

Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.

Prediction of proteinase cleavage sites in polyproteins of coronaviruses and its applications in analyzing SARS-CoV genomes

Feng Gao^a, Hong-Yu Ou^a, Ling-Ling Chen^{a,b}, Wen-Xin Zheng^a, Chun-Ting Zhang^{a,*}

^aDepartment of Physics, Tianjin University, Tianjin 300072, PR China

^bLaboratory for Computational Biology, Shandong Provincial Research Center for Bioinformatic Engineering and Technique, Shandong University of Technology, Zibo 255049, PR China

Received 7 July 2003; revised 19 September 2003; accepted 22 September 2003

First published online 2 October 2003

Edited by Takashi Gojobori

Abstract Recently, we have developed a coronavirus-specific gene-finding system, ZCURVE_CoV 1.0. In this paper, the system is further improved by taking the prediction of cleavage sites of viral proteinases in polyproteins into account. The cleavage sites of the 3C-like proteinase and papain-like proteinase are highly conserved. Based on the method of traditional positional weight matrix trained by the peptides around cleavage sites, the present method also sufficiently considers the length conservation of non-structural proteins cleaved by the 3C-like proteinase and papain-like proteinase to reduce the false positive prediction rate. The improved system, ZCURVE_CoV 2.0, has been run for each of the 24 completely sequenced coronavirus genomes in GenBank. Consequently, all the non-structural proteins in the 24 genomes are accurately predicted. Compared with known annotations, the performance of the present method is satisfactory. The software ZCURVE_CoV 2.0 is freely available at http://tubic.tju.edu.cn/sars/.

© 2003 Published by Elsevier B.V. on behalf of the Federation of European Biochemical Societies.

Key words: Coronavirus; Severe acute respiratory syndrome; SARS-coronavirus; Polyprotein; Cleavage site

1. Introduction

Due to the severity of a life-threatening disease, referred to as severe acute respiratory syndrome (SARS), the World Health Organization (WHO) has issued a global alert for the illness. SARS apparently began in Guangdong province of China in November 2002, and has spread to Hong Kong, Singapore, Vietnam, Canada, the USA and several European countries [1–6]. By early June 2003, more than 700 SARSrelated deaths were recorded by WHO (http://www.who.int/ csr/sars/country/en/).

A novel coronavirus, called SARS-coronavirus or SARS-CoV, has been proved to be the cause of SARS. The coronaviruses (order *Nidovirales*, family *Coronaviridae*, genus *Coronavirus*) are members of a family of large, enveloped, positivestranded RNA viruses that replicate in the cytoplasm of animal host cells [7]. There are three groups of coronaviruses; groups I and II contain mammalian viruses, while group III contains only avian viruses. The viruses are associated with a variety of diseases in humans and domestic animals, including gastroenteritis and diseases of the upper and lower respiratory tract. Many researchers have analyzed the phylogeny of SARS-CoV and concluded that it is not closely related to any of the previously characterized coronaviruses and forms a distinct group (group IV) within the genus Coronavirus [7,8]. At the time this paper was written, there were 12 strains of SARS-CoV complete genome sequences available from Gen-Bank [7–9]. Among these genomes, six have been annotated manually, and the remaining six have not been annotated yet. The genomic organization of SARS-CoV is that of a typical coronavirus, with the order of the characteristic genes being replicase [rep], spike [S], envelope [E], membrane [M], nucleocapsid [N] from the 5' to the 3' terminus. SARS-CoV also encodes a number of non-structural proteins located between S and E, between M and N, or downstream of N with unknown functions. We have developed a coronavirus-specific gene-finding system ZCURVE_CoV 1.0 [10], which is especially suitable for gene recognition in SARS-CoV genomes. The software has the advantages of simplicity, reliability, high accuracy and quickness and can be obtained freely at the website http://tubic.tju.edu.cn/sars/. The system ZCURVE_CoV 1.0 has been run for each of the 12 SARS-CoV genomes. In addition to the polyprotein chains Orfla and Orf1b and the four genes encoding the major structural proteins, S, E, M and N, respectively, ZCURVE_CoV 1.0 also predicts five to six putative proteins between 39 and 274 amino acids in length, with unknown functions in SARS-CoV genomes. However, the cleavage sites of viral proteinase in replicases are not predicted in ZCURVE_CoV 1.0.

