## SEQuENCE AND STRUCTURE OF THE CORONAVIRUS PEPLOMER PROTEIN

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## INTRODUCTION

Coronaviruses display a characteristic fringe of large (17-20 nm), clubshaped peplomers, each consisting of a di- or trimer of the peplomer protein (Cavanagh et al. 1983). The peplomer protein, E2, plays an important role during the infection proces. It mediates the binding of virions to the host-cell receptors and is involved in membrane fusion. In addition, the E2 protein appears to be a major inducer of protective immunity to coronaviral infection (reviewed by Sturman and Holmes, 1983).

In the case of mouse hepatitis virus (MHV) and avian infectious bronchitis virus (IBV), the peplomer protein is synthesized as an Nglycosylated precursor of about 180K (reviewed by Siddell et al., 1982). This precursor is proteolytically cleaved, yielding two products of 80 to 90K, which remain noncovalently associated. Cleavage of the MHV peplomer protein is thought to be essential for the cell fusion activity (Sturman et a1., 1985).

The peplomer proteins of the feline infectious peritonitis virus (FIPV) and the closely related porcine transmissible gastroenteritis virus (TGEV) differ from those of MHV and IBV in two respects. Firstly, proteolytic cleavage does not occur, although FIPV is fully capable to induce cell-fusion. Secondly, the peplomer proteins of FIPV and TGEV are larger, about 210K (Sidde11 et a1., 1982; Boyle et a1., 1984; Jacobs et al., 1986).

MHV, IBV and the FIP/TGE viruses are representatives of three separate antigenic clusters within the coronaviridae family (Siddell et al., 1983). Comparison of the nucleocapsid proteins of TGEV, MHV and IBV revealed a homology of about $27 \%$ in all three cases (Kapke and Brian, 1986), indicating a high degree of divergence.

In this report, we present a comparison of the peplomer proteins of IBV strain M41 (Niesters et al., 1986), MHV strain A59 and FIPV strain 791146. Because of the low overall homology (see below), the sequences that have been conserved in all three proteins are likely to be essential for common structural and/or functional features.

Cloning and sequencing of the peplomer genes of FIPV 79-1146 (de Groot et al, in prep.) and MHV A59 (Luytjes et al., in prep.) will be presented in detail elsewhere. The primary structure of the peplomer proteins was deduced from the nucleotide sequences. Apoproteins of 1162 , 1324 and 1452 amino acid residues were predicted for IBV (Niesters et al., 1986), MHV and FIPV, respectively. At the N-terminal end of the deduced peplomer sequences a hydrophobic, 17-20 residue segment is found (Fig. 2). Binns et al. (1985) reported that this segment is absent in the mature peplomer protein of IBV. These findings suggest that the peplomer proteins are synthesized with a transient, N-terminal signal peptide (Wickner and Lodish, 1985). About 60 to 70 residues upstream of the C-terminal end, a distinct, hydrophobic region is found (Fig. 2), which most probably serves as a transmembrane anchor (Binns et a1. 1985; Niesters et al., 1986).

As apparent from a Diagon comparison (Staden, 1982) of the amino acid sequences of the FIPV and MHV peplomer proteins (Fig. 1), most conserved residues are found in the C-terminal $60 \%$ of the protein.

FIPV $\rightarrow$


Fig. 1. Diagon comparison of the amino acid sequences of the FIPV and MHV peplomer proteins (proportional matching, span length: 21, minimal score: 221; Staden, 1982). The arrows indicate regions with an apparent repetitive character.

A more detailed comparison was made by combining FASTP alignments (Lipman and Pearson, 1985), Diagon plots and visual inspection (shown in Fig. 3). A schematic representation is given in Fig. 2.

