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The largest replicative protein of coronaviruses is known as p195 in the avian infectious bronchitis virus (IBV) and p210 (p240) in the mouse hepatitis virus. It is autokathatically released from the precursors pp1a and pp1ab by one zinc-finger-containing papain-like protease (PLpro) in IBV and by two paralogous PLpros, PL1pro and PL2pro, in mouse hepatitis virus. The PLpro-containing proteins have been recently implicated in the control of coronavirus subgenomic mRNA synthesis (transcription). By using comparative sequence analysis, we now show that the respective proteins of all sequenced coronaviruses are flanked by two conserved PLpro cleavage sites and share a complex (multi)domain organization with PL1pro being inactivated in IBV. Based upon these predictions, the processing of the human coronavirus 229E p195/p210 N terminus was studied in detail. First, an 87-kDa protein (p87), which is derived from a pp1a/pp1ab region immediately upstream of p195/p210, was identified in human coronavirus 229E-infected cells. Second, in vitro synthesized proteins representing different parts of pp1a were autokathatically processed at the predicted site. Surprisingly, both PL1pro and PL2pro cleaved between p87 and p195/p210. The PL1pro-mediated cleavage was slow and significantly suppressed by a non-proteolytic activity of PL2pro. In contrast, PL2pro, whose proteolytic activity and specificity were established in this study, cleaved the same site efficiently in the presence of the upstream domains. Third, a correlation was observed between the overlapping substrate specificities and the parallel evolution of PL1pro and PL2pro. Collectively, our results imply that the p195/p210 autoprocessing mechanisms may be conserved among coronaviruses to an extent not appreciated previously, with PL2pro playing a major role. A large subset of coronaviruses may employ two proteases to cleave the same site(s) and thus regulate the expression of the viral genome in a unique way.

All positive-stranded RNA viruses infecting vertebrates, but also many other RNA viruses, employ proteolytic processing as the major regulation mechanism in virus genome expression. The virion RNA enters ribosomes and directs the synthesis of one or two multidomain protein precursors (polypeptides) that, in a controlled fashion, are proteolytically processed by viral and, sometimes, cellular proteases to produce intermediate and mature products. This processing proceeds in cis and in trans at interdomain junctions that contain the specific signals recognized by proteases (1–3).

We have been studying the protease-mediated regulation of viral gene expression using human coronavirus strain 229E (HCoV), which belongs to the Coronaviridae family. Based on a similar polygenic genome organization, common transcriptional and (post)-translational strategies, and a conserved array of nonstructural domains, the Coronaviridae have been united with the Arteriviridae in the order Nidovirales (Fig. 1A) (4, 5). With genome sizes of up to 32 kilobases, coronaviruses have the largest genomes among RNA viruses, whereas the related arteriviruses are much smaller (13–16 kilobases). The positive-stranded genomic RNA of nidoviruses contains 5′- and 3′-nontranslated regions as well as 6–12 ORFs that, in some cases, partially overlap each other. One of the most striking features of the nidovirus life cycle is the specific mode of genome transcription, which results in the synthesis of a 3′-coterminal, nested set of 4–8 sg mRNAs. Except for the smallest transcript, these sg mRNAs are structurally polycistronic (Fig. 1A), but generally, only the most 5′-proximal ORF is translated (reviewed in Ref. 6).

The two largest ORFs (ORF1a and ORF1b), which encompass the 5′-proximal two-thirds of the genome, are believed to encode all the protein functions required for nidovirus RNA synthesis (7, 8). The ORF1b-encoded polypeptide, which includes the putative RdRp activity and a recently established RNA helicase activity that is associated with a unique zinc finger structure (9, 10), is only produced if a ribosomal frameshift from ORF1a into ORF1b takes place during translation (11). This translational strategy is expected to yield two extremely large polypeptides, pp1a and pp1ab, of about 450 and 750 kDa, respectively. To date, pp1a and pp1ab have not been detected in vivo, most probably because they are cotranslationally and autocatalytically processed into numerous processing intermediates and mature nonstructural proteins. Both the
The coronavirus pp1a can be divided into an N-terminal region that is processed by one or two accessory papain-like proteases (Fig. 1B) and a C-terminal region that is processed by the previously identified processing products and the corresponding cleavage sites (P1 and P1′ residues indicated). The following abbreviations are used: PL, papain-like protease; PL1, papain-like protease 1; X, domain conserved in coronaviruses, alphaviruses, rubiviruses, and hepatitis E virus (56); PL2, papain-like protease 2; 3CL, 3CL-like protease; RdRp, RNA-dependent RNA polymerase; Z, putative zinc finger; HEL, NTPase/RNA helicase; C, conserved domain specific for nidoviruses (4).

number and the origin of most of these proteins remain to be determined for coronaviruses (reviewed in Ref. 12).

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by PCR from pBS-J1266 plasmid DNA (47) using primers 134 and 135. The upstream primer contained a BamHI restriction site, and the downstream primer contained a translation stop codon followed by a PstI restriction site. The PCR product was digested with BamHI and PstI and ligated with BamHI/PstI-digested pMal-c2 DNA (New England Biolabs, Frankfurt, Germany). The resulting plasmid, pMal-H2, encoded the specified ORF1a amino acids fused to the Escherichia coli maltose-binding protein (MBP). The plasmid was used to transform competent E. coli TB1 cells, and the bacterial fusion protein was expressed and purified as described previously (46, 48). The HCoV-specific polypeptide, which contained 211 amino acids of pp1a/pp1ab and is predicted by six N-terminal vector-derived amino acids, was released from MBP by cleavage with endoprotease Xa (Amersham Pharmacia Biotech) and used to immunize rabbits as described previously (46). The resulting antisera was designated α-H2.