The coronavirus replicases are encoded by two large, 5'proximal open reading frames (ORFs) that comprise approximately two-thirds of the genome. Polyproteins ORF1a and ORF1b are connected by a ribosomal frameshift site, which is believed to occur at the conserved 'slippery sequence', UUUAAAC. It results in the translation of an ORF1a protein and a carboxyl-extended ORF1ab frameshift protein, which are also known as replicase polyproteins pp1a and pp1ab [11]. The ORF1a and ORF1ab translation products are polyprotein precursors, which are cleaved by viral proteinases, resulting in a minimum of 13 non-structural proteins, including a 3C-like proteinase, an RNA-dependent RNA polymerase, an ATPase/helicase and other function-unknown non-structural proteins [11]. These proteins in turn are responsible for replicating the viral genome as well as generating nested transcripts that are used in the synthesis of viral proteins. In this paper, all the putative non-structural proteins resulting

0014-5793/03/\$22.00 © 2003 Published by Elsevier B.V. on behalf of the Federation of European Biochemical Societies. doi:10.1016/S0014-5793(03)01091-3

^{*}Corresponding author. Fax: (86)-22-2740 2697.

E-mail address: ctzhang@tju.edu.cn (C.-T. Zhang).

Tal	ble	1

The lengths for 11 non-structural proteins^a cleaved by the 3C-like proteinase

Genome	The length of non-structural proteins (aa)												
	nsp2	nsp3	nsp4	nsp5	nsp6	nsp7	nsp9	nsp10	nsp11	nsp12	nsp13		
TOR2	306	290	83	198	113	139	932	601	527	346	298		
HCoV-229E ^b	302	279	83	195	109	135	927	597	518	348	300		
MHV ^b	303	287	92	194	110	137	928	600	521	374	299		
BCoV	303	287	89	197	110	137	928	603	521	374	299		
IBV ^b	307	293	83	210	111	145	940	600	521	338	302		
TGEV	302	294	83	195	111	135	929	599	519	339	300		
PEDV	302	280	83	195	108	135	927	597	517	339	301		
Average length ^c	304	287	85	198	110	138	930	600	521	351	300		
Standard deviation	2.07	5.87	3.76	5.59	1.60	3.60	4.67	2.15	3.26	16.07	1.35		

^aThese proteins are cleaved by the 3C-like proteinase within polyprotein 1ab derived from the seven coronavirus genomes annotated by NCBI. ^bThe cleavage sites have been confirmed by experimental evidence in these genomes.

^cThe genomes that have maximum lengths for nsp2–13 except nsp8 are IBV, TGEV, MHV, IBV, TOR2, IBV, BCoV, TOR2, MHV (BCoV) and IBV respectively. The genomes that have the minimum lengths for nsp2–13 except nsp8 are HCoV-229E (TGEV, PEDV), HCoV-229E, TOR2 (HCoV-229E, IBV, TGEV, PEDV), MHV, PEDV, HCoV-229E (TGEV, PEDV), HCoV-229E (PEDV), HCoV-229E (PEDV), HCoV-229E (PEDV), HCoV-229E (PEDV), IBV and TOR2, respectively.

from the cleavage by viral proteinases in the polyproteins are precisely predicted using ZCURVE_CoV 2.0.

