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Papain-Like Protease 2 (PLP2) from Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV): Expression, Purification, Characterization, and Inhibition[†]

Yu-San Han,[‡] Gu-Gang Chang,[§] Chiun-Gung Juo,^{||} Hong-Jen Lee,[‡] Shiou-Hwei Yeh,[⊥] John Tsu-An Hsu,[‡] and Xin Chen^{*,‡}

Division of Biotechnology and Pharmaceutical Research, National Health Research Institutes, Miaoli County, Taiwan 350, Republic of China, Faculty of Life Science, National Yang-Ming University, Taipei, Taiwan 112, Republic of China, Institute of Biomedical Sciences, Academia Sinica, Taipei, Taiwan 115, Republic of China, Division of Molecular and Genomic Medicine, National Health Research Institutes, Miaoli County, Taiwan 350, Republic of China

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ABSTRACT: Viral proteases are essential for pathogenesis and virulence of severe acute respiratory syndrome coronavirus (SARS-CoV). Little information is available on SARS-CoV papain-like protease 2 (PLP2), and development of inhibitors against PLP2 is attractive for antiviral therapy. Here, we report the characterization of SARS-CoV PLP2 (from residues 1414 to 1858) purified from baculovirus-infected insect cells. We demonstrate that SARS-CoV PLP2 by itself differentially cleaves between the amino acids Gly180 and Ala181, Gly818 and Ala819, and Gly2740 and Lys2741 of the viral polypeptide pp1a, as determined by reversed-phase high-performance liquid chromatography analysis coupled with mass spectrometry. This protease is especially selective for the P1, P4, and P6 sites of the substrate. The study demonstrates, for the first time among coronaviral PLPs, that the reaction mechanism of SARS-CoV PLP2 is characteristic of papain and compatible with the involvement of the catalytic dyad (Cys)-S[−]/(His)-Im⁺H ion pair. With a fluorogenic inhibitor-screening platform, we show that zinc ion and its conjugates potently inhibit the enzymatic activity of SARS-CoV PLP2. In addition, we provided evidence for evolutionary reclassification of SARS-CoV. The results provide important insights into the biochemical properties of the coronaviral PLP family and a promising therapeutic way to fight SARS-CoV.

Severe acute respiratory syndrome (SARS)¹ is a life-threatening atypical pneumonia caused by SARS coronavirus (SARS-CoV) (1, 2). SARS-CoV belongs to the coronaviruses (order *Nidovirales*, family *Coronaviridae*, genus *Coronavirus*), which are enveloped positive-stranded RNA viruses (3–5). Its RNA is 29.7 kb in length and encodes structural proteins and two large nonstructural polypeptides, pp1a (486

kDa) and pp1ab (790 kDa) (3, 4). These two nonstructural polypeptides undergo co-translational processing that results in various nonstructural proteins (nsp's) (Figure 1). It is generally believed that the processing of pp1a and pp1ab is performed by two viral-encoded proteases, 3C-like protease (3CL) and papain-like protease 2 (PLP2), encoded on pp1a (Figure 1) (6–8). SARS-CoV and infectious bronchitis virus (IBV) contain only one PLP domain, whereas both murine hepatitis virus (MHV) and human coronavirus 229E (HCoV-229E) have two paralogous PLP domains, PLP1 and PLP2 (9–18). The classification of the PLP1 or PLP2 domain is based on its location in the nsp. The following domains are usually organized in the sequential order: acidic (Ac), PLP1, X, PLP2, and Y domains (6, 14). Because the PLP domains of both SARS-CoV and IBV are located between the X and Y domains, they are considered orthologues of PLP2 (Figure 1) (6, 14). These viral proteases are essential for viral pathogenesis and virulence. Therefore, one possible approach to containing and resolving the global threat of a SARS epidemic is the development of efficacious anti-SARS drugs targeting these proteases. However, little information is available on the enzymatic properties and the substrate specificity of SARS-CoV PLP2. Nor do we have information on the differences between coronaviral PLP1 and PLP2 for achieving inhibition specificity.

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* To whom correspondence should be addressed: Division of Biotechnology and Pharmaceutical Research, National Health Research Institutes, Miaoli County, Taiwan 350, Republic of China. Telephone: 88637 246166 ext. 35718. Fax: 88637 586456. E-mail: xchen@nhri.org.tw.

[‡] Division of Biotechnology and Pharmaceutical Research, National Health Research Institutes.

[§] National Yang-Ming University.

^{||} Academia Sinica.

[⊥] Division of Molecular and Genomic Medicine, National Health Research Institutes.

¹ Abbreviations: SARS, severe acute respiratory syndrome; PLP2, papain-like proteinase 2; 3CL, 3C-like protease; SARS-CoV, severe acute respiratory syndrome coronavirus; IBV, infectious bronchitis virus; MHV, murine hepatitis virus; HCoV-229E, human coronavirus 229E; FMDV, foot-mouth disease virus; BCoV, bovine coronavirus; nsp, nonstructural protein; RP-HPLC, reverse-phase high-performance liquid chromatography; MALDI-TOF MS, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry; Abz, *o*-aminobenzoyl; EDDNP, *N*-ethylenediamine-2,4-dinitrophenylamide; FRET, fluorescence resonance energy transfer.

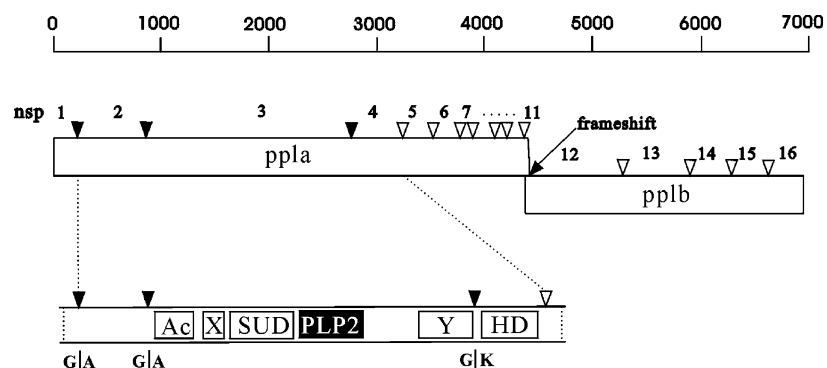


FIGURE 1: Predicted cleavage sites of SARS-CoV PLP2. The amino acid numbering corresponding to the SARS-CoV nonstructural polypeptide (nsp) is marked in the top panel. pp1a and pp1b are the two nonstructural polypeptides before they are cleaved and processed by viral proteases. In the middle panel, the filled triangles mark the sites predicted to be cleaved by SARS-CoV PLP2, and the open triangles mark the sites predicted to be cleaved by SARS-CoV 3CL protease. Numbers 1–16 denote the nsp generated after proteolytic cleavage by SARS-CoV PLP2 or 3CL. The lower panel shows the domain organization of nsp's containing the SARS-CoV PLP2 domain. On this nsp's, Ac, X, SUD, PLP2, Y, and HD stand for the acidic, X, SARS-unique, papain-like protease 2, Y, and hydrophobic domains, respectively (6). The cleavage sites at Gly180–Ala181 (G/A), Gly818–Ala819 (G/A), and Gly2740–Lys2741 (G/K) are marked.