**Metabolic Labeling, Cell Lysis, and Immunoprecipitation—**Infection or mock infection of MRC-5 cells was done essentially as described previously (49). Briefly, 3 × 10⁶ MRC-5 cells were mock-infected or infected with HCoV at a multiplicity of 10 plaque-forming units per cell. Radioactive labeling of newly synthesized proteins was done for 2.5 h, between 7 and 9.5 h postinfection, with 100 μCi of [³⁵S]methionine per ml. Before labeling, the cells were washed twice with methionine-free minimal essential medium supplemented with 2% dialyzed fetal bovine serum. The cells were lysed in 1 ml of lysis buffer (46). One hundred microliters of a cell lysate was mixed with 400 μl of immunoprecipitation buffer (46) and 5 μl of preimmune serum or 5 μl of α-H2 serum. After 60 min at 4 °C, 25 μl of protein A-Sepharose CL-4B (P9424; Sigma) was added to isolate the immune complexes, which were washed and eluted as described previously (46). The immunoprecipitated proteins were analyzed by SDS-polyacrylamide gel electrophoresis in a 10–17% gradient gel and autoradiography.

**Expression of pp1a/pp1ab Amino Acids by in Vitro Transcription—**Previously, the HCoV PL2pro coding sequence has been found to be non-clonable in E. coli (47). We therefore used PCR-based methods to express this region of the HCoV genome. If not otherwise specified, we used a DNA template that had been isolated from a recombinant vaccinia virus, vHCoV-inf-1, carrying a complete cDNA copy of the HCoV genome (30). The nucleotide sequences of all PCR products used in vitro RNA synthesis were determined to exclude any PCR-derived nucleotide misincorporations. The amino acid sequences of the proteins analyzed in this study are summarized in Fig. 2, and the primers used to generate appropriate DNA templates for in vitro RNA synthesis are given in Table I. To produce proteins pp717–1285, pp717–1436, pp717–1910, and pp717–1285_C1054A, nucleotides 2006–3451 and 3454–6022 were amplified from vHCoV-inf-1 genomic DNA using primers 111 and 113 and 107. The two PCR products were digested with BsoI, purified, and ligated together using T4 DNA ligase. The ligated product was then used as a template for second round PCR amplification with outside primers 111 and 107. The resulting 3,614-base pair PCR product was used as a template for RNA synthesis.

Codon and Deletion Mutagenesis—to generate pp717–1285 C1054A, the coding sequence of pp1a/pp1ab amino acids 717–1285 was amplified by PCR using primers 111 and 107. The PCR product was digested with BsoI and ligated into the EcoRI site of plasmid pBST-51. The resulting plasmid, pBST-111–103, was subjected to site-directed mutagenesis by in vitro recombination PCR (48, 52) using primers 112 and 113. The resulting plasmid was designated pBST-111–103 C1054A. Both pBST-111–103 C1054A DNA and the parental pBST-111–103 DNA were linearized with EcoRI and used as templates for RNA synthesis.

To generate pp717–1285 VM, an in vivo recombination PCR was done using primes 188 and 189 and pBST-111–103 DNA as a template. The resulting plasmid, pBST-111–103 VM, was linearized with EcoRI and used as a template for RNA synthesis. pp717–1285_C1054A, nucleotides 2441–3912 were amplified from pBST-111–103 C1054A plasmid DNA and used as a template to amplify nucleotides 2441–3912 using primers 111 and 139. In a separate reaction, nucleotides 3881–6022 were amplified from vHCoV-inf-1 genomic DNA used by primers 165 and 107. The two PCR products were digested with BsoI, purified, and ligated together using T4 DNA ligase. The ligated product was then used as a template for second round PCR amplification with outside primers 111 and 107. The resulting 3,614-base pair PCR product was used as a template for RNA synthesis.

Codon and Deletion Mutagenesis—to generate pp717–1285 C1054A, the coding sequence of pp1a/pp1ab amino acids 717–1285 was amplified by PCR using primers 111 and 107. The PCR product was digested with BsoI and ligated into the EcoRI site of plasmid pBST-51. The resulting plasmid, pBST-111–103, was subjected to site-directed mutagenesis by in vitro recombination PCR (48, 52) using primers 112 and 113. The resulting plasmid was designated pBST-111–103 C1054A. Both pBST-111–103 C1054A DNA and the parental pBST-111–103 DNA were linearized with EcoRI and used as templates for RNA synthesis.

To generate pp717–1285 VM, an in vivo recombination PCR was done using primes 188 and 189 and pBST-111–103 DNA as a template. The resulting plasmid, pBST-111–103 VM, was linearized with EcoRI and used as a template for RNA synthesis. pp717–1285 VM, nucleotides 171–1285 in which each of the three valine residues, Val⁷⁸⁸, Val⁹⁰⁶, and Val⁹⁰⁸, was replaced with methionine (V900M, V906M, and V908M). To generate pp717–1910 C1054A-VM, an in vivo recombination PCR was done using primes 112 and 113 and pBST-111–103 VM plasmid DNA as a template. The resulting plasmid, pBST-111–103 C1054A-VM, served then as a template to amplify nucleotides 2441–3912 using primers 111 and 139. In a separate reaction, nucleotides 3881–6022 were amplified from vHCoV-inf-1 genomic DNA by using primers 165 and 107. The two PCR products were digested with BsoI, purified, and ligated together using T4 DNA ligase. The ligated product was then used as a template for second round PCR amplification with outside primers 111 and 107. The resulting 3,614-base pair PCR product was used as a template for RNA synthesis.