2. Materials and methods

Seven genomic sequences of coronaviruses and the annotation information were downloaded from the NCBI RefSeq project. These coronaviruses include avian infectious bronchitis virus (IBV) (NC_001451), bovine coronavirus (BCoV) (NC_003045), human coronavirus 229E (HCoV-229E) (NC_002645), murine hepatitis virus (MHV) (NC_001846), porcine epidemic diarrhea virus (PEDV) (NC_003436), SARS coronavirus TOR2 (TOR2) (NC_004718) and transmissible gastroenteritis virus (TGEV) (NC_002306). The above genomes have been annotated by NCBI and the sequences of mature peptides are available. According to the annotation, a total of 77 sites cleaved by the 3C-like proteinase and 17 sites cleaved by the papainlike proteinase were extracted from the above seven genomes. Octapeptides cleaved by the 3C-like proteinase and 12-mer peptides cleaved by the papain-like proteinase were used to train the corresponding positional weight matrix (PWM) [12]. The cleavage site is at the center of the octapeptide or 12-mer peptide. The length distribution of non-structural proteins within ORF1ab was also derived from the annotated genomes. At the time this paper was written, there were 24 complete sequences of coronavirus genomes available in the GenBank database, of which 12 are SARS-CoVs and 12 are other groups of coronaviruses. The former comprises SARS-CoV TOR2 (NC_004718), Urbani (AY278741), HKU-39849 (AY278491), CUHK-W1 (AY278554), BJ01 (AY278488), CUHK-Su10 (AY282752), SIN2500 (AY283794), SIN2748 (AY283797), SIN2679 (AY283796), SIN2774 (AY283798), SIN2677 (AY283795) and TW1 (AY291451), whereas the latter comprises IBV (NC_001451), BCoV (NC_003045), bovine coronavirus strain Mebus (BCoVM) (U00735), bovine coronavirus isolate BCoV-LUN (BCoVL) (AF391542), bovine coronavirus strain Quebec (BCoVQ) (AF220295), HCoV-229E (NC_002645), MHV (NC_001846), murine hepatitis virus strain ML-10 (MHVM) (AF208067), murine hepatitis virus strain 2 (MHV2) (AF201929), murine hepatitis virus strain Penn 97-1 (MHVP) (AF208066), PEDV (NC_003436) and TGEV (NC_002306).

The mature peptides cleaved by the 3C-like proteinase are highly conserved in length among different groups of coronaviruses, while others cleaved by the papain-like proteinase are not so conserved. The lengths of all the non-structural proteins cleaved by the 3C-like proteinase within polyprotein 1ab are listed in Table 1, while the lengths for the non-structural proteins cleaved by the papain-like proteinase are listed in Table 2. The average length and standard deviation for each kind of non-structural proteins are calculated. As shown in Tables 1 and 2, the lengths of the non-structural proteins cleaved by the 3C-like proteinase are highly conserved, while the lengths and the number of the papain-like cysteine proteinase cleavage products (abbreviated as PCP CP) appear to be irregular. Since the NCBI annotations are not always correct, the annotations of cleavage products of

the papain-like proteinase may be incomplete. It is observed that the size of the annotated PCP CP3 of SARS-CoV, MHV and IBV is approximately the sum of the sizes of PCP CP3 and PCP CP4 of other mammalian coronaviruses listed in Table 2. Therefore, the PCP CP3 of SARS-CoV, MHV and IBV may be further cleaved, i.e. it is possible that another papain-like proteinase cleavage site is present in the PCP CP3 of SARS-CoV, MHV and IBV. Based on the above analysis, a cleavage model of the papain-like proteinase is presented schematically in Fig. 1. According to this model, all coronaviruses have four non-structural proteins cleaved by the papain-like proteinase. Consequently, the cleavage products of the papain-like proteinase predicted by this model show the conservation in both their length and number. The average length and standard deviation for each papain-like proteinase cleavage product are estimated based on the genomes of BCoV, HCoV-229E, TGEV and PEDV, in which four of the papain-like proteinase cleavage products are annotated (see Table 2). Fig. 2A,B shows the conservation sites cleaved by the 3Clike proteinase and papain-like proteinase, respectively. It can be seen that both the 3C-like proteinase and papain-like proteinase have conserved cleavage sites. The same arrangement order of the cleavage products in polyprotein 1ab, similar sizes of non-structural proteins and the conserved residues in the cleavable peptides form the basis of the present algorithm to predict cleavage sites of polyproteins. Here, the method is described briefly as follows.

First, ORF1ab and the slippery sequences are identified using ZCURVE_CoV 1.0. Subsequently, the predicted ORF1ab is translated into amino acid sequence. Starting from the C-terminus of the predicted ORF1ab polyprotein, the candidate cleavage site of nsp13 is searched within a particular region using the sliding-window tech-

Table 2

The lengths for the non-structural proteins^a cleaved by the papainlike proteinase

Genome	Length (aa)									
	PCP CP1	PCP CP2	PCP CP3	PCP CP4						
IBV	_	673 ^b	2106	_						
TOR2	179	639	2422	_						
MHV	247 ^b	585 ^b	2501	_						
BCoV	246	605	1899	496						
HCoV-229E	111 ^b	786	1587 ^b	481						
TGEV	108	771	1509	490						
PEDV	110	785	1622	480						
Average length ^c	144	737	1654	487						
Standard deviation ^c	68.18	88.10	169.87	7.63						

^aThese proteins are cleaved by the papain-like proteinase within polyprotein lab derived from the seven coronavirus genomes annotated by NCBI.

