.

RESULTS

Conserved amino acid sequences among four potyvirus HC-Pro proteolytic domains

We have employed a genetic approach to identify the class of proteinase to which potyvirus HC-Pro belongs.
By identifying essential Ser, His, Cys, or Asp residues within the proteinase, and assuming that HC-Pro functions by a mechanism described for one of the four major proteinase classes, the catalytic nature of the enzyme may be deduced. Working on the assumption that HC-Pro from several potyviruses will possess a highly conserved three-dimensional structure and catalytic mechanism, we aligned the sequences of the proteolytic domain of HC-Pro of TEV, tobacco vein mosaic virus (TVMV), potato virus Y (PVY), and plum pox virus (PPV) to identify those positions with invariant residues among all four viruses (Fig. 1).

The proteolytic domains of each virus were highly conserved in sequence; nearly half of the positions (75 of 155) were invariant, and no insertions or deletions were detected. The TEV sequence was most similar to that of TVMV (66%), although extensive similarity was evident with PVY (62%) and PPV (60%). Given that the entire HC-Pro sequences of TEV and TVMV display 54% similarity (Domier et al., 1987) the proteolytic domain (carboxyl-terminal) appears to be more conserved than the helper component domain (amino-terminal).

The Gly–Gly cleavage site recognized by TEV HC-Pro in vitro is constant in each viral sequence, allowing the prediction that the carboxyl-terminus of HC-Pro will originate from position 713 in the TVMV polyprotein, 766 in PPV, and 740 in PVY. Of the 29 TEV positions occupied by Ser, His, Cys, or Asp, only 17 were conserved among all four viruses.

**Mutagenesis of conserved amino acid residues**

Oligonucleotides were synthesized to generate amino acid substitutions at the 17 conserved positions containing Ser, His, Cys, or Asp, as well as 2 nonconserved Ser positions (designated S3 and S6; Table 1). Except for His positions, substitutions that maintained the chemical nature of the original amino acid residue were selected, since these would be predicted to disrupt less the overall structure of the protein but would still eliminate proteolytic activity if the substitution were imposed at an active site residue. Thus, Ser residues were converted to Thr in mutants S1 to S7, Cys residues were changed to Ser in mutants C12 and C13, and Asp residues were converted to Glu in mutants D15 to D20. The choice of substituting His residues with Ser in mutants H8 to H11 was made not on the basis of high chemical similarity, but because His to Ser substitutions are most common at positions in HC-Pro where one of the potyviral sequences contains a nonconserved His. We reasoned, therefore, that imposition of Ser at His positions could maintain the general structure of the proteinase, although the weakly basic character of His would be lost. All mutations with the
The exception of S1 were placed in pTL7S-2027, which harbors TEV cDNA representing a genome region encoding a 30-kDa segment of the polyprotein (Fig. 2). This segment encompasses the proteolytic domain (20-kDa) of HC-Pro, the Gly–Gly cleavage site, and a 10-kDa fragment of the protein adjacent to HC-Pro within the polyprotein. Expression of a similar construct in vitro yields products of 20 and 10 kDa, which result from autoproteolytic cleavage of the 30-kDa synthetic precursor polyprotein (Carrington et al., 1989a,b). Mutation S1 was generated in pTL7S-1527, which is similar to pTL7S-2027 except that it encodes a 37-kDa (rather than 20-kDa) segment of the polyprotein (Fig. 2).

Synthetic transcripts were generated from wild-type pTL7S-1527 and pTL7S-2027 and subjected to cell-free translation in the message-dependent, rabbit reticulocyte lysate system. As shown previously, translation resulted in proteolytic products of 37 and 10 kDa using wild-type pTL7S-1527 transcripts and in proteolytic products of 20 and 10 kDa using wild-type pTL7S-2027 transcripts (Fig. 3). Of the 19 mutants tested, only 2 were deficient in proteolytic activity. Alterations of pTL7S-2027 affecting Cys-649 (mutant C12) and His-722 (mutant H10) completely inhibited formation of the 20- and 10-kDa products and promoted accumulation of the intact 30-kDa precursor. Little or no effect on proteolytic activity was detected in the other 17 mutant HC-Pro derivatives tested. Although it appears that mutant D19 was partially defective in proteolytic activity, this result was not reproducible in subsequent experiments (for example, see below).