On the basis of sequence comparisons, all coronaviral PLPs, including SARS-CoV PLP2, are thought to be papain-like cysteine proteases with a catalytic dyad composed of Cys and His residues (Figure 2). The sequence homology of the coronaviral PLP domains is less than 25%, and the homology is even lower when compared with papain (14). Therefore, it is not known whether these coronaviral PLPs are similar to cellular papain structurally or catalytically. Mutation at either one of the dyad residues renders the PLPs inactive, as confirmed by its failure to cleave the predicted substrate *in vivo* in cotransfection studies (7, 12, 14, 17, 19–21). Cys1651 and His1812 are predicted to be the catalytic dyad residues of SARS-CoV PLP2 (7, 20). Mutation to Ala at either site abolishes the activity of SARS-CoV PLP2 (7, 20). In these previous studies, the enzymatic activities associated with coronaviral PLPs, including SARS-CoV PLP2, were demonstrated by the *in vivo* cotransfection of two plasmids encoding the PLP and substrate polypeptides or by the *in vitro* transcription and translation of substrate and PLP, followed by Western blot analysis or isotope labeling of the translated polypeptides to detect the cleavage products (7, 9–20, 22, 23). Even though a couple of coronaviral PLPs have been purified to homogeneity from *Escherichia coli* cells as either MBP fusion or by itself, their activities were determined on the substrate synthesized and mixed in the reticulocyte lysate (12, 23). Whether PLP protease cleaves its substrate unassisted or whether its enzymatic activity requires other cellular components *in vivo* remains unclear.

Sequence alignment suggests that SARS-CoV PLP2 cleaves the first three sites on pp1a between Gly180 and Ala181, Gly818 and Ala819, and Gly2740 and Lys2741 of SARS-CoV (Figure 1) (6–8, 20). An *in vitro* transcription and translation experiment showed that cleavage most likely occurs at the Gly818–Ala819 sites (7). Furthermore, cleavage products of the predicted sizes were observed in SARS-CoV-infected Vero-6 cells and in cotransfection study (20). Although these two recent studies support the proposition that SARS-CoV PLP2 is the enzyme responsible for the cleavages, neither the exact cleavage sites nor the substrate specificity of SARS-CoV PLP2 were investigated. Furthermore, limited information is available on the catalytic properties of coronaviral PLP proteases, possibly because

of difficulties in expressing and purifying sufficient amounts of the active protein and the lack of a sensitive and quantitative enzymatic assay with purified enzyme and substrate *in vitro*. In this study, we purified SARS-CoV PLP2 encoding residues 1414–1858 out of nsp3 of SARS-CoV from baculovirus-infected insect cells. Reversed-phase high-performance liquid chromatography (RP-HPLC) coupled to mass spectrometry was established to determine its enzymatic activity, substrate specificity, and exact cleavage sites. The catalytic properties were investigated with a steady-state kinetic technique using a fluorogenic substrate. Finally, a fluorescence-based inhibitor-screening platform was designed to screen for the inhibitors of SARS-CoV PLP2.

MATERIALS AND METHODS

Materials. Grace medium was from Invitrogen. His-Bind Fractogel was from Novagen. Source 15S beads and Superdex 200 HR prepac column were from Amersham Pharmacia. DIG Glycan Differentiation Kit was from Roche. Protease inhibitors were from Sigma. The oligopeptide substrates used in RP-HPLC were synthesized by Kelowna International Scientific (Taiwan).

Plasmid Construction. cDNA encoding a fragment of SARS-CoV PLP2 (amino acids 1414–1858) was PCR-amplified from the reverse transcription product of the SARS viral genome (SARS-TW1 strain) with the following primers CGGGATCCTGAACTCTCTAAATGAGCCGC and CGGAATTCTTACGACACAGGCTTGATGGTTG. This region includes the PLP2 activity domain and part of SUD (Figure 1). The *Bam*HI and *Eco*RI sites in the primers were used to facilitate cloning. The cDNA fragment was cloned into a modified version of the pBacPAK8/His2 vector (24). The resulting plasmid encodes a 51-kDa fusion protein of SARS-CoV PLP2 with a 6× His tag at its N terminus. The insert was sequenced to verify cloning.

Insect Cell Culture, DNA Transfection, Virus Selection, and Amplification. Sf9 cells were grown at 27 °C in Grace medium supplemented with 10% fetal bovine serum. Transfection of DNA into Sf9 cells and selection and amplification of the recombinant virus were carried out as described previously (24). To express the protein, Sf9 cells were infected at a multiplicity of infection of 3, and the cells were harvested 48 h after infection.

	10	20	30	40	50
PAPAIN	IPEYVDWRQKGAVTPVKNNQSGSCG	SWAFSAVVTIEGIIKIRTGNLNEYSE	QELLDCDR--		
SARS_PLP2	-----KWKFPPQVGGLTSIKWAD	NNC YLSSVLLAL LQ QLEVKNF	---APALQEAYYRARAGD		
BCoV_PLP2	-----KWQVVFNGKYFTFKQAN	NNC FVNVSVCL MLQ SLNLKFK	---IVQWQEAWLEFRSGR		
MHV_PLP2	-----KWPVVVCGNYFAFKQSN	NNC YINVACL MLQ HLSLKFH	---KWQWQEAWNEFRSGK		
IBV_PLP2	-----KWNVQYRDNFLILEWRD	NCW ISSAIVL LQ AAKIRFKG	---FLTEAWAKLLGGD		
HCoV_229E_PLP2	-----YESAVVNGIRVLKTS	NNC WVNAVCI LQ YSKPHFIS	---QGLDAAWNKFVLGD		
HCoV_229E_PLP1	-----EMPFEELNGLKILKQLD	NNC WVNSV MLQ LTGILDG	-----DYAMQFFKMGR		
BCoV_PLP1	-----EPEFVKVLDLYVPKATR	NNC WLRSVLAV MQ LPCQFK	---DKNLQDLWVLYKQY		
MHV_PLP1	-----ETHFKVCGFYSPAERT	NCW LRSTLIV MQ SLPLEFK	---DLEMQKLWLSYKSSY		
		*			
PAPAIN	RSYGCNGGYPWSALQLVAQYG	-IHYRNTYPYEGVQRYCRSREKGPYA	-----		
SARS_PLP2	AANFCALILAYSN--	K TVGELGDVRETMT HLQ HANLES	AKRVLN VV--CKHCGQ KTTTL		
BCoV_PLP2	PARFVSLVLAKGG- FKF	-GDPADSRDFLRVVFSQVDLTGAICDFEIA	-- CK--CGVKQEQR		
MHV_PLP2	PLRFVSLVLAKGS- FKF	-NEPSDSTDFMRVVLREADLSGATCDFEFV	-- CK--CGVKQEQR		
IBV_PLP2	PTDFVAWCYASCT- AKV	-GDFSANWLLANLAEHFDADYTNAFLKKRVSCN	- CGIKSYEL		
HCoV_229E_PLP2	VEIFVAFVYYVARL MK --	-GDKGAEDTLTKLSKYLAEAQVQLEHYS	- SCVECD AKFKNS		
HCoV_229E_PLP1	VAKMIERCYTAE--	QCIRGAMGDVGLCM Y RLKDLHTGFMVMDYK	---- CS--CT--SGRL		
BCoV_PLP1	SQLFVDTLVNKIPANIVVPQGGYVADF	Y WFLTLCDWQCWAYW-K	---- CIKCD--LALKL		
MHV_PLP1	NKEFVDKLVKSVPKSIILPQGGYVADF	Y FFLSQCSFKAYANW-R	---- CLKCD--MDLKL		
PAPAIN	-----	AKTDGVRQVQPYNEGALLYSIANQ	-PVSVV		
SARS_PLP2	TGVEAVMY-MGTLSYDNLTGVSI	P -VCGRDATQYLVQQESSFVMSAPP	---AEYKLQ		
BCoV_PLP2	TGVDAMMH-FGTLSREDLEIGYTV	D -SCG-KKLIHCVRFDPFLICSNT	---ASVKLP		
MHV_PLP2	KGVDAVMH-FGTLDKGDLAKGYT	IAC--TCG--NKL VHCTQLNVPFLICS	NKP---EGKKLP		
IBV_PLP2	RGLEACIQPVRATNLLHFKTQYSN	- CPTCG ANNTDEVIEASLPYLLL	FATDG-PATVDCD		
HCoV_229E_PLP2	VASINSAT-VCASVKRDGVQVGY	-- C--VHG IKYYSRVRSVRGRAIIVS	VEQLEPCAQSRL		
HCoV_229E_PLP1	EESGAVLF--CTPTKKAFFPYGT	--- CLNC NAPRMCTIRQLQGTII	IFVQKQPEFVNPVSVFV		
BCoV_PLP1	KGLDAMFF-----YGD-V--	VSHV C--KCG--ES MVLIDVDVPFTA	HFALKDKLFCAFITK		
MHV_PLP1	QGLDAMFF-----YGD-V--	VSHV C--KCG--TG MTLLSADIPYTLHFGLR	DDKFCAFYTP		
PAPAIN	LEAAGKDFQLYRGGIFVGP	CGNKV-- DH AVAAVGYPNYILIKNS	-----WGTGWGENGYIR		
SARS_PLP2	QGTFLCA-----NEYTGN	YQ CGH YTHITAKETLYRI--	DGA HLTKMSEYKGPVTDVIFYKE		
BCoV_PLP2	KG-VGSA-----NIFKG	-DKV GHY VHVKCEQSYQLY--	DAS NVKKVTDVTGNLSDCLYLK		
MHV_PLP2	DD-V-VA-----ANI	FTG-GSL GHY THVKCKPKYQLY--	DAC NVSKVSEAKGNFTDCLYLK		
IBV_PLP2	EDAVGTV-----VFVG	STNS GH CYTQAAGQAFDNLAK	DRK FG-KKSPYITAMYTR----		
HCoV_229E_PLP2	LSGVAYT-----AFSG	VPDK GHY -TVYDTAKKSMY--	DGD RFVKHDLSSLVSVVMVG		
HCoV_229E_PLP1	VKPVCS-----IFRG	AVSC GHY QTNIYSQNL	CV-- DGF GVNKIQPWTNDALNTICIK		
BCoV_PLP1	RS-VYKA-----ACV	VDVND SH MAVVDGKQI----	DDH RITSITSDKFDIFIIGHGMS		
MHV_PLP1	RK-VFRA-----ACV	VDVND SH MAVVDGKQI----	DGK VVTKFNGDKYDFMVGHGMA		
		*			