^bThese cleavage products have been confirmed by experimental evidence.

^cThe average length and standard deviation are calculated based on the genomes of BCoV, HCoV-229E, TGEV and PEDV.



Fig. 1. Comparison between the N-terminal sequences of the polyprotein labs in MHV and BCoV is shown schematically. The additional cleavage site in the annotated PCP CP3 predicted by the present method for MHV is situated at the corresponding position where the PCP CP3 and PCP CP4 are cleaved in BCoV. Cleavage sites that have been annotated by NCBI are indicated by black arrows, while the cleavage site predicted by the present method is indicated by an open arrow.

nique. The distance between the scanning region center and the C-terminus of polyprotein 1ab should be equal to the average length of nsp13. Denoting the center position by c, a window with an octapeptide size slides from the positions $c-3\delta$ to $c+3\delta$, where δ is the standard deviation of the length distribution for nsp13 (see Table 1). Given an octapeptide within the region $S = X_4 X_3 X_2 X_1 X_{1'} X_{2'} X_{3'} X_{4'}$, where X_i (i=4, 3, 2, 1, 1', 2', 3', 4') represents the amino acid at the position P_i , the score of the octapeptide is computed as

Score
$$(\mathbf{X}_4 \mathbf{X}_3 \mathbf{X}_2 \mathbf{X}_1 \mathbf{X}_{1'} \mathbf{X}_{2'} \mathbf{X}_{3'} \mathbf{X}_{4'}) = \prod_{i=4}^{4'} f(i, \mathbf{X}_i)$$
 (1)

where $f(i, X_i)$ (*i*=4, 3, 2, 1, 1', 2', 3', 4') is the frequency of amino acid X_i occurring at the position P_i , which is an element in the corresponding positional weight matrix. The site with maximum score is selected as a candidate site. Consequently, the cleavage site of nsp12|13 is determined and nsp13 is found.

Prediction of other cleavage sites is performed in a recurrent way. Once the cleavage site of nsp12|13 is determined, the next cleavage site to be predicted is nsp11|12, then nsp10|11, and so forth until nsp1|2. Generally, if the site of nspk|(k+1) is determined, the next target is to predict the site of nsp(k-1)|k, where k = 12, 11, ..., 2, but $k \neq 8$ (see the explanation below). For clarity, take k = 6 as an example, where the site of nsp6|7 is known. First, the center position and the sliding window used for identifying the site of nsp5|6 need to be determined. The center position c is situated upstream of the site of nsp6|7. The distance between the center position c and the site of nsp6|7 should be equal to \overline{l}_6 , which is the average length of nsp6. In Table 1, we find $\bar{l}_6 = 110$ aa and the standard deviation δ of the length distribution for nsp6 is 1.6. A window with an octapeptide size thus slides from the position $c-3\delta \approx c-5$ to $c+3\delta \approx c+5$. Second, the site with the highest score is predicted to be the candidate site of nsp5|6. Note that in some cases the scores may be zero because of the limited training samples. In this case, a very small quantity (0.001) is assigned to the zero elements in the positional weight matrix. Also note that the nsp7|8 site is cleaved in polyprotein 1a, while the nsp7|9 site is cleaved in polyprotein 1ab. Therefore, the cleavage sites of nsp7|8 and nsp7|9 are in fact the same, leading to the result of $k \neq 8$. Furthermore, if the following two conditions are satisfied, besides the site with the maximum score, the site with the second maximum score is also taken into account: (i) Gln and Leu are found at the P_1 and P_2 positions, respectively; (ii) the distance between the two sites is less than five amino acid residues. This procedure considers the prediction of two adjacent cleavage sites in the scanning window. Consequently, two alternative cleavage sites annotated by NCBI are also found in the genomes of MHV and BCoV. Note that such cases occur rarely in the genomes studied.

