VACCINIA VIRUS HEMAGGLUTININ
A Novel Member of the Immunoglobulin Superfamily

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Vaccinia virus hemagglutinin (VVHA) has long been recognized as a lipid-linked glycoprotein that can not only agglutinate chicken as well as human erythrocytes, but can also play an important role in the intercellular dissemination of the virus (1). The binding element on the surface of the erythrocytes and the mechanism responsible for the virus spreading remain elusive for >40 yr. Being a nonvion protein unnecessary for the virus replication and expressed on the surface of the infected cells, VVHA has been used in recent years as a selection marker in constructing the recombinant vaccinia virus (2), which has the potential to develop polyvalent vaccines. It is of vital importance to elucidate its structure and its influence on the cytopathology and virulence of the virus. The VVHA gene from a nonvaccine strain (WR strain) of vaccinia virus was mapped within the Sal I/Hind III region at the right edge of the Hind III A fragment of the genome and was sequenced recently (3). For safety’s sake, the better choice of vaccinia virus in developing vaccines for human use should be a less virulent vaccine strain (4). To provide a molecular basis for understanding the function of VVHA and its interaction with the erythrocytes and the infected cells, we now sequenced and analyzed the VVHA gene from the Chinese vaccine strain (Tiantan Strain) of vaccinia virus.

Materials and Methods

Bacterial Strains, Bacteriophages, and Plasmids. Escherichia coli K12 JM103 was originally obtained from Boehringer Mannheim Biochemicals, Penzberg, FRG, and was grown on M9 minimal-medium plates, as recommended by the supplier. Bacteriophage M13 mp18/ mpl9 RF and plasmids pAT153 and pUC19 were purchased from Pharmacia Fine Chemicals, Uppsala, Sweden. Plasmid pV630, containing the Sal I G fragment of the Tiantan strain of vaccinia virus, was constructed in our laboratory. The sequence coding for VVHA was confirmed by the restriction map compared with that of the WR strain, and also by the loss of HA trait in the recombinant viruses with a galactosidase gene inserted into the region (Hao Yuwen, manuscript in preparation).

Enzymes and Chemicals. Restriction enzymes were mainly purchased from New England Biolabs, Beverly, MA. Deoxyribonucleoside triphosphates (dNTPs, ddNTPs, α-[32P]dATP, and α-[35S]dATP-α-S) were obtained from Amersham International, Amersham, UK. A modified T7 DNA polymerase (Sequenase) and all the other chemicals used in nucleotide
sequencing were from United States Biochemical Corp., Cleveland, OH. All enzymes and chemicals were used according to the manufacturer's recommendations.

**Nucleotide Sequencing.** The 1.8-kb SalI/Hind III fragment of plasmid pV630 was subcloned into plasmid pUC19. Overlapping fragments were then taken from the subclone and inserted into M13 mp18 and mp19 in both orientations. The nucleotide sequence was determined on both strands by the M13 dideoxy chain termination method using a modified T7 DNA polymerase (5). The nucleotide and peptide sequences were analyzed on an IBM PS-2 computer with the CalTech software package developed by Dr. Alan Goldin from the California Institute of Technology, Pasadena, CA, and kindly provided by the Molecular Biology Computer Research Resources (MBCRR), Boston, MA. The routine to perform dot-matrix analysis was written by G. Gutman and B. Ward from the University of California, Irvine, CA. The FASTA program (6), developed and kindly provided by Dr. W. R. Pearson from the University of Virginia, Charlottesville, VA, was used to search the National Biomedical Research Foundation (NBRF) protein database release 12.0 (7) obtained from MBCRR.

**Results**

The VVHA gene from the Tiantan strain is located at the right edge of the Sal I G fragment of the virus genome. The 1,458-bp nucleotide sequence starting from the right terminus of the Sal I G region reveals a single open reading frame with 315 amino acids (Fig. 1). Of them, 11 nucleotides and eight deduced amino acids were found to be different from those in the WR strain (3).

A search in the NBRF protein sequence database revealed proteins belonging to the Ig superfamily with similarities to VVHA. In addition to a 22% overall identity between the first 110 residues of VVHA and the human Ig CH chain V-I region, consensus residues were found clustered around the two conserved cystein residues. Sequence alignment was then extended to other members of the Ig superfamily (Fig. 2). Most of the residues that are conserved among the IgV domains (8) are also relatively invariant in the VVHA molecule. The best example is that although there is only one tryptophan in the deduced 315-amino acid VVHA molecule, its position is very similar to that of the conserved tryptophan in the V region (9). Moreover, the size of the similar region (100 residues) and the distance between the two cysteins (70 residues) resemble those of the Ig V region (8). It is reasonable to suggest that VVHA contains an Ig-like domain of ~100 amino acids at its NH2 terminus, with a three-dimensional structure characteristic of the Ig-like fold (9).