To generate pp717–1910 C1054A-VM, nucleotides 2441–3912 were amplified from pBST-111–103 C1054A plasmid DNA by using primers 111 and 139, and nucleotides 3881–6022 were amplified from vaccinia virus vF10 DNA² by using primers 165 and 107. The recombinant vaccinia virus vF10 contained a cDNA copy of the HCoV ORF1a in which codon 1702, TGG, has been changed to TTG. The two PCR products were digested with BsoI and ligated together with T4 DNA ligase. The ligated product was then used as a template for second PCR amplification with outside primers 111 and 107. The resulting DNA, which was produced from the purified PCR template encoded the pp1a/pp1ab amino acids 717–1910 in which active-site residues of both PL1pro and PL2pro have been deleted. The proteins tested for proteolytic activity are depicted on the left, and the black lines designate these proteins in relation to their positions in pp1a and pp1ab. See “Experimental Procedures” for a description of the generation of each expression construct.

² V. Thiel, unpublished observations.
The RNA derived from the purified PCR template encoded the pp1a/pp1ab amino acids 717–1910 in which the PL1pro catalytic Cys residue has been replaced with Ala (C1054A). To generate pp717–1910_A1054–1061, nucleotides 2006–3451 and 3476–6022 were amplified by PCR from vHCoV-inf-1 genomic DNA in two separate reactions by using the primer pairs 137/214 and 215/39, respectively. The two products were digested with BsaI and ligated together with T4 DNA ligase. The ligated product was then used as a template for second round PCR amplification with primers 111 and 107. The RNA derived from the purified PCR template encoded the pp1a/pp1ab amino acids 717–1061 and 1082–1910, i.e. residues 1054–1061 have been removed from PL1pro.

To generate pp717–1910_C1054–1061, nucleotides 2006–5392 and 5396–6022 were amplified by PCR from vHCoV-inf-1 genomic DNA in two separate reactions by using the primer pairs 137/214 and 215/39, respectively. The two products were digested with BsaI and ligated together with T4 DNA ligase. The ligated product was then used as a template for second round PCR amplification with primers 111 and 107.

The RNA derived from the purified PCR template encoded the pp1a/pp1ab amino acids 717–1910 in which PL1pro catalytic Cys residue has been replaced with Ala (C1701A). To generate pp717–1910_C1701–1708, nucleotides 2006–5393 and 5426–6022 were amplified by PCR from vHCoV-inf-1 genomic DNA in two separate reactions by using the primer pairs 137/214 and 215/39, respectively. The two products were digested with BsaI and ligated together with T4 DNA ligase. The ligated product was then used as a template for second round PCR amplification with primers 111 and 107.

The RNA derived from the purified PCR template encoded the pp1a/pp1ab amino acids 717–1700 and 1709–1910, i.e. residues 1701–1708 have been removed from PL1pro.

**SARS N-terminal Protein Sequence Analysis**—The proteins pp717–1285_VM and pp717–1910_C1054_VM were produced by in vitro transcription in the presence of [35S]methionine as described above. After incubation of the translation reactions for 160 min, the products were separated by electrophoresis in SDS-polyacrylamide gels and transferred electrophoretically to polyvinylidene difluoride (PVDF) membranes (162–0180, Bio-Rad). The areas of the membranes containing the N-terminal cleavage products were identified by autoradiography and isolated. The bound proteins were then subjected to 16 cycles of Edman degradation by use of a pulsed liquid protein sequencer (ABI 476A, Applied Biosystems, Inc., Weiterstadt, Germany). The eluate from each cycle was mixed with scintillation mixture, and the radioactivity was measured.

**RESULTS**

The Coronavirus p195/p210 Proteases Are Flanked by Conserved PLpro Cleavage Sites and Share a Conserved Five-domain Organization.—The N-terminal part of the replicative polyproteins pp1a/pp1ab of coronaviruses is poorly conserved (14, 47). The proteolytic domains responsible for the processing of this region are part of the largest pp1a/pp1ab cleavage product, known as p195 in IBV and p210 in MHV. These proteases are autocatalytically processed by non-identical mechanisms that, in the case of p195, involve a single PLpro activity and, in the case of p210, both the PL1pro and PL2pro activities (15, 16, 20, 22). Because of these differences, it remained unknown whether the N and C termini of p195/p210 are conserved in coronaviruses. Also, although we suspected that the IBV PLpro may be an ortholog to PL2pro (13, 14), its relationship with the pair of PLpros conserved in all other coronaviruses remained unresolved. By using software for generating global alignments (ClustalX) and local alignments (Dialign2 and Macaw), we have produced a coronavirus-wide multiple sequence alignment which included p195/p210 together with flanking sequences (Fig. 3). The conserved features identified by this alignment include two cleavage sites at the N and C termini and five domains in the order Ac, PL1pro, X, PL2pro and Y, where Ac is the N-terminal domain and Y is the C-terminal domain as defined in this study (see below).