Repeating the above procedure 11 times, all of the mature peptides cleaved by the 3C-like proteinase are identified one by one. Then, the papain-like proteinase cleavage products are searched within the remaining regions of polyprotein 1ab. A similar recurrent procedure is performed to search for the papain-like proteinase cleavage sites. The scores of 12-mer peptides are calculated as described above. The center position and the size of the sliding window used to search for the papain-like proteinase cleavage sites are determined in a way similar to that used for the 3C-like proteinase. The sites associated with the maximum scores in the corresponding scanning regions are predicted to be cleavage sites. Consequently, three papain-like proteinase cleavage sites are predicted for each genome.



Fig. 2. Conservation of the sites cleaved by coronavirus proteinases. Two separate multiple, gap-free alignments around the P1|P1' positions of the sites cleaved by the 3C-like proteinase (A) and papain-like proteinase (B) in the training set are converted to logo presentations in which the size of an amino acid is proportional to its conservation at the specific position and the sampling size. The amino acid conservation is measured in bits of information plotted on a vertical axis whose upper limit is determined by the natural diversity of amino acids (20) expressed as a logarithm of 2 [16]. Seventy-seven sites cleaved by the 3C-like proteinase were used to generate the logo in A, and 17 sites cleaved by the papain-like proteinase were used to generate the logo in B.

3. Results and discussion

Replicase polyprotein processing is carried out by two or three ORF1a-encoded viral proteinases. Coronaviruses encode a chymotrypsin-like proteinase, 3C-like proteinase, which is analogous to the main picornaviral proteinase, 3C proteinase [11]. As mentioned above, the cleavage sites of the 3C-like proteinase are highly conserved. As shown in Fig. 2A, the P_1 position of the peptide sequence is exclusively occupied by Gln. Leu is dominant at the P_2 position (more than 75%) and Val, Ser, Thr and Pro are clearly favored at the P_4 position. At the $P_{1'}$ position, small, aliphatic residues (Ser, Ala, Asn, Gly and Cys) are found, of which the content of Ser is more than 50%. There are no highly favored residues at the P_3 , $P_{2'}$, $P_{3'}$ and $P_{4'}$ positions. The length distributions of each of the 11 non-structural proteins cleaved by the 3C-like proteinase in the annotated genomes are listed in Table 1. Of these non-structural proteins, nsp2 is the putative 3C-like proteinase; nsp3 contains a hydrophobic domain; nsp7 is known as a growth-factor-like protein; nsp9 is the putative RNAdependent RNA polymerase; nsp10 contains a metal ionbinding domain and NTPase/helicase domain. Recently the mRNA cap-1 methyltransferase function has been assigned to nsp13 [13]. The functions of other non-structural proteins are unknown. Moreover, coronaviruses also encode one (group III) or two (groups I and II) papain-like proteinases,

which are analogous to the foot and mouth disease virus leader proteinase. SARS-CoV appears to contain only one papain-like proteinase domain in the predicted gene product of ORF1a [7]. For the papain-like proteinase, the cleavage sites are also conserved, but not as conserved as those of the 3C-like proteinase. Gly and Ala are found at the P₁ position and Gly accounts for more than 75%. At the P₂ and P_{1'} positions, Gly is also the dominant residue, which accounts for more than 45% and 50%, respectively. No residues exceed 40% at other positions. In this study, similar sizes of non-structural proteins and conserved cleavage sites form the basis of the present algorithm.

The performance of the algorithm is satisfactory by comparing the predicted results with known annotations. Although all the SARS genomes have been annotated by in silico analysis so far, some annotations for other coronaviruses, such as IBV, MHV and HCoV-229E, are supported by experimental evidence [11]. The jack-knife (leave-one-out) test has been performed here to ensure the validation of the prediction results for the cleavage sites of the 3C-like proteinase. By the jack-knife test, each genome out of the seven genomes under study is singled out in turn, and used as a testing genome. The remaining six genomes are used as the training set. Based on the data derived from the six training genomes, the cleavage sites of the 3C-like proteinase in the testing genome are predicted and evaluated. The jack-knife test was

Table 3

Comparison	n of the	predicted	results t	for	TGEV	and	PEDV	with	those	annotated	by	NCBI ^a
------------	----------	-----------	-----------	-----	------	-----	------	------	-------	-----------	----	--------------------------