To further characterize these HC-Pro derivatives, and to identify mutants that exhibited relatively subtle effects on the rate of proteolytic cleavage, reaction progress curves were generated by pulse–chase analysis for each mutant proteinase. Translation products were synthesized during a 7-min period in the presence of [35S]methionine, followed by addition of excess non-labeled methionine to terminate incorporation. Samples were withdrawn at various times and analyzed by SDS–PAGE. As shown previously (Carrington et al., 1989b), proteolysis of the synthetic polyproteins encoded by wild-type pTL7S-1527 and pTL7S-2027 transcripts proceeded rapidly, with greater than 50% conversion of polyprotein precursor to product immediately after the pulse period (Fig. 4). Consistent with results shown in Fig. 3, HC-Pro mutants with substitutions of Cys-649 (C12) and His-722 (H10) exhibited no proteolytic activity over the time-course examined. The processing kinetics measured for all other mutant proteinases were nearly indistinguishable from those of the wild-type enzyme, although a few reacted slightly slower (e.g., mutant C13 at Cys-684) and a few slightly faster (e.g., mutant S5 at Ser-729). The percentage processing determined at most time-points for each mutant, other than C12 and H10, was within 20% of wild-type in all cases.

**DISCUSSION**

We have conducted a genetic analysis of the proteolytic domain of potyvirus proteinase HC-Pro. Sequences encoding Ser, His, Cys, and Asp residues that were conserved within the HC-Pro molecule from four separate potyviruses were mutagenized in vitro and expressed using a cell-free translation and proteolytic processing assay. The results unambiguously show that only 2 of 19 mutant proteinases, containing amino acid substitutions at Cys-649 and His-722, were proteolytically nonfunctional. Kinetic analysis of processing reactions has demonstrated that the targeted amino acid residues in the remaining 17 mutant proteinases were nonessential to the catalytic mechanism, even though they were strictly conserved among the potyviruses TEV, TMV, PVY, and PPV.

Each of the major groups of proteins contains a unique assemblage of amino acid residues that constitute the active site (see Introduction). Given the spectrum of essential residues we have identified among those mutagenized in this study, we can rule out the possibility that HC-Pro belongs to the serine-type or...
FIG. 2. Genetic map of the tobacco etch virus (TEV) genome, structure of plasmids pTL7SN-1527 and pTL7S-2027, and positions of mutated sequences. The 5'- and 3'-nontranslated regions in the genome diagram are indicated by straight lines, whereas the long open reading frame spanning most of the RNA is shown by the rectangular box. The genome-linked protein (VPg) is shown as the shaded square box, and is attached to the 5'-terminus of the viral RNA (Hari, 1981). The vertical lines show the positions of sequences encoding cleavage sites, while the black square indicates the coding sequence for the Gly-Gly dipeptide cleaved by the HC-Pro (Carrington et al., 1989a). The sizes (in kilodaltons) of the processed TEV proteins are shown above the diagram. The proteolytic domain of HC-Pro is shown by the shaded region within the HC-Pro segment. DNA fragments (enlarged diagram) representing segments of the TEV genome were inserted into vectors pTL7SN (to generate pTL7SN-1527) and pTL7S (to generate pTL7S-2027), which carry cDNA representing the 5'-nontranslated region and initial coding sequence (striped box) of the TEV genome, and a bacteriophage SP6 promoter. Mutation S1 was introduced into pTL7SN-1527, whereas the remaining 18 mutations were placed within pTL7S-2027. TEV nucleotide numbers are shown on the scale at the bottom. Abbreviations: HC-Pro, helper component-protease; Cl, cylindrical inclusion protein; VPg, genome-linked protein; Pro, protease; Pol, polymerase; Cap, capsid protein; Poly(A), polyadenylated tail; Ser, serine; Thr, threonine; His, histidine; Cys, cysteine; Asp, aspartic acid; Glu, glutamic acid.

FIG. 3. Expression and proteolytic activities of wild-type and mutagenized TEV HC-Pro derivatives. Synthetic transcripts were generated from wild-type plasmids pTL7SN-1527 (designated 1527) and pTL7S-2027 (designated 2027) and from plasmids carrying point mutations affecting single amino acid substitutions in the HC-Pro proteolytic domain (designated S1, S2, . . . , D19, D20). The exact position mutagenized in each construct is given in Table 1 and Figs. 1 and 2. Transcripts were translated in the rabbit reticulocyte lysate system in the presence of [35S]methionine, and the radiolabeled products were detected by SDS PAGE and autoradiography. Note that mutants H10 (affecting His-722) and C12 (affecting Cys-649) are defective in proteolytic activity and thus accumulate the precursor polyprotein. Sizes (in kilodaltons) of the radiolabeled proteins are given at the left and right sides.
aspartic acid-type proteinase families, since no Ser or Asp residues were required for proteolytic activity of HC-Pro. If the conserved Asp residue identified by Bazan and Fletterick (1988) in the 3C- and 2A-type viral proteinases functions as an active-site component in a manner analogous to Asp-102 in serine proteinases, we can likewise eliminate the possibility that HC-Pro belongs to the "viral cysteine proteinase" group. Since only one His residue (His-722) was necessary for proteolytic activity, it also appears unlikely that HC-Pro belongs to the metalloendoproteinase group. To the contrary, the finding that only Cys and one His residue were required for proteolytic activity fits precisely with the expectation for a cysteine-type proteinase. Indeed, the cysteine proteinase family is the only group not excluded by our data.