FIGURE 2: Alignment of the papain-like domains (PLPs) from papain and coronaviral PLPs. The catalytic dyad residues are marked with an asterisk underneath. Dyad and other conserved residues are marked in bold. The amino acid sequences are from the following Genbank accession numbers: P00784 (papain, amino acids 1–188), AY291451 (SARS-CoV-TW1, amino acids 1632–1847 for PLP2), Q9PYA3 (MHV, amino acids 1050–1251 for PLP1 and 1644–1855 for PLP2), Q66198 (BCoV, amino acids 1055–1257 for PLP1 and 1652–1863 for PLP2), Q05002 (HCoV-229E, amino acids 1035–1239 for PLP1 and 1683–1897 for PLP2), and P27920 (IBV, amino acids 1255–1469), respectively.

Purification and Characterization of SARS-CoV PLP2 Protease. The insect cell pellets were resuspended and

sonicated in equilibration buffer containing 20 mM Na₂HPO₄-NaH₂PO₄, 0.5 M NaCl, 20 mM imidazole, and 7 mM

β -mercaptoethanol at pH 8.0. The cell lysate was cleared by centrifugation at 15000g for 15 min and then loaded onto a His-Bind Fractogel affinity column. The column was washed with equilibration buffer containing 20 mM imidazole then with equilibration buffer containing 90 mM imidazole, before elution with equilibration buffer containing 250 mM imidazole. The buffer of the eluate was changed to buffer A containing 20 mM Na_2HPO_4 – NaH_2PO_4 and 7 mM β -mercaptoethanol at pH 6.2, using an Amicon Ultra-15 centrifugal filter tube and then loaded onto a Source 15S column. The Source 15S column was washed with buffer A and then eluted with a gradient of 0–1 M NaCl in buffer B containing 20 mM Na_2HPO_4 – NaH_2PO_4 and 7 mM β -mercaptoethanol at pH 6.2. Western blot analysis was carried out as described with anti-His antibody from Serotec (U.K.) (25). Gel-filtration chromatography was performed with a Superdex 200 HR column as described previously (26). The stability of PLP2 at different temperatures was studied by incubating the enzyme for 30 min at various temperatures before determining its initial rate at 37 °C. The enzyme and substrate concentrations used in the reaction were 0.1 and 20 μM , respectively.

RP-HPLC, Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry (MALDI–TOF MS), and Liquid Chromatography Tandem Mass Spectrometry (LC–MS/MS). The cleavage reaction included the peptide substrate and the purified SARS-CoV PLP2 enzyme at concentrations of 1 mM and 1 μM , respectively, in buffer containing 20 mM Na_2HPO_4 – NaH_2PO_4 and 7 mM β -mercaptoethanol at pH 6.8. The reaction was carried out at 37 °C before the addition of 50 μL of 0.1% trifluoroacetic acid to stop the reaction. The product was analyzed on an Agilent 1100 HPLC system with a Zorbax C18 column (4.6×250 mm), using a 0–80% linear gradient of 90% acetonitrile with 0.1% trifluoroacetic acid. The cleaved products were collected and analyzed with a Voyager DE-STR Biospectrometry Workstation (PerSpective Biosystems, Framingham, MA). α -Cyano-4-hydroxycinnamic acid was used as a matrix. For the cleavage products of Gly180–Ala181 oligopeptide, the fraction was further analyzed on a LC–MS/MS system with a Zorbax column (2.1×250 mm) (LCQ DECA Xpplus, ThermoFinnigan, San Jose, CA), using the same gradient condition in RP-HPLC by replacing trifluoroacetic acid with acetic acid.

Kinetic Constant Measurement. *o*-Aminobenzoic acid (Abz) and *N*-ethylenediamine-2,4-dinitrophenyl amide (EDDNP) were chosen as the fluorescent donor and quencher (27), respectively, to label the N and C termini, respectively, of the peptide substrate FRLKGGAPIKGV, to produce the internally quenched fluorescent substrate Abz-FRLKGGAPIKGV-EDDNP (AnaSpec, San Jose, CA). Enhanced fluorescence emission upon substrate cleavage was monitored at the excitation and emission wavelengths of 320 and 420 nm, respectively (Fluoroskan Ascent, ThermoLabsystems, Sweden). Fluorescence Intensity was converted into amounts of hydrolyzed substrates with a standard curve drawn from the fluorescence measurements of well-defined concentrations of each substrate after complete hydrolysis by SARS-CoV PLP2.