Although there are some common sequence motives, the alignments in the N-terminal parts (residues $1-471$ of FIPV, 1-398 and 652-724 of MHV, 1206 of IBV) must be considered as tentative and may not be significant. In fact, apart from their supposed common origin, there is no real indication that the $N$-terminal domains, which presumably constitute the distal bulbous part of the peplomer, have the same three-dimensional structure. This is underlined by the distribution of the cysteine residues. In the C-terminal segments, corresponding to residue 689-1291 of FIPV, most cysteines are conserved in all three proteins. Their conservation indicates that they are probably involved in disulfide linkages important for the overall structure of the peplomer proteins. In contrast, there is no convincing matching of cysteines in the regions aligned to residues 1471 of FIPV. Moreover, differences in the N-terminal regions account largely for the observed differences in molecular weights of the peplomer apoproteins.

As shown in Fig. 2, potential $N$-glycosylation sites are mainly found in regions of low homology. These sites appear to be particularly abundant in the $N$-terminal part of the protein and the low homology region immediately upstream of the transmembrane anchor. It may be noted that the peplomer protein of MHV contains less glycosylation sites than the peplomer proteins of FIPV and IBV (21 versus 35 and 29, respectively).

The presumptive transmembrane anchor is preceded by a highly conserved amino acid motive, KWPWYVWL, and followed by a peculiar, noncharged region, which is remarkably rich in cysteines (Fig. 3). We can only speculate about the function of these cysteine residues, e.g. acylation or membrane anchoring of the protein by disulfide bridges. However, the clustering of cysteines is clearly a typical feature of the coronavirus peplomer protein.


Fig. 2. Homology and potential glycosylation sites (NXS or NXT, except $X=P r o)$ of peplomer sequences. Regions are considered highly homologous if two sequences are at least $30 \%$ identical.

| FIPV: GTALKYLGTLPPSVKEIAISKWGHFYINGYNFFSTFPIGCISFNLTTGVSGAFWTIAYTS | 468 |
| :---: | :---: |
| MHV : GSISVD- | 391 |
| FIPV: YTEALVQVENTAIKNVTYCNSHINN-IKCSQLTANLNNGFYPVASSEVGFVNKSVVLLPS IBV: --- | 527 260 |
|  | 398 |
| FIPV: FFTYTAVNITID----LGMKLSGYGQPIASTLSNI-TLPMQDNNTDVYCIRSNQFSVYVH <br> IBV: NSVNTTTFTLHN----FTFHNETGANPNPS̄GVQNILTYQTQTAQSGYYNFNFSELSESFV̄Y | 582 315 |
| MHV: RQVDLQLGNSGFLQTANYKIDTAATSCQLHYTLPKNNVTINNHNPSSWNRRYGENDAGVF | 458 |
| FIPV: STCKSSLWDNIFNQDCTDVLEATAVIKTGTCPFSFDKLNNYLTFNKFCLSL <br> IBV: KE--STNFMYGSYHPSC̄NFRLETINNGLWF--------NSLSVSIAYGPLOGGCKQSVF | 633 363 |
| MHV: GKNQHDVVY̌AQQCFTVRSSYCPCAQPDIVSPCTTQTKPKSAFVNVGDHCEGLGVLEDN-- | 516 |
| FIPV: --_------SPVGAN-CKFDVAARTRT--NEQVVRSLYVIYEEGDNIVGVPSDNSGL-H IBV: SGRATCCYAYSYGGPSLCKGVYSGELDL-- $\overline{\text { NFECGLLVYVTKSGGSRIQTATEPPVITRH }}$ | 679 421 |
| MHV: ------CGNADPHKGCICANNSFIGWSHDTCLVNDRCQIFANILLNGINSGTTCSTDLQL | 570 |
| FIPV: DLSVLHLDSCTDYNIYGRTGVGIIRR-T------NS--TLLSGLYYTSLSGDLLGFKNVS IBV: NYNNITL̄NTC̄VDYNIYGRTGOGFITTNVTDSAVSYNYLADA--GLAILDTSGSID $\overline{I F V V O G ~}$ | 730 479 |
| MHV: PNTEVVTGIEVKYDLYGITGOGVFKEVK-- $\overline{\text { A }}$ YYYNSWQTLLYDVN----GNLNGFRDLT | 623 |
| FIPV: D-GVIY-SVTPC-DVSAQAAVIDGAIVGAM--TSIN--SELLGLTHWTTTPNFYYYSIY | 782 |
| IBV: EYĞLTYYKV̄PCEDVNQQFVVSGGKLVGIL--TSRNETGS | 529 |
| MHV : TNK- TY - TIRSCYSGRVSAAFHKDAPEPALLYRNINCSYVFSNNISREENPLNYFDSYLG | 681 |
|  | 827 |
|  | 571 |
| MHV : CVVNADNRT---DEALPNCDLRMGAG-LCVDYSKSRRAHRSVSTGYRLTTFEPYTPMLVN | 737 |
| FIPV: --VQPI--STGNVTIPTNFTISVQVEYMQVYTTPVSIDCARYVCNGNPRCNKLLTQYVSA IBV: --VAPLLNVTENVLIPNSFNLTVTDEYIQTRMDKV̄QINCLQYVCGNSLDCRDLFQQYGPV MHV: DSVQ $\bar{S} V D G L Y E-M Q I P T N F T I G \overline{H E E F I Q T R S P K V T I D C A A F V C G D N T A C R Q \bar{L}} \mathrm{LVEYGSF}$ | 885 631 796 |