If HC-Pro derives from the cysteine-type proteinase family, the enzyme has evolved substantially since divergence from its cellular counterparts. The overall length of the proteolytic domain of HC-Pro is no longer than 155 residues as defined by deletion studies (Carrington et al., 1989a), whereas the cellular cysteine proteinases possess over 200 residues (Kamphius et al., 1985). The distance between the putative active-site Cys-649 and His-722 is 73 residues, while the active-site residues of papain (type member of the cysteine-type proteinases) are separated by 134 positions. However, the relative placements of putative active-site residues along the polypeptide chain are similar. In papain, the reactive Cys residue occupies position 25, while in HC-Pro Cys-649 resides within 30 amino acid residues from the amino-terminal boundary of the proteolytic domain. Comparisons of the sequence environments surrounding the essential Cys and His residues of cellular cysteine-type proteinases and HC-Pro suggest a distant relationship. A strictly conserved Gly residue is located two positions to the amino-side of the catalytic Cys of several cellular cysteine proteinases, as well as Cys-649 of HC-Pro from the four sequenced potyviruses. Additionally, the consensus sequence Lys-Xaa-Xaa-His-Ala-Val-(3 Xaa)-Gly, which centers on the active site His of cellular cysteine proteinases, bears resemblance to the consensus sequence Lys-Xaa-Xaa-His-Val-Val-(3 Xaa)-Gly,
which surrounds the essential His residue of HC-Pro from the four potyviruses. Direct proof that HC-Pro is assigned correctly to the cysteine proteinase family will require high-resolution structural analysis using X-ray crystallography, although protein modeling using a cysteine proteinase of known structure (e.g., papain) as a guide may yield clues as to the validity of this hypothesis.

Although a cysteine-type proteinase comparable to HC-Pro is not encoded by the genomes of picornaviruses or comoviruses, a possible analog has been identified as a product of the coronavirus genome. Translation of the 5′-proximal cistron of coronaviral genomic RNA yields a large polyprotein precursor termed F1–F2 (Boursnell et al., 1987; Brierley et al., 1987). Two proteinase domains within the polyprotein have been identified by computer-assisted sequence similarity searches (Gorbalenya et al., 1989b). Remarkably, one exhibits similarity to the 3C family of viral proteinases, which includes the potyviral Nla (49-kDa proteinase of TEV). The other, located near the amino-terminus of the polyprotein, strongly resembles a cysteine-type proteinase from Streptococcus pneumoniae. The sequence similarity is particularly noteworthy surrounding the putative active-site Cys residue. Compared to potyvirus HC-Pro, the coronavirus avian infectious bronchitis virus (IBV) protein contains the sequence Gly–His–Cys–Tyr, surrounding the putative active-site Cys residue (underlined), which is nearly identical to the sequence Gly–Tyr–Cys–Tyr surrounding the essential Cys-649 of TEV HC-Pro. Interestingly, the relative positions of these proteinase domains within their respective polyproteins are similar. Further comparisons between these proteins will be possible once the biochemical activity and substrate specificity of the IBV putative proteinase are established and its role in polyprotein processing is understood.

It is important to point out that these experiments have identified Cys-649 and His-722 of HC-Pro as residues important for autolytic activity at its carboxyl-terminus. When a polypeptide representing the initial 97-kDa of the TEV polyprotein is synthesized in vitro, proteolysis at the amino-terminus of HC-Pro (to yield products of 31 and 56 kDa) fails to occur, even though HC-Pro-mediated cleavage at its carboxyl-terminus proceeds efficiently to yield products of 87 and 10 kDa (Carrington et al., 1989a). This has led to speculations that a third proteolytic activity is required for complete HC-Pro maturation, or that auxiliary factors are necessary to complement the activity of HC-Pro or the 49-kDa proteinase. It is possible that HC-Pro possesses two distinct proteolytic activities, one of which catalyzes cleavage at the carboxyl-terminus (the focus of this study) while the other processes at the amino-terminus. This might explain the presence of the sequence motif Gly–Asn–Ser–Gly (positions 623–626 in the TEV polyprotein), which is invariant in all potyviral HC-Pro sequences determined to date, and which is very similar to the highly conserved sequence surrounding the active-site Ser residue in cellular serine-type proteinases. However, one still must concomitantly postulate the involvement of a host factor or separate viral protein to affect proteolysis by this putative activity, since cleavage at the amino-terminus of HC-Pro fails to occur in vitro.

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