Block A is the only conserved sequence block that was identified upstream of PL1pro. This block is part of a newly recognized domain that varies in size among the different coronaviruses (−150–240 amino acids). Because this domain is highly enriched in acidic Asp and Glu residues, it was named Ac (acidic domain). PL1pro was previously identified in all coronaviruses except IBV. Our alignment shows that IBV may in fact encode a deviant form of PL1pro, which, like its counterparts in other coronaviruses, is located between the Ac and X domains and contains a conserved sequence around the catalytic Cys residue (block B). We consider this domain to be enzymatically defective because, in contrast to other coronaviruses, the other conserved sequences essential for proteolytic activity (e.g. catalytic His residue and zinc finger) are missing in the IBV sequence. The assignment of this domain as a PL1pro remnant is consistent with the PL2pro-like features observed for the IBV PLpro. Thus, both the coronavirus PL2pros and the IBV PLpro occupy similar positions in pp1a/
FIG. 3. Multiple sequence alignment of the p195/p210 regions of coronavirus replicase polyproteins. An initial draft of this alignment was generated using the Dialign2 program (35) and subsequently improved with the ClustalX program (34). The alignment was further checked...
pp1ab; they are embedded between nonconserved regions of variable sizes that, on the upstream side, separate the X domain from PL2pro/PLpro and, more downstream, PL2pro/PLpro from the Y domain. Furthermore, immediately upstream of the catalytic Cys residue, both the PL2pros and PLpro (but not the PL1pros), share a moderately conserved region of ~80 amino acids, which includes the particularly conserved block C. These observations suggest that the p195/p210 proteins of all coronaviruses have a uniform domain organization and include two PLpros. In IBV, one of these proteases (PL1pro) is proteolytically defective and the other one (PL2pro) is proteolytically active. (Henceforth, we use PL2pro rather than PLpro to refer to the proteolytically active papain-like protease of IBV).

The largest conserved domain identified in this study encompasses a region of ~450–490 amino acids at the C terminus of all coronavirus p195/p210 proteins. It was named Y domain. This domain contains two highly hydrophobic stretches and 11 conserved Cys/His residues (14), all in the N-terminal ~180 amino acids. These structural features lead us to predict that the Y domain may anchor p195/p210 into membranes and bind Zn$^{2+}$ or similar metal ions. Overall, the multdom domain organization of p195/p210 implies that the protein is multifunctional.

The conservation of the cleavage sites flanking p195/p210 was recognized on the basis of the immediate proximity of these sites to sequence blocks whose identification proved to be statistically rigorous. The cleavage sites previously identified at the p65 \( \rightarrow \) p210 junction in MHV (Ala \( \rightarrow \) Gly) and the p87 \( \rightarrow \) p195 junction in IBV (Gly \( \rightarrow \) Gly) matched one another in a region upstream of the conserved sequence block A (Fig. 3). In HCoV and porcine transmissible gastroenteritis virus (TGEV), previously uncharacterized Gly-Gly dipeptides with similar positions were identified as putative cleavage sites. Likewise, all coronaviruses have putative cleavage sites of similar composition (Gly \( \rightarrow \) Gly, Ala \( \rightarrow \) Ala, Ala \( \rightarrow \) Gly, or Ser \( \rightarrow \) Gly) in the vicinity of the C terminus of block D. Strikingly, after this analysis was performed, one of these predicted sites, Gly \( \rightarrow \) Gly, proved to be cleaved by PL2pro at the p195 \( \rightarrow \) p41 junction in IBV (16). Similarly, the preliminary mapping data obtained recently for the MHV PL2pro p210 \( \rightarrow \) p44 cleavage site (22) are compatible with the location of the predicted Gly \( \rightarrow \) Ala scissile bond.

In this study, we were specifically interested in relating the implications of the sequence analysis to the understanding of the autocatalytic release mechanisms of coronavirus p195/p210. This protein is predicted to be released from the N-terminal region of the predicted protein, with a calculated molecular mass of 87,345. It was shown in Fig. 4, revealed two major proteins that had apparent molecular masses of 87 (p87) and 230 kDa (p230). The proteins were specifically precipitated by immune serum from metabolically labeled lysates from 3\( \times \)10$^6$ mock-infected (M) (lanes 1 and 3) or HCoV-infected (I) (lanes 2 and 4) MRC-5 cells. The proteins were analyzed by SDS-polyacrylamide gel electrophoresis in a 10–17% acrylamide gradient gel after immunoprecipitation with \( \alpha \)-H2 antiserum (lanes 3 and 4) or the corresponding preimmune serum (lanes 1 and 2). Antiserum \( \alpha \)-H2 recognizes the HCoV ORF1a-encoded amino acids 112–322. The cells were labeled from 7 to 9.5 h postinfection with 100 \( \mu \)Ci of [\( ^{35} \)S]methionine per ml. Sizes of molecular mass markers (CFA 626; Amersham Pharmacia Biotech) with masses in kilodaltons as well as the 230- and 87-kDa processing products, p230 and p87, respectively, are indicated.