The kinetic constants were measured with this fluorogenic peptide substrate or the substrate without the fluorescent label as described previously (26, 28). The concentrations of the

enzyme and substrate used were 0.1 μM and 0.2–3 times the K_m value (0.125–1 mM), respectively. The buffers used at different pH values were 20 mM acetate buffer at pH 5.25, 20 mM phosphate buffer at pH 6–8, 20 mM Tris–HCl buffer at pH 8.65, or 20 mM 3-cyclohexylamino-1-propanesulfonic acid (CAPS) at pH 9.2–9.8, all containing 7 mM β -mercaptoethanol. The change in ionic strength because of the pH adjustment was negligible with respect to the total ionic strength. Peak area was calculated by integration. The initial rate was measured with less than 10% substrate depletion for the first 20 min to calculate the kinetic parameters using the Michaelis–Menten equation (29, 30). The pH–log k_{cat} or pH–log(k_{cat}/K_m) profiles were fitted to the following equation for two protonation sites (29, 30):

$$\log k = \log \frac{C}{1 + \frac{[\text{H}^+]}{K_{a1} + \frac{K_{a2}}{[\text{H}^+]}}}$$

in which k is the observed kinetic constant (k_{cat} or k_{cat}/K_m) at various pH values, C is the pH-independent limiting k value, and K_{a1} and K_{a2} are the macroscopic dissociation constants for the molecular groups responsible for the two ionization steps. All computer-fitted work was performed with the Sigma Plot 5.0 program (Jandel, San Rafael, CA).

Inhibitor-Screening Platform with the Fluorogenic Substrate. An inhibitor-screening platform was set up in a 96-well plate containing 50 nM SARS-CoV PLP2 and 20 μM fluorogenic substrate Abz-FRLKGGAPIKGV-EDDNP in buffer (20 mM Na_2HPO_4 – NaH_2PO_4 at pH 6.8) without β -mercaptoethanol. The enhanced fluorescence emission upon substrate cleavage was monitored at the excitation and emission wavelengths of 320 and 420 nm, respectively, in the presence or absence of the chemical compounds assayed (Fluoroskan Ascent).

RESULTS

Expression, Purification, and Characterization of SARS-CoV PLP2. Because the minimum PLP activity for other coronaviral PLPs was previously identified within a domain of 230–300 residues (12, 19, 21, 23), our construct encodes the region from amino acid 1414 to 1858 of SARS-CoV pp1a with a calculated molecular weight of 51 kDa. We refer to this region as SARS-CoV PLP2 throughout the text. Whether SARS-CoV PLP2 is glycosylated and whether glycosylation is important for its activity were unknown. Therefore, we chose to express the protein in baculoviral-infected insect cells, which have been used to express glycosylated proteins and other proteins not readily expressible in *E. coli* cells (31). The purified protease ran approximately at the calculated size of 51 kDa on SDS–PAGE (Figure 3A). In gel-filtration chromatography, the PLP2 was eluted at a position corresponding to a 56.7-kDa protein with a Stokes' radius of 3.3 nm (inset of Figure 3B). The result indicates that purified SARS-CoV PLP2 exists as a monomer in solution. Next, we investigated whether the purified SARS-CoV PLP2 was glycosylated because there are several putative N- and O-glycosylation sites (data not shown). From DIG Glycan Differentiation Kit (Roche), we detected none of the following glycosylation forms: mannose, *O*-linked disaccharide

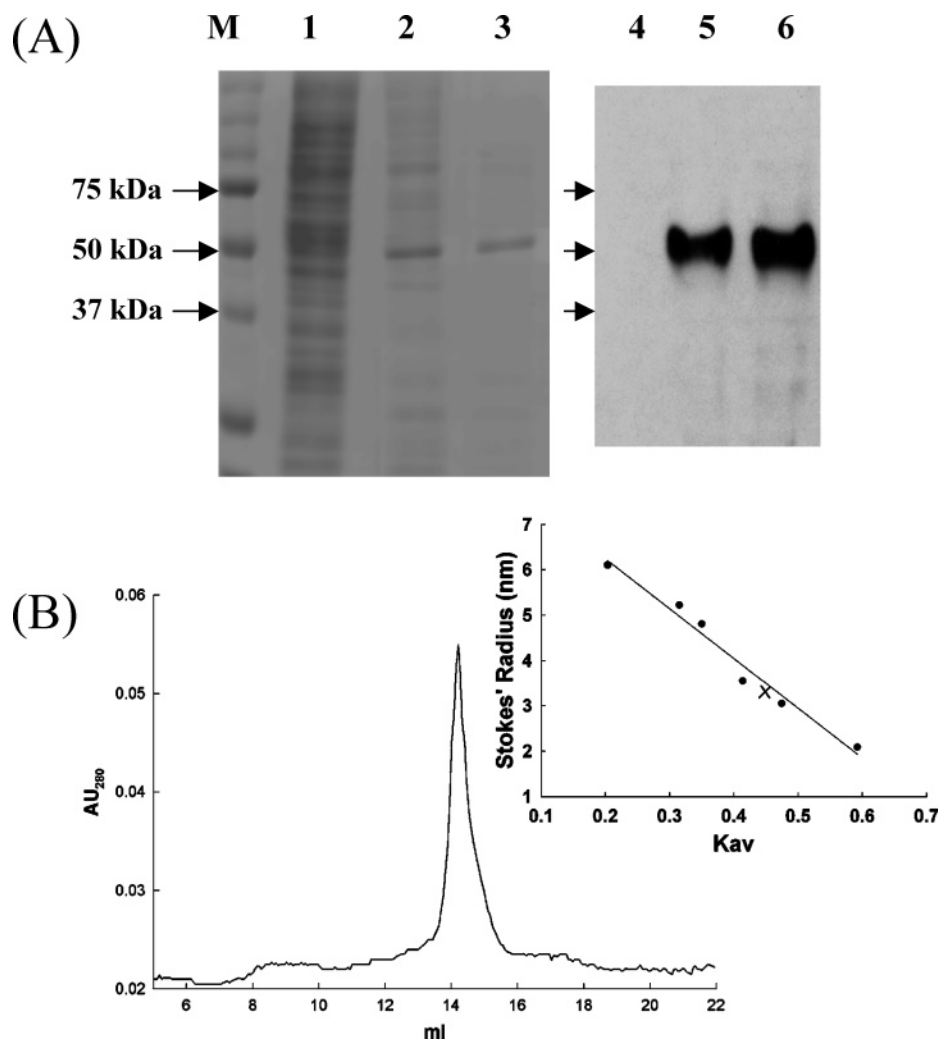


FIGURE 3: Purified SARS-CoV PLP2 is monomeric. (A) Purification of SARS-CoV PLP2. M indicates the molecular weight markers. Lanes 1–3 are a Coomassie-blue-stained gel of insect cell lysate, with the eluate from His-Bond Fractogel and the eluate from a Source 15S column, respectively. The amounts of the proteins loaded in lanes 1–3 are 10, 4, and 2 μ g, respectively. Lanes 4–6 are a Western blot of lanes 1–3, probed with anti-His antibody (see the Materials and Methods). (B) SARS-CoV PLP2 is monomeric by the gel-filtration experiment. The elution profile with a Superdex 200 HR column is shown. The x axis shows the elution position (in milliliters), and the y axis shows the absorbance measurement at 280 nm to detect the presence of the protein. The inset is the calibration curve for the column. The X in the inset marks the elution position for SARS-CoV PLP2.

galactose- β (1–3) *N*-acetylgalactosamine, *N*-linked galactose- β (1–4)-*N*-acetylglucosamine, or *O*-linked *N*-acetylglucosamine (data not shown). Furthermore, the purified PLP2 runs at the expected size on SDS–PAGE (Figure 3A) suggesting that this purified PLP2 is not glycosylated. The activity of the protease was significantly reduced after incubation at temperatures higher than 37 °C for 30 min (data not shown), indicating that SARS-CoV PLP2 is inherently thermolabile. This is consistent with the observation that SARS-CoV cannot tolerate high temperatures.

Intrinsic Cleavage Activity of SARS-CoV PLP2. Using the purified SARS-CoV PLP2, we developed an assay method constituted only of enzyme and substrate and analyzed by RP-HPLC and mass spectrometry. The substrates are 12-mer oligopeptides containing the predicted Gly180–Ala181, Gly818–Ala819, and Gly2740–Lys2741 cleavage sites, RELNGGAVTRYV, FRLKGGAPIKGV, and ISLKG-GKIVSTC, respectively.