Number	Genome	Location	(bp)	Location (aa)		Length (aa)	Cleavable peptide	Feature
		Start	Stop	Start	Stop	-		
1	TGEV	315	638	1	108	108	_	PCP CP1
	NC_002306	639	2951	109	879	771	KIARTG RGAIYV	PCP CP2
		2 9 5 2	7 4 7 8	880	2 388	1 509	YNKMGG GDKTVS	PCP CP3
		7 479	8 948	2 389	2878	490	VSPKSG SGFFDV	PCP CP4
		8 949	9854	2879	3180	302	STLQ SGLR	nsp2
		9855	10736	3 1 8 1	3 4 7 4	294	VNLQ AGKV	nsp3
		10737	10985	3 4 7 5	3 5 5 7	83	STVQ SKLT	nsp4
		10 986	11 570	3 558	3752	195	TILQ SVAS	nsp5
		11 571	11903	3 7 5 3	3863	111	TKLQ NNEI	nsp6
		11 904	12308	3 864	3 998	135	VRLQAGKP	nsp7
		12 309	15094	3 999	4927	929	TSMQ SFTV	nsp9 ^b
		15095°	16891°	4928	5 5 2 6	599	TVLQ AAGM	nsp10
		16892 ^c	18 448 ^c	5 527	6045	519	IGLQ AKPE	nsp11
		18 449 ^c	19465 ^c	6 0 4 6	6384	339	KALQ SLEN	nsp12
		19 466 ^c	20 365°	6385	6684	300	PQLQ SAEW	nsp13
2	PEDV	297	626	1	110	110	_	PCP CP1
	NC_003436	627	2981	111	895	785	FGRRGG NIVPVD	PCP CP2
		2 982	7847	896	2517	1 622	FKKKGG GDVKFS	PCP CP3
		7 848	9 287	2 5 1 8	2997	480	ANKKGA GLPSFS	PCP CP4
		9 288	10193	2 998	3 299	302	STLQ AGLR	nsp2
		10 194	11033	3 300	3 579	280	VNLQ GGYV	nsp3
		11 034	11 282	3 580	3 6 6 2	83	SSVQ SKLT	nsp4
		11 283	11867	3 6 6 3	3857	195	SMLQ SVAS	nsp5
		11 868	12192	3858	3965	108	VKLQ NNEI	nsp6
		12 191	12 596	3 966	4100	135	VRLQAGKQ	nsp7
		12 597	15376	4101	5027	927	SIMQ STDM	nsp9 ^d
		15 377	17167	5 0 2 8	5624	597	AVLQ SAGL	nsp10
		17 168	18718	5625	6141	517	SDLQANEG	nsp11
		18719	19735	6142	6480	339	NNLQ GLEN	nsp12
		19 736	20 638	6481	6781	301	PQLQ ASEW	nsp13

^aNote that of the 24 coronavirus genomes, the predicted results by ZCURVE_CoV 2.0 are in complete agreement with those annotated by NCBI, except for the genomes of TGEV and PEDV, in which the predicted results are different from those annotated by NCBI. In this table the reasons for these conflicts are analyzed.

^bThis conflict with the annotation is caused by the problematic annotation.

^cThe locations are different from the annotation, which is caused by a questionable additional insertion of an amino acid residue in nsp9.

^dThis conflict with the annotation is caused by the non-standard frameshift.