MIVLVTCLLLLCSYHTVLSTTNNECIO-VNVTOLAGNENLIRDFLFSNFKEEGSVVVGGY FIPV: MIVLVTCLLLLCSYHTVLSTTNNECIO-VNVTQLAGNENLIRDFLFSNFKEEGSVGGY
 MHV: M--LFVFILFLPSC--LGYIGDFRCIQLVN-SNGANVSAPS----ISTETVEVSQGLGTY Mill
FIPV: YPTEVWYNCSRTARTTAFQYFNNIHAFYFVMEAMENSTGNARGKP-LLFHVHGEPVSVII
 MHV: YVLDRVY-LNATLLLTGY_YPVDGSKFRNLALTGTNSVSLSWFEPPYLNQFNDGIFAKVQ
FIPV: ----SAYRDDVQQRPLLKHG-LVCITKNRHINYEQFTSNQWNSTCTGADRKIPFSVIPTFIPV: --$\stackrel{\infty}{\infty}$ FIPV: DNGTKIYGLEWNDDFVTAYISGRSYHLNINTNWFNNVTLLYSRSSTATWEYSAAYAYQGV 232 IBV: DTTVFVTHCYKYDGCPITGMLQKNF-LRVSAMKNGQ---LFYNLTVSVAKYPTFKSFQCV 161 MHV: TNGNKLIGFWHTDVKPPPICVLKRNFTLNVNADAF---MFHFYQHGGTFY--AYYADKPS 222 FIPV: S--NFTYYKLNNTNGLKTYELCEDYEHCTGYATNVFAPTSGGYIPDGFS--FNNWFLLTN FIPV: - - - 288 $\stackrel{\infty}{\stackrel{\infty}{N}}$
FIPV: SSTFVSGRFVTNQPLLINCLWPVPSFGVAAQEFCFEGAQFSQCNGVSLNNTVDVIRFNLN 348
 MHV: TSAVDCASSYTSEIKCKTQSM-LPSTGV----------YELSGYTVQPVGVVYRRVAN 325 FIPV: FTADVQSGMGATVFSLNTTGGVILEISCYSDTVSESSSYSYGEIPEGITDGPRYCYVLYN
MHV: LPACNIEEWLTARSVPSPLNWERKTFQNC.YFNLSSLLRYVQAESLECNNIDASKVYGRCF ©


$$
\begin{aligned}
& \text { Fig. 3. Alignments of the peplomer sequences of FIPV (strain } \\
& \text { 79-1146), IBV (strain M41) and MHV (strain A59). } \\
& \text { Tentative or arbitrary alignments are indicated by } \\
& \text { extra spacing between sequence lines. Identical re- } \\
& \text { sidues are underlined. Thke cleavage sites in the } \\
& \text { peplomer precursor proteins of MHV and IBV are } \\
& \text { indicated by arrowheads (Cavanagh et al., } 1986 \text {, L. } \\
& \text { Sturman, pers, commun.), the presumptive signal } \\
& \text { peptides and transmembrane anchors by black bars. }
\end{aligned}
$$