As shown in Figure 4A, after 12 h of digestion, only a tiny fraction (5%) of the Gly180–Ala181 substrate was cleaved. Under the same conditions, cleavage at the Gly818–

Ala819 and Gly2740–Lys2741 sites was 100 and 29%, respectively (parts B and C of Figure 4). The percentage cleavage of the Gly818–Ala819 substrate reached 70% after 2 h of digestion (Figure 4D). Under the same condition, only 7% of the Gly2740–Lys2741 sites was cleaved, whereas there was no cleavage detected on the Gly180–Ala181 site (data not shown). All of the peaks in the chromatograms were collected and confirmed by MALDI–TOF MS or LC–MS/MS. The mass results unequivocally demonstrated that cleavage occurred at the expected peptide bonds (data not shown). The 12 h of digestion for the Gly180–Ala181 oligopeptide resulted in a nonspecific cleavage product coeluted with the predicted cleavage product AVTYRV. We only detected the presence of AVTYRV but not RELNGG. These results demonstrated that SARS-CoV PLP2 by itself is active, with a preference for the Gly818–Ala819 site *in vitro*.

Next, we wanted to determine the requirements for substrate length and composition with oligopeptides ranging from a 6-mer to an 18-mer that encompass the Gly818–Ala819 site. The numbering of the P–P' sites is based on

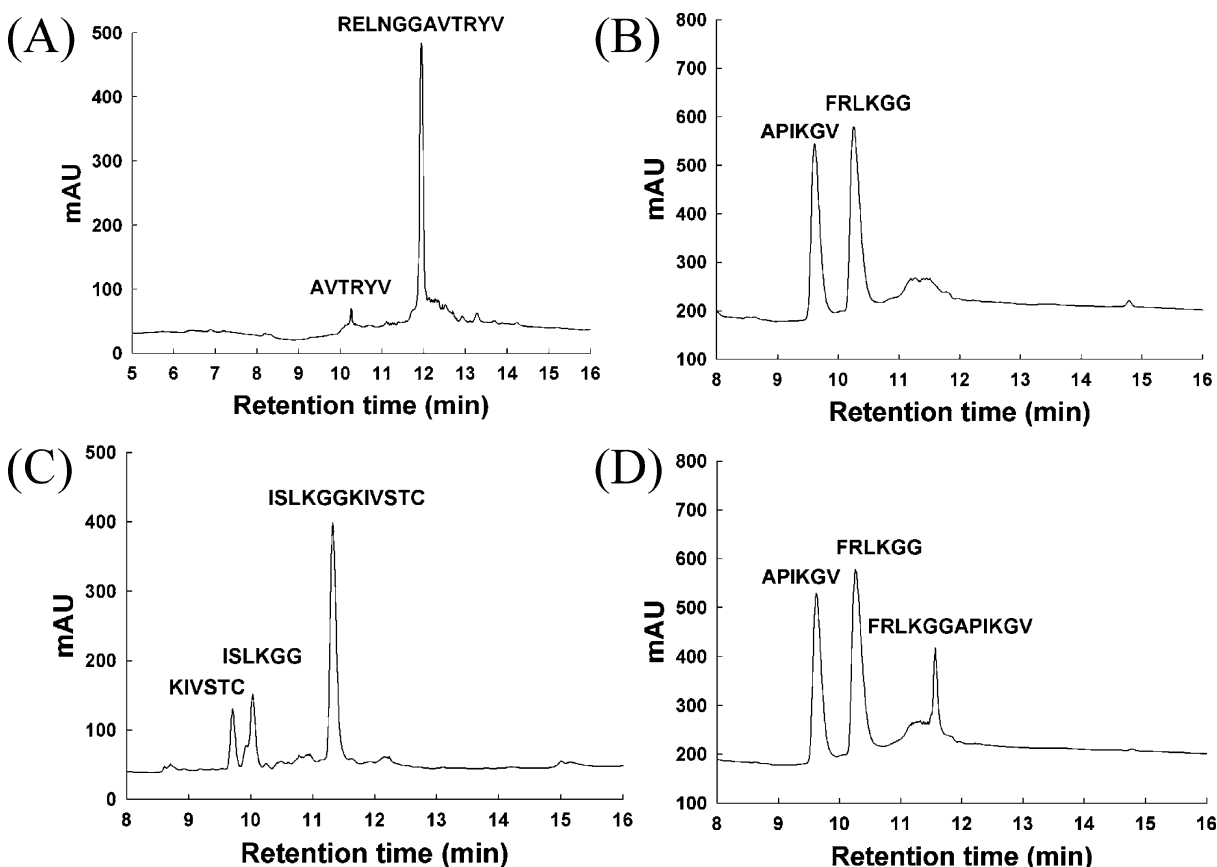


FIGURE 4: RP-HPLC analysis of the sites cleaved by SARS-CoV PLP2. The panels are the HPLC elution profiles of the hydrolyzed products of the oligopeptides RELNGGAVTRYV (A), FRLKGGAPIKGV (B and D), and ISLKGGKIVSTC (C), after incubation with purified SARS-CoV PLP2 for 12 (A–C) or 2 (D) h. The cleavage sites are confirmed by mass spectrometry (data not shown), as indicated on the peaks.

Table 1: Optimal Length of the Substrates for Cleavage by SARS-CoV PLP2

length	peptide substrate																		cleavage % ^a	
	P9	P8	P7	P6	P5	P4	P3	P2	P1	P1'	P2'	P3'	P4'	P5'	P6'	P7'	P8'	P9'	2 h	12 h
18 ^b	N	N	V	F	R	L	K	G	G	A	P	I	K	G	V	T	F	G	60 ± 1	100
16 ^b		N	V	F	R	L	K	G	G	A	P	I	K	G	V	T	F		63 ± 1	100
14			V	F	R	L	K	G	G	A	P	I	K	G	V	T			68 ± 9	100
12				F	R	L	K	G	G	A	P	I	K	G	V				70 ± 7	100
11			V	F	R	L	K	G	G	A	P	I	K						53 ± 4	100
10				F	R	L	K	G	G	A	P	I	K						50 ± 3	100
10					R	L	K	G	G	A	P	I	K	G					15 ± 4	56 ± 6
9				F	R	L	K	G	G	A	P	I							36 ± 6	85 ± 5
8						L	K	G	G	A	P	I	K						11 ± 4	45 ± 4
6							K	G	G	A	P	I							nd ^c	nd

^a Percentages of the cleavage is the average of three independent measurements using three different batches of enzyme. ^b Because of their low solubility, 0.5 mM, instead of 1 mM, of the 18- and 16-mer oligopeptides were used in the assay. ^c Cleavage was not detectable.

Berger and Schechter, with cleavage occurring between the P1 and P1' sites (Table 1) (32). As shown in Table 1, oligopeptides ranging from a 10-mer to an 18-mer, including a 10-mer with six residues in the P positions and four in the P' sites, i.e., FRLKGG-APIK, were excellent substrates for cleavage and were over 50 and 100% cleaved, respectively, after 2 and 12 h of digestion. Interestingly, the 10-mer oligopeptide with five residues at both the P and P' sites, RLKGG-APIKG, was not efficiently cleaved, with only 56% cleavage after 12 h of incubation (Table 1). Moreover, a 9-mer with six residues in the P sites and three residues in the P' sites, FRLKGG-API, was a much better substrate than the 10-mer with five residues in the P sites. This 9-mer was 36 and 85% cleaved after 2 and 12 h of digestion,

respectively (Table 1). These results indicate that six residues in the P sites are critical for efficient recognition and cleavage by the enzyme. Consistent with this, shorter oligopeptides (8- and 6-mer) lacking the P6 residue were poorly cleaved. For the 8-mer peptide, 11 and 45% were cleaved after 2 and 12 h, respectively, while there was no cleavage at all for the 6-mer (Table 1). Therefore, the P6 residue seems to contribute significantly to substrate recognition and cleavage by SARS-CoV PLP2.