The borders of the domains are tentative. The alignments of the PL1pro and PL2pro regions were based on results of our previous analysis (32). For two regions that are located between domains X and PL1pro, and PL2pro and Y, respectively, no consistent alignments have been produced. Therefore, only the sizes of these regions are indicated. The pp1a position of the rightmost residue in an alignment row is indicated at the right side. The shading of individual residues in the alignment was done according to a four-level conservation; black background and white letters, grey background and white letters, and black letters, respectively, indicate residues that are conserved in 100, 80, and 60% of the sequences. Groups of conserved amino acids are as follows: IVLM; FYW; KRH; DNQE; ST; AG. According to the Macaw, four blocks, which are labeled with letters from A to D above the alignment and are discussed in the main text, are statistically significant for the entire pp1a searching space: A, \( p = 1.1 \times 10^{-22}; \) B, \( p = 2.1 \times 10^{-20}; \) C, \( p = 4.2 \times 10^{-27}; \) and D, \( p = 3.9 \times 10^{-45}. \) Two hydrophobic regions predicted to be trans-membrane domains (40) are marked with dashed lines and denoted with TM1 and TM2, respectively. Other highlights are as follows: +, catalytic Cys and His residues of PLpros; #, postulated metal-chelating Cys and His residues of the PLpro zinc fingers; @, conserved Cys and His residues of domain Y; ©, cleavage sites of PLpros. MHVA and MHJV, MHV strains A59 and JHM. The National Center for Biotechnology Information sequence ID: IBV, 138147; HCoV, 464694; TGEV, 872319; MHVA, 453423 (nucleotide); MHVJ, 286058 (corrected according to Ref. 57); all sequences are for proteins unless otherwise specified.

![Detection of an ORF1a-encoded 87-kDa cleavage product in HCoV-infected cells.](image-url)
A product was determined by autoradiography. The isolated protein was transferred to PVDF membranes, the position of the C-terminal cleavage site was determined. The predicted sites, Gly111 → Asn112 and Gly927 → Gly928, were either analyzed directly (lanes 1 and 2) or the same sequence with an active-site replacement of the catalytic Cys1054 (C1054A, lanes 3 and 4). The translation reactions were done as described under “Experimental Procedures,” and the reaction products were either analyzed directly (lanes 1 and 2) or after further incubation for 120 min (lanes 2 and 4). The positions of full-length precursor proteins and cleavage products are indicated.

| Time: 40′ | 120′ |
|-----------|------|
| 0′  120′  |
| 120′  120′|

**Fig. 5.** Evidence for a second PL1pro cleavage site in the HCoV pp1a and pp1ab polyproteins. A, in vitro translation reactions of capped RNAs derived from EcoRI-linearized plasmids pBST-111–103 and pBST-111–103 C1054A. The respective RNAs encode the pp1a/1ab amino acids 717–1285 (WT, lanes 1 and 2) or the same sequence with an active-site replacement of the catalytic Cys1054 (C1054A, lanes 3 and 4). The translation reactions were done as described under “Experimental Procedures,” and the reaction products were either analyzed directly (lanes 1 and 2) or after further incubation for 120 min (lanes 2 and 4). The positions of full-length precursor proteins and cleavage products are indicated. B, a protein called pp717–1285_VM was translated in a reticulocyte lysate in the presence of [35S]methionine. Except for three amino acid substitutions (Val717 to Pro1910 or truncated versions of it (Fig. 2). In total, four basic constructs and eight mutated variants were designed. All constructs shared the predicted HCoV pp1a/pp1ab amino acids 695–1285. By using these PCR templates, capped RNAs were generated and translated in reticulocyte lysates to characterize the cleavage at the predicted junction by SDS-polyacrylamide gel electrophoresis and N-terminal protein sequence analysis.

**HCoV PL1pro Can Cleave the Conserved Site at the Predicted p87 ↓ p195/p210 Junction—**First, the involvement of PL1pro in the cleavage of the p87 ↓ p195/p210 junction was addressed. To this end, the proteolytic processing of pp717–1285, a protein that contained PL1pro but lacked the X and PL2pro domains, was characterized. We analyzed the processing of the wild-type protein and a mutant, pp717–1285 and pp717–1285 C1054A, in which the catalytic Cys1054 of PL1pro was replaced with Ala. Previously, this mutation was proved to block the PL1pro-mediated cleavage at the p87/p195/p210 junction (25, 32). The observed stability of the precursors might be due to the inability of PL1pro to process this site or an incorrect assignment of this HCoV site by the computer-assisted analysis (Fig. 3). Alternatively, the PL1pro-mediated cleavage at this site could be inhibited by p87, as was previously observed for another coronavirus, MHV (20). (It should be noted that these aspects have not been discussed in the original studies (25, 32) as they are brought to light by the computer-based analysis described in Fig. 3).