Tal	bl	le	4

The predicted results by the present method for BCoVL and SARS-CoV BJ01

Number Genome		Location	Location (bp)		n (aa)	Length (aa)	Cleavable peptide	Feature	
		Start	Stop	Start	Stop	_			
1	BCoVL	211	948	1	246	246	_	PCP CP1	
	AF391542	949	2763	247	851	605	IRGYRG VKPLLY	PCP CP2	
		2 764	8 4 6 0	852	2750	1 899	WRVPCA GRRVTF	PCP CP3	
		8 461	9 948	2751	3 2 4 6	496	FSLKGG AVFSYF	PCP CP4	
		9 949	10857	3 2 4 7	3 549	303	SFLQ SGIV	nsp2	
		10858	11718	3 5 5 0	3 8 3 6	287	IKLQ SKRT	nsp3	
		11719	11985	3837	3 9 2 5	89	SQFQ SKLT	nsp4	
		11986	12 576	3926	4122	197	TVLQ ALQS ^a	nsp5	
		12 577	12906	4123	4 2 3 2	110	TVLQ NNEL	nsp6	
		12907	13317	4 2 3 3	4 369	137	VRLQ AGTA	nsp7	
		13 318	16100	4 3 7 0	5 297	928	TTVQ SKDT	nsp9	
		16101	17909	5 298	5 900	603	AVMQ SVGA	nsp10	
		17910	19472	5901	6421	521	TRVQ CSTN	nsp11	
		19473	20 594	6422	6 795	374	TKLQ SLEN	nsp12	
		20 595	21 491	6 796	7 094	299	PRLQ AASD	nsp13	
2	BJ01	246	782	1	179	179	_	PCP CP1	
	AY278488	783	2 699	180	818	639	TRELNG GAVTRY	PCP CP2	
		2 700	8 465	819	2 7 4 0	1 922	FRLKGG APIKGV	PCP CP3	
		8 466	9965	2741	3 2 4 0	500	ISLKGG KIVSTC ^b	PCP CP4	
		9 966	10883	3 2 4 1	3 546	306	AVLQ SGFR	nsp2	
		10884	11753	3 547	3 8 3 6	290	VTFQ GKFK	nsp3	
		11754	12002	3837	3919	83	ATVQ SKMS	nsp4	
		12 003	12 596	3 9 2 0	4117	198	ATLQ AIAS	nsp5	
		12 597	12935	4118	4 2 3 0	113	VKLQ NNEL	nsp6	
		12936	13 352	4 2 3 1	4 369	139	VRLQ AGNA	nsp7	
		13 353	16147	4 3 7 0	5 301	932	PLMQ SADA	nsp9	
		16148	17950	5 302	5 902	601	TVLQ AVGA	nsp10	
		17951	19 531	5903	6429	527	ATLQ AENV	nsp11	
		19 532	20 569	6430	6775	346	TRLQ SLEN	nsp12	
		20 570	21 463	6776	7 0 7 3	298	PKLQ ASQA	nsp13	

^aThe alternative cleavage site predicted by the present method is at QALQ|SEFV (Gln-3928|Ser-3929).

^bCompared with the annotation, this cleavage site is predicted additionally by the present method.

finished by repeating the above procedure seven times. Consequently, the predicted results by the jack-knife test are found to be as good as those by a self-consistency test mentioned previously, suggesting that the prediction results are reliable.

The prediction results for TGEV and PEDV, which are different from the annotations of NCBI RefSeq projects, are listed in Table 3. The prediction results for other genomes can be obtained from the supplementary materials (http://tubic. tju.edu.cn/sars/). The coronavirus -1 frameshift site [14] is believed to occur at the 'slippery sequence', UUUAAAC. This assumption has been supported by experimental evidence [15]. But the annotated frameshift sites are not always consistent with this pattern, as in the case of PEDV, whose frameshift site lies upstream of the UUUAAAC sequence according to the annotation. This may be due to the questionable annotation. For example, the genomes of MHV and BCoV were originally annotated by the authors as the ones having a nonstandard frameshift site, however, these conclusions were then corrected by the re-annotations of NCBI as the ones having standard frameshift sites. In light of this, we adopt UUUA-AAC as the standard slippery sequence.

Using the present method, only few false positive predictions exist in the prediction results. The tedious calculations for deriving the cutoff value can be avoided by restricting the sizes of the scanning regions and only selecting the site with the maximum score within this region. The annotated cleavage sites often correspond to the highest scores measured by the PWM method. However, the sites scored high by the PWM method do not always correspond to the cleavage sites and vice versa. Restricting the scanning regions for each of the cleavage sites is more efficient to reduce the false positive prediction rate. For the prediction of the 3C-like proteinase cleavage sites, there are only two conflicts between the predicted results and the annotations, which are marked in Table 3. The first conflict lies in the locations of non-structural proteins downstream of nsp9 in TGEV, which may be due to the problematic annotation. The length of amino acid sequences for ORF 1ab (315-20368 bp) should be 6684 aa, instead of 6685 aa, which is annotated by NCBI. The questionable additional insertion of an amino acid residue in nsp9 causes one conflict of location errors. The second is caused by a nonstandard frameshift site in PEDV, which causes the difference of five amino acid residues between the non-standard frameshift site and the standard frameshift site. For this reason, the octapeptide predicted by the present method is SIMQ|STDM instead of the annotated SIMQ|STDY.