Catalytic Properties of SARS-CoV PLP2. The kinetic constants were determined for the optimal substrate Gly818–Ala819, FRLKGGAPIKGV. The apparent k_{cat} and K_m values were 1.5 min⁻¹ and 185 μM, respectively. We then designed a fluorogenic substrate, Abz-FRLKGGAPIKGV-EDDNP,

Table 2: Kinetic Parameters of SARS-CoV PLP2 at Different pH Values^a

pH	k_{cat} (min^{-1})	K_{m} (μM)	$k_{\text{cat}}/K_{\text{m}}$ ($\mu\text{M}^{-1} \text{min}^{-1}$)
5.25	1.4 ± 0.2	21.4 ± 1.1	0.06 ± 0.01
6.05	11.9 ± 1.1	64.0 ± 0.3	0.17 ± 0.03
6.82	20.6 ± 2.4	65.4 ± 5.4	0.31 ± 0.01
7.42	19.8 ± 2.3	77.2 ± 2.1	0.27 ± 0.03
8.01	21.8 ± 1.8	72.0 ± 4.3	0.30 ± 0.02
8.65	22.0 ± 3.1	71.4 ± 8.0	0.30 ± 0.01
9.20	14.0 ± 0.5	68.0 ± 0.6	0.21 ± 0.02
9.80	6.0 ± 0.4	70.4 ± 1.5	0.08 ± 0.01

^a Kinetic constants were measured as described in the Materials and Methods.

which has little fluorescent emission because of the fluorescence resonance energy transfer (FRET) between Abz and EDDNP (27). When this peptide substrate is cleaved between the Gly and Ala by SARS-CoV PLP2, the FRET disappears and fluorescence intensity increases (27). The apparent k_{cat} and K_{m} values for this fluorogenic substrate at pH 7.4 were $20 \pm 2 \text{ min}^{-1}$ and $77 \pm 2 \mu\text{M}$, respectively (Table 2). Using fluorescent substrate is a much more sensitive and convenient method for determination of the enzymatic property, as observed before for other proteases (33). From this method, we determined the k_{cat} and K_{m} values of the enzyme at different pH values ranging from pH 5.25 to 9.8. As shown in Table 2, the enzymatic activity of SARS-CoV PLP2 ($k_{\text{cat}}/K_{\text{m}}$ values) was optimal at pH 6.8–8.6. Lowering the pH to 5.25 affected both the K_{m} and k_{cat} values, with a more significant effect on k_{cat} . The k_{cat} value decreased almost 15-fold, whereas there was only a 3-fold increment in the K_{m} value. At pH 9.8, there was a 3-fold decrease in k_{cat} , with no effect on K_{m} (Table 2). These results indicate that the activity of SARS-CoV PLP2 is greatly modulated by pH, reflecting the stability of the thiolate–imidazolium ion pair at different pH values.

These kinetic data were further analyzed by fitting them to an equation involving the two protonation forms (see the Materials and Methods). Both the catalytic constant (k_{cat}) and the specificity constant ($k_{\text{cat}}/K_{\text{m}}$) of the enzyme displayed bell-shaped activity versus pH dependence, consistent with the existence of two macroscopic molecular ionizable groups associated with maximal catalysis (Figure 5). The apparent pK_{a} values for these ionizable residues were obtained with an excellent fit in both cases. From the pH– $\log k_{\text{cat}}$ plot, a constant C value of $25 \pm 3 \text{ min}^{-1}$, equivalent to the pH-independent k_{cat} value, was obtained. This is the true k_{cat} value, higher than the apparent value ($20 \pm 2 \text{ min}^{-1}$) determined in the regular assay at a fixed pH of 7.4 (Table 2). From the pH– $\log k_{\text{cat}}$ plot, two macroscopic molecular pK_{a} values of 6.4 ± 0.1 and 9.3 ± 0.2 were obtained, most likely attributable to the dyad residues Cys1651 and His1812 of SARS-CoV PLP2. These results indicate that these two residues form the ion pair involved in the catalytic step of enzymatic catalysis and that the enzymatic activity is modulated electrostatically for catalytic competence. These data are consistent with the characteristics of a papain-like cysteine protease (34, 35).

From the pH– $\log(k_{\text{cat}}/K_{\text{m}})$ plot, a constant C value of $0.3 \pm 0.02 \text{ min}^{-1} \mu\text{M}^{-1}$, equivalent to the pH-independent $k_{\text{cat}}/K_{\text{m}}$ value, was obtained. This is the true $k_{\text{cat}}/K_{\text{m}}$ value and is the same as that determined in the regular assay at a fixed

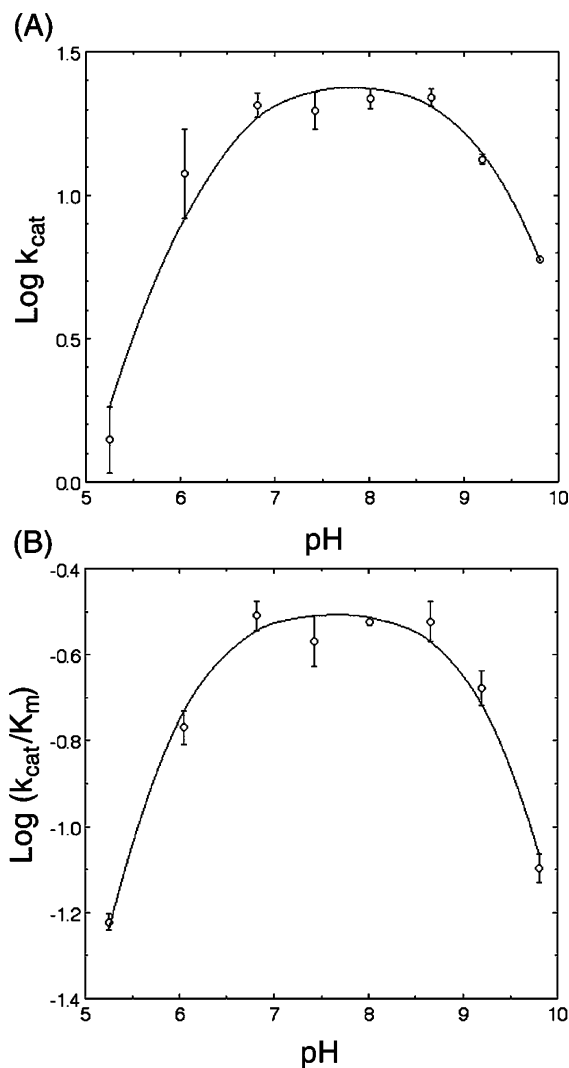


FIGURE 5: Dependence of the kinetic parameters of SARS-CoV PLP2 on pH. (A) Dependence of the catalytic constant (k_{cat}) of SARS-CoV PLP2 on pH. (B) Dependence of the specificity constant ($k_{\text{cat}}/K_{\text{m}}$) of SARS-CoV PLP2 on pH. In both panels, open circles show the data from the kinetic experiments. The line is the computer-generated result of fitting the experimental data shown in Table 2 to an equation for two ionizable groups (see the Materials and Methods).

pH of 7.4 (Table 2). Therefore, the true K_{m} value for the 12-mer peptide substrate should be corrected to $82 \mu\text{M}$. Two apparent pK_{a} values of 5.9 ± 0.1 and 9.4 ± 0.1 were obtained from the pH– $\log(k_{\text{cat}}/K_{\text{m}})$ plot. The former must be deprotonated, and the latter must be protonated to achieve an optimal substrate-binding mode. At this stage, identification of these macroscopic molecular groups, in terms of specific amino acid residues, is not straightforward.