To exclude the potential negative effect of p87 on the p87 ↓ p195/p210 processing in this study, we expressed proteins that contained only small fragments of p87 (~140–180 amino acids) immediately adjacent to the predicted C terminus of this protein. By PCR, we produced four DNAs that contained a T7 RNA polymerase promoter, a Met initiator codon, and different 3′-extensions encoding the pp1a/pp1ab amino acids Val717 to Pro910 or truncated versions of it (Fig. 2). In total, four basic constructs and eight mutated variants were designed. All constructs shared the predicted HCoV pp1a/pp1ab amino acid sequence with an active-site replacement of the catalytic Cys1054. Asn112 and Gly897 → Gly898, cleavage site, preceded by a small domain and followed by the Ac-PL1pro domains. In three of the basic constructs, this minimal sequence was extended to include either the X domain alone or both the X and PL2pro domains. Furthermore, two basic constructs, pp717–1285 and pp717–1910, were subjected to site-directed mutagenesis, for example, to inactivate the PL1pro and PL2pro domains or to allow radiosequence analyses of [35S]methionine-labeled cleavage products (for details on the constructs, see “Experimental Procedures” and Fig. 2). By using these PCR templates, capped RNAs were generated and translated in reticulocyte lysates to characterize the cleavage at the predicted junction by SDS-polyacrylamide gel electrophoresis and N-terminal protein sequence analysis.
The analysis was done by SDS-polyacrylamide gel electrophoresis in a 12% polyacrylamide gradient gel. The positions of full-length precursor, pp717–1285, and the C-terminal pp717–1910 cleavage product are indicated. In vitro proteolytic activity of PL2pro domain and identification of its cleavage site. A, the HCoV ORF1a-encoded amino acids 717–1910 were translated in rabbit reticulocyte lysates in the presence of [35S]methionine. The proteins to be tested for proteolytic activity contained wild-type sequence (lanes 1 and 2) or the same sequence with active-site replacements in PL1pro (C1054A, lanes 3 and 4) and both PL1pro and PL2pro (C1054A and W1702L), respectively. The proteins were translated at 30 °C for 40 min, and after the termination of translation, the reaction products were either analyzed directly (lanes 1, 3, and 5) or after further incubation for 120 min (lanes 2, 4, and 6). The analysis was done by SDS-polyacrylamide gel electrophoresis in a 10–17% polyacrylamide gradient gel. The positions of full-length precursor proteins and cleavage products are indicated. B, a protein called pp717–1910_C1054A-VM was translated in a reticulocyte lysate in the presence of [35S]methionine. Except for a PL1pro-inactivating amino acid replacement (C1054A) and three additional substitutions (V900M, V906M, and V908M), which had been introduced downstream to the predicted cleavage site, this protein contained the HCoV pp1a/pp1ab wild-type sequence from residues 717 to 1910. The translation reaction was incubated for 160 min at 30 °C, and the reaction products were separated on an SDS-10% polyacrylamide gel. After electrophoretic transfer to PVDF membranes, the position of the C-terminal cleavage product was determined by autoradiography. The isolated protein was subjected to 16 cycles of Edman degradation, and the distribution of radiolabeled amino acids was determined by scintillation counting. The amino acid sequence of pp1a and pp1ab from positions 895 to 913 is shown. The amino acids Met996, Met998, and Met986 present in pp717–1910_C1054A-VM are shown in boldface type, and the deduced PL2pro cleavage site is indicated by an arrow.
type parent pp717–1910. Both precursors were cleaved to produce proteins with apparent molecular masses of ~120 kDa (Fig. 6A, lanes 1–4). This result indicated that another (non-PL1pro-mediated) activity may be responsible for cleavage of the pp717–1910_C1054A protein at the p87 ↓ p195/p210 junction. To verify that PL2pro is the protease that mediates this cleavage, the double mutant pp717–1910_C1054A/W1702L was analyzed. In this mutant, both the PL1pro and PL2pro domains were inactivated by active-site replacements: PL1pro by a replacement of the active-site nucleophile Cys1054 with Ala, and PL2pro by a Leu substitution for the highly conserved Trp1702, which is immediately adjacent to the active-site nucleophile Cys1701. The mutated protein proved to be proteolytically inactive (Fig. 6A, lanes 5 and 6), confirming that the observed cleavage is indeed associated with the activity of PL2pro.

Based on the data described above, it was reasonable to believe that PL2pro, like PL1pro, cleaves the (same) Gly897–Gly898 bond. Alternatively, PL1pro and PL2pro might use partly overlapping sites (for example, in the Ala895-Ala-Gly-Gly898 sequence) or the two proteases might cleave separate but adjacent sites in the viral polypeptide. To establish the specificity of PL2pro unequivocally, we determined the newly identified HCoV PL2pro cleavage site by protein sequencing, using the same approach as described above for the determination of the PL1pro cleavage site structure. We produced a [35S]methionine-labeled derivative of pp717–1910_C1054A, in which each of the Val900, Val906, and Val908 residues was replaced with Met (pp717–1910_C1054A-VM). These replacements did not affect the PL2pro-mediated processing pattern of the primary translation product (data not shown). The radiosequence analysis of the C-terminal pp717–1910_C1054A-VM processing product revealed that PL2pro cleaves the pp1a/pp1ab Gly897 ↓ Gly898 peptide bond (Fig. 6B). Hence, our combined data show that the HCoV PL1pro and PL2pro domains cleave the same site in the viral polypeptide in vitro.

PL2pro Dominates Over PL1pro in the Cleavage of the Gly897–Gly898 Peptide Bond—The above findings imply that, by cleavage of the same site, both PL1pro and PL2pro are able to mediate the autoproteolytic release of the protein of which they are part. To gain initial insight into how these activities might be coordinated, the effects of the individual domains in the PL1pro-X-PL2pro constellation on the efficiency of the Gly897 ↓ Gly898 cleavage were analyzed.