## INDICATIONS FOR COILED $\alpha$-HELICES IN THE STALK OF THE PEPLOMER

The three-dimensional structure of the coronaviral peplomer has not yet been determined. Therefore, no data are available on how monomers interact to form a stable multimer. Also, the molecular basis for the typical elongated appearance of the peplomers is unknown. In other elongated protein molecules, like the haemagglutinin of influenza virus and reovirus, long $\alpha$-helices of the monomers interlock in a coiled coil (Wilson et al., 1981; Bassel-Duby et al., 1985). As reviewed by Cohen and Parry (1986) this structure stabilizes the multimer and imparts the elongated character to the molecule. Indicative for $\alpha$-helices forming a coiled coil is a seven-residue 'heptad' repeat in the amino acid sequence ( $\mathrm{a}, \mathrm{b}, \mathrm{c}, \mathrm{d}, \mathrm{e}, \mathrm{f}, \mathrm{g}$ ), in which the residues in the positions a and $d$ generally have an apolar character. In the $\alpha$-helix, these residues are aligned, resulting in a continuous hydrophobic stretch along the axis. Such hydrophobic 'backbones' form the interface between interlocking helices.

Conceivably, a similar structure could also be present in the peplomers of coronaviruses. Systematic examination of the peplomer sequences revealed the presence of two heptad repeats in the C-terminal region of E2 (Fig. 4). The repetitious character of these regions is also obvious in the Diagon plot (Fig. 1). One of the repeats is located immediately upstream of the transmembrane anchor. The presence of this repeat (residues 1328-1380, 1055-1080 and 1214-1251 of the peplomer proteins of FIPV, IBV and MHV, respectively) is well conserved, in spite of the low degree of amino acid conservation in this region.

The other heptad repeat is even longer and located further upstream (Fig. 4; residues 1067-1149, 796-864 and 972-1041 of the E2 proteins of FIPV, IBV and MHV , respectively). In Fig. 5, this repeat is visualized by listing the sequence in alternating rows of four and three residues (a 'helical net'). As indicated in Fig. 4, both heptad repeats coincide in all three proteins with regions devoid of helix-breaking proline residues.


Fig. 4. Indications for long $\alpha$-helices in the peplomer protein.

The two heptad repeats are indicative for the presence of two $\alpha-$ helices in the peplomer protein that are capable to interlock with other helices. For the major repeat a helix with a length of at least 10 nm can be predicted. This helix would be longer than the longest helix of influenza virus haemagglutinin ( 7.5 nm ) and would extend over about half the length of the peplomer ( $17-20 \mathrm{~nm}$ ). It is tempting to speculate that, as in the case of the haemagglutinins, such an $\alpha$-helix interlocks with the $\alpha$-helix(-ces) of (an)other mononomer(s) to form a coiled coil. This structure could stabilize the multimer and account for the characteristic elongated appearance of the stalk of the coronavirus peplomer.

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HEPTADS:


FIPV:


IBV:


MHV :


Fig. 5. Amino acid sequences of peplomer proteins drawn in a helical net, assuming one heptad per two turns. The residues 1060-1192, 775-907 and 951-1084 of the peplomers of FIPV, IBV and MHV, respectively, are listed vertically in alternating rows of three and four residues. Hydrophobic residues are encircled. The hatched bars indicate continuous hydrophobic regions, which may interact with other $\alpha$-helices.

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