Using the cleavage model of the papain-like proteinase presented here, the additional cleavage sites in the annotated PCP CP3 predicted by this method for SARS-CoV TOR2, MHV and IBV are ISLKGG|KIVSTC, FSLKGG|AVFSRM and VEKKAG|GIVSGT, respectively. The predicted cleavable peptides are similar to those annotated by NCBI, for example, the cleavable peptide FSLKGG|AVFSRM in MHV is different from the annotated peptide FSLKGG|AVFSYF in BCoV only at the $P_{5'}$ and $P_{6'}$ positions. Comparison between the N-terminal sequences of the polyprotein 1abs in MHV and BCoV is shown in Fig. 1. The additional cleavage site in the annotated PCP CP3 predicted by this method for MHV is situated at the corresponding position where the PCP CP3 and PCP CP4 are cleaved in BCoV. Cleavage sites that have been annotated by NCBI are indicated by black arrows, whereas that predicted by the present method is indicated by the open arrow. Therefore, the annotated PCP CP3 of SARS-CoV TOR2, MHV and IBV may be a precursor, which can be cleaved further.

Based on the present method, the genomes without annotation have been annotated. To save printing space, only the results of BCoVL and SARS-CoV BJ01 are summarized in Table 4. The detailed annotations for other coronavirus genomes are accessible at http://tubic.tju.edu.cn/sars/.

4. Conclusion

SARS is an extremely severe disease, which has spread to many countries around the world. Evidence shows that SARS is caused by a new coronavirus, i.e. SARS-CoV. A system, called ZCURVE_CoV 1.0, has been developed previously to recognize protein-coding genes in coronavirus genomes, especially suitable for SARS-CoV genomes [10]. Here an improved version of the system, ZCURVE_CoV 2.0, has been developed to identify all the non-structural proteins cleaved by viral proteinases in the polyproteins. Consequently, all the non-structural proteins in the 24 completely sequenced coronavirus genomes are predicted. Compared with the known annotations, including those based on experimental evidence, the performance of the present method is satisfactory. Acknowledgements: We are indebted to Prof. Jingchu Luo of Peking University for the timely updated SARS-related information provided. We are also grateful to both referees for their constructive comments, which are very useful to improve the quality of the paper. Invaluable assistance from Ren Zhang is gratefully acknowledged. The present study was supported in part by the 973 Project of China (Grant 1999075606).

References

- [1] Peiris, J.S. et al. (2003) Lancet 361, 1319-1325.
- [2] Ksiazek, T.G. et al. (2003) New Engl. J. Med. 348, 1953-1966.
- [3] Drosten, C. et al. (2003) New Engl. J. Med. 348, 1967–1976.
- [4] Tsang, K.W. et al. (2003) New Engl. J. Med. 348, 1977–1985.
- [5] Lee, N. et al. (2003) New Engl. J. Med. 348, 1986–1994.
- [6] Poutanen, S.M. et al. (2003) New Engl. J. Med. 348, 1995–2005.
- [7] Rota, P.A. et al. (2003) Science 300, 1394–1398.
- [8] Marra, M.A. et al. (2003) Science 300, 1399-1404.
- [9] Qin, E'd. et al. (2003) Chin. Sci. Bull. 48, 941–948.
- [10] Chen, L.L., Ou, H.Y., Zhang, R. and Zhang, C.-T. (2003) Biochem. Biophys. Res. Commun. 307, 382–388.
- [11] Ziebuhr, J., Snijder, E.J. and Gorbalenya, A.E. (2000) J. Gen. Virol. 81, 853–879.
- [12] von Heijne, G. (1986) Nucleic Acids Res. 14, 4683-4690.
- [13] von Grotthuss, M., Wyrwicz, L.S. and Rychlewski, L. (2003) Cell 113, 701–702.
- [14] Brierley, I., Jenner, A.J. and Inglis, S.C. (1992) J. Mol. Biol. 227, 463–479.
- [15] Nam, S.H., Copeland, T.D., Hatanaka, M. and Oroszlan, S. (1993) J. Virol. 67, 196–203.
- [16] Schneider, T.D. and Stephens, R.M. (1990) Nucleic Acids Res. 18, 6097–6100.