We initially investigated how the X domain or the combination of X and PL2pro affect the PL1pro-mediated cleavage at the p87 ↓ p195/p210 junction. The data shown in Fig. 7A indicate that, irrespective of whether or not the X domain was...
The ubiquitous occurrence of PL1pro and PL2pro in all coronaviruses sequenced to date (Fig. 3) indicates that these enzymes most probably originated from the duplication of a papain-like protease in one of the ancestors of the contemporary coronaviruses. PL1pro and PL2pro have subsequently evolved as part of the pp1a/pp1ab polyproteins that automatically determines the branching of their phylogeny (Fig. 8A). However, when the radial tree was reconstructed using a multiple alignment of coronavirus PLpros, its topology markedly deviated from the one determined for other replicative proteins (compare Fig. 8, A and B). Only the orthologous enzymes of the most closely related viruses, TGEV and HCoV, and MHV and bovine coronavirus (BCoV), respectively, were clustered together (Fig. 8B). More deeply rooted branches of PL1pro and PL2pro proteins were interleaved (Fig. 8B) rather than forming two separate divisions, one for PL1pros and another for PL2pros (Fig. 8A). The tree topology of Fig. 8B was inferred using an NJ algorithm (42). It was supported by results of a bootstrap analysis (Fig. 8B) and was also observed in one of the five most parsimonious trees with the best score after an exhaustive search of the entire tree-space using PAUP*4.0.0d55 (44). The four other most parsimonious trees also deviated from the expected tree presented in Fig. 8A (data not shown). These observations indicate that either our initial assumption of the one-time duplication of a PLpro domain in the coronavirus ancestral lineage is not correct or the evolution of the paralogous PLpros was complicated by homoplasy events that fooled the reconstruction of the genuine topology presented in Fig. 8A. Remarkably, in the inferred PLpro tree shown in Fig. 8B, the PLpros cluster according to the coronavirus genetic grouping (53), bringing together the paralogous enzymes of the same virus (Fig. 8B). This topology makes the second evolutionary scenario most probable as it is compatible with a parallel evolution of the paralogous PL1pro and PL2pro under the pressure of the common substrate that these proteases cleave in HCoV and in other coronaviruses. Thus, the phylogenetic analysis of PLpros supports the results of other analyses (see above) and indicates that the substrate pressure had a significant impact on the structure of the coronavirus PLpros.

**DISCUSSION**

The life cycle of many RNA viruses is driven by the concerted action of several proteases. The proteolytic enzymes mediate...
the production of diverse functional subunits and thus couple (and regulate) the replication, expression, and encapsidation of the virus genome in a timely and spatially coordinated manner. To do this, proteases cleave non-overlapping sets of few sites in the virus-encoded polyprotein(s) (1–3). In this study, we have characterized two sequentially positioned and paralogous proteases of a human coronavirus with an unusual structural organization featuring a \( \text{Zn}^{2+} \) finger embedded between the two domains of a papain-like fold (32). We now show that these proteases also possess unique functional properties as they have overlapping substrate specificities. The coordination of these protease activities may require an extent of complexity not observed elsewhere.

**PL2pro Is the Dominant Force and PL1pro Is Tightly Regulated to Release p195/p210 in Coronaviruses**—To gain insight into the autocatalytic release mechanisms of the largest coronavirus replicative protein, p195/p210, from its polyprotein precursor, we updated a coronavirus-wide multiple alignment of this protein. The results we obtained revealed that p195/p210 has a conserved domain organization and is flanked by conserved cleavage sites. Furthermore, IBV (like other coronaviruses) is predicted to have two PLpro domains (rather than only one PLpro as thought before). The previously characterized IBV papain-like protease, known as PLpro, was shown to be an ortholog of the coronavirus PL2pro domains, and an inactivated remnant of PL1pro was identified at a more upstream position in the viral polyprotein. We then sought to connect the revised domain organization of p195/p210 with the available experimental data and found that, strikingly, the processing mechanisms at the N terminus of p195/p210 vary among different coronaviruses. Thus, it emerged that the conserved peptide bond at the N terminus of p195/p210 is cleaved by different proteases in two coronaviruses, by PL1pro in MHV (20) but by PL2pro in IBV (15). Furthermore, this bond was apparently not cleaved by PL1pro in vitro in HCoV (25). We attempted to reconcile the results of our theoretical analysis and the published data and to this end, performed a comprehensive characterization of this cleavage reaction in HCoV. By using *in vitro* translation of synthetic RNAs in reticulocyte lysates, it was established that both PL1pro and PL2pro cleave the predicted Gly\(^{897} \downarrow \text{Gly}^{898}\) bond at the N terminus of p195/p210. Although p195/p210 itself remains to be identified in HCoV-infected cells, its upstream neighbor in the polyprotein, p87, which is released by cleavage of the same bond, was detected in this study. This result strongly suggests that our observations do not reflect an *in vitro* artifact.

We then characterized mutants of PL1pro and PL2pro and found specific conditions under which the proteolytic activities of PL1pro and PL2pro, respectively, were evident. In a precursor containing PL1pro but lacking PL2pro, the p195/p210 N-terminal cleavage was mediated by PL1pro. If the HCoV PL1pro was expressed in combination with PL2pro from the same RNA template, the N-terminal cleavage of p195/p210 was significantly stimulated (Fig. 7A). Both results are consistent with similar phenomena reported recently for MHV translcleavage assays (50) (see below). Subsequently, a more detailed analysis of the enhancement of the HCoV PL2pro activity led us to conclude that PL2pro is able to cleave the p195/p210 site on its own. Moreover, it became evident that PL2pro is capable of silencing the PL1pro activity. These conclusions were not reached in a similar study of MHV (50), which failed to positively identify the proteolytic activity of PL2pro. We believe that the apparent discrepancy between the HCoV and MHV data results from technical reasons and does not reflect virus-specific differences in the p195/p210 processing mechanism.