Substrate Specificity of SARS-CoV PLP2. We then investigated the substrate specificity of SARS-CoV PLP2 by focusing on the P6 to P1' sites, because the corresponding residues in other coronaviral PLPs affect the cleavages as shown before (11, 21, 22, 36, 37) and in this study (Table 1). The optimal substrate Gly818–Ala819 was used for comparison. As shown in Table 3, substitution at either the P1 (Gly) or P4 (Leu) position with Ala or Lys (G818A, L815A, or L815K) was not tolerated at all, with no cleavage observed after 12 h of digestion. A significant decrease in cleavage, from 70 to around 20% after 2 h, or from 100 to around 50% after 12 h, was observed when the P2 (Gly) or

Table 3: Substrate Specificity of SARS-CoV PLP2

source	peptide substrate ^a							cleavage % ^b	
	P6	P5	P4	P3	P2	P1	P1'	2 h	12 h
Gly818–Ala819	F	R	L	K	G	G	A	70 ± 7	100
A819G							G	65 ± 10	100
A819N							N	62 ± 12	100
G818A						A		nd ^c	nd
G817A					A			16 ± 3	46 ± 7
G816A				A				71 ± 10	100
G816Q				Q				58 ± 8	100
L815A			A					nd	nd
L815K			K					nd	nd
R814A		A						32 ± 6	82 ± 3
R814S		S						33 ± 5	85 ± 2
F813A	A							17 ± 2	56 ± 5
F813V	V							20 ± 5	64 ± 7

^a The substrates used in the study were all 12-mers. Only the P6–P1' sites are shown. Substitutions were as indicated. The other sites (not shown) are the same as in the original Gly818–Ala819 oligopeptide FRLKGGAPIKGV. ^b Percentage cleavage is the average of three independent measurements using three different batches of enzyme. ^c Cleavage is not detectable.

Table 4: Summary of the Substrates of Some Coronaviral PLPs

virus	peptide substrate ^a								protease ^c	references
	AA ^b	P6	P5	P4	P3	P2	P1	P1'		
SARS-CoV	175	R	E	L	N	G	G	A	PLP2	this paper
SARS-CoV	813	F	R	L	K	G	G	A	PLP2	this paper
SARS-CoV	2735	I	S	L	K	G	G	K	PLP2	this paper
MHV	2832	F	S	L	K	G	G	A	PLP2	22
BCoV	2745	F	S	L	K	G	G	A	PLP2 ?	38
HCoV-229E	892	F	T	K	A	A	G	G	PLP1/2	14
IBV	668	V	V	C	K	A	G	G	PLP2	15
IBV	2260	V	E	K	K	A	G	G	PLP2	16
MHV	242	L	K	G	Y	R	G	V	PLP1	36, 37
MHV	827	W	R	F	P	C	A	G	PLP1	11
BCoV	241	I	R	G	Y	R	G	V	PLP1 ?	38
BCoV	846	W	R	V	P	C	A	G	PLP1 ?	38
HCoV-229E	106	G	K	R	G	G	G	N	PLP1	12
HCoV-229E	892	F	T	K	A	A	G	G	PLP1/2	14

^a Genbank accession numbers are AY291451 (SARS-CoV-TW1), Q9PYA3 (MHV), Q66198 (BCoV), Q05002 (HCoV-229E), and P27920 (IBV), respectively. The horizontal line separates the coronaviral PLP2s from PLP1s. ^b The starting position of the amino acid corresponds to the P6 site. ^c The question mark denotes unknown or unconfirmed information.

P6 (Phe) position was substituted with Ala or Val (G817A, F813A, or F813V) (Table 3). The substitution of P5 (Arg) with either Ala or Ser (R814A or R814S) resulted in moderately reduced cleavage, whereas substitution of P3 (Lys) with either Ala or Gln (K816A and K816Q) did not affect cleavage at all. Furthermore, substitution at the P1' (Ala) position with either Gly or Asn (A819G and A819N) did not affect cleavage at all. Therefore, the P1, P2, P4, and P6 positions are important for optimal substrate recognition and cleavage by SARS-CoV PLP2, also consistent with the observation presented in Table 1.

Cleavage of the MHV and BCoV Substrates by SARS-CoV PLP2. The amino acid sequences are poorly conserved among the confirmed sites of cleavage by coronaviral PLP1s from MHV and HCoV-229E, except that the P1 site is either Gly or Ala (Table 4). Interestingly, we identified a consensus cleavage motif by coronaviral PLP2s, quite different from those recognized by PLP1s. SARS-CoV and MHV PLP2s share the consensus motif F/I-R/S-L-K-G-G from position

Table 5: SARS-CoV PLP2 Cleaves the Substrates of MHV and BCoV

source	peptide substrate ^a							cleavage % ^b	
	P6	P5	P4	P3	P2	P1	P1'	2 h	12 h
Gly180–Ala181	R	E	L	N	G	G	A	nd ^c	5 ± 1
Gly818–Ala819	F	R	L	K	G	G	A	70 ± 7	100
Gly2740–Lys2741	I	S	L	K	G	G	K	7 ± 1	29 ± 4
MHV	F	S	L	K	G	G	A	21 ± 5	80 ± 2
BCoV	F	S	L	K	G	G	A	27 ± 4	85 ± 9
HCoV-229E	F	T	K	A	A	G	G	nd	nd
IBV	V	E	K	K	A	G	G	nd	nd

^a The substrates used in the study were all 12-mers. Only positions P6–P1' are shown. The sequences have the following GenBank accession numbers: AY291451 (SARS-CoV-TW1), Q9PYA3 (MHV), Q66198 (BCoV), Q05002 (HCoV-229E), and P27920 (IBV). ^b Percentage cleavage is the average of three independent measurements using three different batches of enzyme. ^c Cleavage was not detectable.

P6 to P1, while PLP2s from HCoV-229E and IBV share the conserved motif A-G-G at the corresponding P2–P1–P1' positions, with the residues at the remaining positions divergent (Table 4). Notably, one of the sites in BCoV predicted to be cleaved by its PLP has the sequence F-S-L-K-G-G from position P6 to P1 (38), matching the consensus sequence of F/I-R/S-L-K-G-G postulated in this study (Table 4). We were interested to know whether SARS-CoV PLP2 cleaves these homologous sites of MHV and BCoV.

At least 80% of the MHV or BCoV substrate was cleaved in a 12 h incubation, compared with 100% cleavage at the Gly818–Ala819 site of SARS-CoV (Table 5). These two substrates from MHV and BCoV were even better *in vitro* than the other two sites of SARS-CoV, Gly180–Ala181 and Gly2740–Lys2741. After 2 h of digestion, SARS-CoV PLP2 cleaved about 21 and 27% of the MHV and BCoV substrates, respectively, whereas it cleaved only 7% of its own Gly2740–Lys2741 site and did not cleave the Gly180–Ala181 site at all (Table 5). Furthermore, SARS-CoV PLP2 did not cleave the sites from HCoV-229E and IBV even after 12 h of digestion (Table 5). Therefore, SARS-CoV PLP2 shares substrate homology with those of MHV and BCoV, and it cleaves these homologous substrates from them. The epidemiological and evolutionary implications of these results are discussed below.