Thus, in the MHV study, the proteolytic activities of proteins containing both the PL1pro and PL2pro domains were demonstrated in respect to the equivalents of the p9\( \downarrow \text{p87} \) and p87\( \downarrow \text{p195/p210} \) sites in bimolecular reactions. Furthermore, the design of these previously tested proteases differed from that of the proteins characterized in our study in monomolecular reactions. Teng et al. (50) observed a significant stimulation of the cleavages in the presence of PL2pro, and in a separate experiment (Fig. 7A in (50)), the cleavages were blocked in a nonconservative (His-to-Pro) PL1pro active-site mutant. Although the latter result was interpreted (50) to argue against an involvement of PL2pro in the cleavage, we consider this conclusion premature since no data on the corresponding PL2pro active-site mutant(s) were presented in that paper. Likewise, in two MHV mutants in which PL1pro was deleted, the C terminus of the deletion was placed downstream of Block C, that is within the predicted N-terminal region of PL2pro (see Fig. 3). As a result, PL2pro was unintentionally truncated, which, according to our model of p195/p210, predetermined the processing-negative phenotype of those mutants. The ability of PL1pro and PL2pro to cleave the p87\( \downarrow \text{p195/p210} \) junction in trans has yet to be characterized for HCoV.

The results discussed above and the other published data (20, 25, 32) suggest that the PL1pro activity at the N terminus of p195/p210 is tightly down-regulated by upstream and downstream domains in both MHV and HCoV. Regardless of the mechanisms of these effects, which remain to be elucidated, these observations indicate that PL1pro may have a very short time frame to exert its proteolytic activity in cis. We therefore suggest that PL2pro releases the N terminus of the p195/p210 proteins in HCoV and other coronaviruses and dominates over PL1pro in this cleavage reaction (Fig. 9), although we acknowledge that the PL2pro activity at this site remains to be formally proved for MHV (and some other coronaviruses). In contrast, the processing at the p195/p210 C terminus may be mediated by PL2pro alone (Fig. 9). This site was shown to be effectively processed by PL2pro in IBV (16) and MHV (22), although the ability of PL1pro to cleave this site was not yet rigorously tested for any coronavirus. Future studies on the p195/p210 C-terminal cleavage site structure should also resolve a slight uncertainty of our computer-assisted prediction, which was due to the low complexity and weak conservation in this region, about the precise location of the scissile bond (Fig. 3). This advance would allow us to correlate the structure of the three cleavage sites with the type of the cognate protease(s) in the N-terminal part of the coronavirus pp1a/pp1ab proteins (Fig. 9).

The characterization of the HCoV p87\( \downarrow \text{p195/p210} \) cleavage *in vitro* proved to be a significant technical challenge, since template DNAs containing the “non-clonable” PL2pro coding sequence had to be produced. *In vitro* ligation and PCR approaches (combined with extensive nucleotide sequencing) finally allowed us to analyze the processing of large size precursors containing both HCoV papain-like proteases. The spectrum of constructs (Fig. 2) allowed us to discriminate between the activities of PL1pro and PL2pro. However, more experiments are yet to be done to address a possible involvement of other conserved and non-conserved domains of p195/p210 as well as other replicative proteins in the modulation of the PL1pro and PL2pro activities. Also, since the cellular environment may be involved in the control of specific proteolytic activities, future studies of the p195/p210 autoprocessing in the natural setting using reverse genetics are required to understand the full complexity of these processes. Also, these studies might reveal variations among coronaviruses.

**Evolution of PL1pro and PL2pro and Their Substrates in Coronaviruses**—The PL1pro-inactive/PL2pro-active organi-
Paralogous proteases with different specificities among cellular enzymes; and the entero-/rhinovirus 2A and 3C proteases, employing similar chymotrypsin-like folds and recognizing different sites, illustrate this trend in viruses (reviewed in Refs. 3 and 54). Surprisingly, PL1pro and PL2pro of HCoV (and presumably other coronaviruses) retained overlapping substrate specificities despite a profound divergent evolution elsewhere in the genome. This conservation involves yet-to-be-identified determinants in PL1pro and PL2pro, although it can already be noted that all proteolytically active coronavirus PLpros share a unique zinc finger that was shown to be essential for the proteolytic activity of the HCoV PL1pro (32). The PL1pro and PL2pro alignments (Fig. 3) revealed that only few positions are occupied by lineage-specific amino acid residues, and this unusual pattern could be linked to the selective pressure of a common substrate, driving the parallel evolution of PL1pro and PL2pro. Accordingly, the expected topology of the coronavirus PLpro tree (Fig. 8A) was not readily reconstructed using the conventional algorithms (Fig. 8B).

\[ p195/p210 \text{ Is a Multifunctional Protein with a Unique Regulation of Expression} \]

The cleavage of the same site by two proteases may provide a specific selective advantage since it creates an additional level of regulation in processes that involve (and consume) p195/p210. The unique character of this regulation might be dictated by the exceptional complexity of the domain organization of p195/p210. It is conceivable that the p195/p210 processing may have different kinetics if either PL1pro/PL2pro or PL2pro alone mediates this reaction. Kinetic parameters could affect the localization of specific products in the cell and/or their interactions with other partners. The p195/ p210 product may be anchored in membranes through hydrophobic regions of the C-terminal Y domain or other mechanisms (55) and, furthermore, may be involved in the regulation of transcription through the PLpro-associated zinc finger domains. Finally, the fact that field isolates of BCoV obtained from two different tissues of the same animal were recently shown to selectively accumulate non-synonymous mutations in the p195/p210 protein points to yet another functional aspect of this multidomain protein. Taken together, these data suggest that the above-mentioned or other activities of this multifunctional protein may have profound effects on the host cell or

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4 J. Ziebuhr, V. Thiel, and A. E. Gorbalenya, unpublished data.
even the entire organism. It is thus tempting to speculate that
the sophisticated two-protease regulation at the N terminus of
p195/p210 might be involved in specific coronavirus-host
interactions.

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