An Effective Inhibitor of SARS-CoV PLP2 by Inhibitor-Screening Platform. We designed a fluorogenic inhibitor-screening platform using only nanomolar amounts of SARS-CoV PLP2, which is suitable for adaptation to high throughput screening. Interestingly, zinc ion inhibited the protease activity potently with the IC₅₀ value of 1.3 μM (data not shown). Zinc conjugates, including *N*-ethyl-*N*-phenyldithiocarbamic acid Zn and hydroxypridine-2-thione Zn were also effective in inhibiting SARS-CoV PLP2, with the IC₅₀ values of 3.3 and 3.7 μM, respectively. The inhibition is specific because other divalent metals, such as Mg, Mn, Ca, Ni, and Co, had no effects on the activity of SARS-CoV PLP2 at 10 μM (data not shown). Cu ion at 10 μM weakly inhibited the activity of the PLP2 to 70% (data not shown). Although the mechanism of this inhibition by zinc is not yet understood, we used this as a control to investigate the inhibitory effects of some commercially available and known cysteine protease inhibitors. None of the known cysteine protease inhibitors, including E64 at 0.1 mM, *N*-ethylmaleimide at 1 mM,

cystatin at 10 $\mu\text{g/mL}$, leupeptin at 0.1 mM, antipain at 0.1 mM, and chymostatin at 1 mM, were effective in inhibiting SARS-CoV PLP2 (data not shown). These results suggest that the catalytic dyad of SARS-CoV PLP2 is not readily accessible to these inhibitory compounds, and zinc ion is an effective inhibitor of SARS-CoV PLP2.

DISCUSSION

The controlled proteolysis of the replicase polypeptides by viral proteases is essential for viral virulence and biogenesis. In this study, we demonstrated that, despite low sequence homology (Figure 2), SARS-CoV PLP2 catalytically is a "papain-like" enzyme with similar catalytic mechanism as that of papain, providing the first clue on the reaction mechanism of SARS-CoV PLP2. Catalytically, SARS-CoV PLP2 is different from cathepsins, papain-like cysteine proteases also belonging to the papain superfamily but with a narrow acidic pH optima (39, 40). It will be interesting to determine whether other coronaviral PLPs share similar enzymatic properties with SARS-CoV PLP2. Even though the catalytic dyad is ubiquitously present among coronaviral PLPs and papain (34, 35, 41–43), the residues around the Cys and His dyad are quite different (Figure 2). The nucleophilic Cys is preceded by Asn and followed by a hydrophobic (Y, F, or W) residue in coronaviral PLPs, while there is a signature motif GSCWAFS at the Cys active site of papain. In coronaviral PLPs, the dyad residue His is preceded by a Gly (or Cys and Ser for PLP1s of BCoV and MHV, respectively) and followed by several hydrophobic residues, while Asp precedes the active-site His of papain. Structurally, acidic groups nearby the dyad might contribute electrostatically to the modulation of the catalytically competent dyad ion pair (Cys-His⁺) (34). The wide pH–activity profiles indicate that this ionized thiolate–imidazolium ion pair is relatively more stable than is its neutral form (35). PLP2s from both MHV and SARS-CoV are not inhibited by most commonly used cysteine protease inhibitors (see ref 13 and this study). In contrast, PLP1 from MHV is sensitive to these inhibitors, as shown by transfection studies (13, 19, 37, 44, 45). These data suggest that the substrate-binding sites or the environments of the catalytic dyad in coronaviral PLP2s might be different significantly from those in PLP1s or other E64-sensitive cysteine proteases, raising a possibility that structurally PLP1s might differ from their PLP2 counterparts.

We demonstrate that SARS-CoV PLP2 by itself is active, independent of contributions from the cellular environment or other cellular factors. Mass spectrometry to determine the cleavage site is a sensitive alternative compared to the previously used method (14, 36, 46). Employing the method of HPLC coupled with mass spectrometry, we not only confirm without a doubt the cleavage sites by SARS-CoV PLP2 but also detect the enzymatic activity of SARS-CoV not observable *in vivo* as described in Hartcourt et al. (20). In our study, cleavage at the Gly2740–Lys2741 site was detected by RP-HPLC (Figure 4C), although it was not detected in an *in vivo* study with a slightly longer PLP2 (residues 1540–2204) (20). Only when the C-terminal region was extended to include a hydrophobic domain (residues 1540–2425) did cleavage of this Gly2740–Lys2741 site occur *in vivo* (20). Our study shows that the intrinsic activity of SARS-CoV PLP2 can process all three predicted sites with

differential activity and that the extent of cleavage differs from that observed in the *in vivo* study (20). Purified SARS-CoV PLP2 marginally cleaved the Gly180–Ala181 site *in vitro*, with the greatest activity observed at the Gly818–Ala819 site (Figure 4 and Table 5). However, the Gly180–Ala181 site was readily cleaved *in vivo*, seemingly with the greatest efficiency, whereas the other two sites, Gly818–Ala819 and Gly2740–Lys2741, were less efficiently processed (20). The different results *in vitro* (this study) and *in vivo* (20) are explainable because previous studies on other coronaviral PLPs have shown that the length and boundary of PLP domains affect the efficiency of the cleavage on the substrates *in vivo* (14, 23). Thus, the region outside the activity domain and/or the cellular localization of SARS-CoV PLP2 contributes to its activity observed *in vivo*.

SARS-CoV PLP2 is capable of cleaving the PLP2 substrate sites from MHV and BCoV but not those from HCoV-229E and IBV (Table 5). Accordingly, we speculate that the site from BCoV, FSLKGGG, is cleaved by its PLP2 rather than by its PLP1 (Table 5). On the basis of the common properties shared with MHV and BCoV on substrate homology and cleavage and insensitivity to E64, our study supports that SARS-CoV diverges early in evolution from MHV and BCoV (6, 47). Therefore, rather than classifying SARS-CoV as an independent group 4 as originally proposed (3, 4), our data provide evidence to classify SARS-CoV phylogenetically as group 2b relative to group 2a, which includes MHV and BCoV (6, 47). Supporting this, both viral proteins and RNA in the replicative machinery of MHV and SARS-CoV are similarly located in the membrane compartment *in vivo*, as demonstrated by electron microscopy and immunostaining experiments (20, 48–50). The information provided in this study is important for SARS-CoV-related research and epidemiology because it validates the use of MHV (and BCoV) as a suitable model with which to characterize SARS-CoV genes/proteins and functions (47).

Different from MHV PLP2 (22) and uniquely for SARS-CoV PLP2, P4 (Leu) is critical for substrate recognition and cleavage (Table 3). In addition, the P6 site might contribute through anchoring of the substrate to the active site (Table 1). Thus, it is likely that the peptidomimetic compound with a longer chain might work as an inhibitor to inhibit the enzymatic activities. Importantly, we found that SARS-CoV PLP2 is the molecular target for zinc inhibition. Because none of the structures of coronaviral PLPs is available, how the high concentration of zinc ion affects the activity of SARS-CoV PLP2 awaits further study. It is interesting to note that zincum gluconicum (Zenullose or ZICAM sold over the counter), a zinc salt and a homeopathic therapy, is effective in treating the common cold, a disease caused by rhinoviruses, although its molecular target is not yet known (51). In addition, zinc acetate is added as a supplement for the treatment of Wilson's disease, indicating that zinc is safe to use in humans (52). Previously, zinc and some zinc conjugates are found to be effective inhibitors for SARS-CoV 3CL protease (53). Therefore, using high dosage of zinc or zinc conjugates might be effective for inhibiting SARS-CoV replication *in vivo* through the simultaneous inhibition of SARS-CoV PLP2 and 3CL protease. Our detailed study of substrate preference and fluorogenic inhibitor-screening platform will also assist in the design and discovery of novel pharmacophores specific for PLP2 inhibition.

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