The most intriguing finding in the self-comparison of the VVHA sequence is that two tandem repeating units exist head to tail in the middle of the VVHA gene and deduced peptide (Fig. 3, A and B). They were located in a region from 170 to 240 residues at the amino acid level, just after the Ig-like domain. These two units share significant sequence homology with each other but show little similarity to proteins belonging to the Ig superfamily. They might possibly have evolved from the duplication of a gene fragment unrelated to the Ig superfamily. It is not known whether this region has a useful viral function.

The deduced protein sequence (Fig. 1) and its hydrophobicity plot (Fig. 4 A) demonstrate that VVHA should be a typical transmembrane glycoprotein (Fig. 4 B). The first 16 amino acids of VVHA comprise a hydrophobic region rich in leucine.

1 The following sequences, each with an entry name given in parentheses, are fully referenced in release 12.0 of the NBRF protein database (7): human Ig CH chain V-I region (LIHUNG), human Ig H chain V-II region (MHHUMC), rabbit poly-Ig receptor (QRBBG), human CD4 (RWHUT4), and human TCR β chain (RWMSCS).
which is probably a signal peptide to be cleaved off the mature protein. At the COOH
terminus, another hydrophobic region is followed by a hydrophilic tail rich in basic
residues. This unit is most likely the transmembrane-cytoplasmic portion of VVHA.
Between the two hydrophobic regions are one Ig-like domain and two tandem repeating
units.

Figure 1. Nucleotide and deduced amino acid sequence starting from the Sal I G fragment
of the genome of the Ti-ant strain of vaccinia virus. The putative signal sequence and the probable transmembrane portion of the molecule are indicated, respectively, by dashes and asterisks below the amino acid sequence. Five potential N-linked glycosylation sites are also underlined. These sequence data have been submitted to the EMBL/GenBank Data Libraries.
Discussion

The concept of an Ig-like domain as the primordial, yet versatile structure involved in intercellular recognition in higher eukaryotes has been strongly reinforced by the sequences of many newly identified members of the Ig superfamily (10). The homophilic adhesion of the neural cell adhesion molecule (N-CAM) and the binding of CD2 to LFA-3 may represent the basic model for the interaction within the superfamily. Considering that the VVHA has an Ig-like domain exposed on the cell surface and that VVHA is responsible for the hemagglutination, the intercellular spreading, and perhaps the release of the virus (3), we believe that VVHA will be another case in support of the above model. Among the superfamily members, LFA-3 and rat OX-45, whose equivalent in humans is called Blast-1 (11), were found to be...
expressed on the surface of erythrocytes. The LFA-3 antigen was also shown to mediate adhesion between T cells and erythrocytes by interacting with CD2. It would be of interest to see whether VVHA would trigger the virus-induced hemagglutination by recognizing an as yet unidentified Ig-related ligand on the erythrocytes.

It is generally accepted that all members of the Ig superfamily share a common ancestry (10). Comparisons of the amino acid sequence between VVHA and other superfamily members demonstrate that the Ig-like domain in VVHA is structurally more similar to the Ig V domain, and that the sequence flanking the Ig-like domain is perhaps dissimilar to proteins belonging to the Ig superfamily. This prompts us to consider that vaccinia virus had captured an exon encoding an Ig V domain from the eucaryotic cell when interacting with the host immune system and converted it through evolution to the VVHA molecule of its own. It is noteworthy that monoclonal autoantibodies against intermediate filaments or Thy-1.2 antigen produced by clones established after immunization with lysates from cells infected by vaccinia virus were shown to crossreact with VVHA (12). Elucidation of the influence of VVHA on the cytopathogenesis and virulence of the vaccinia virus requires further study of the potential molecular mimicry between VVHA and other members of the Ig superfamily, including the myelin-associated glycoprotein MAG and the major glycoprotein of peripheral myelin P0 found on neural tissues (10).

Summary

Striking similarities between vaccinia virus hemagglutinin (VVHA) and proteins belonging to the Ig superfamily clearly indicate that VVHA, a 315-amino acid glycoprotein expressed on the surface of the infected cells, is a novel viral protein that can be added to the expanding list of the Ig superfamily. Its deduced amino acid sequence contains one Ig-like domain at the NH2 terminus, followed by two tandem
repeating units and a hydrophobic region, suggestive of membrane spanning. The results offer an opportunity for the further study of the probable evolutionary and possible functional relationship between VVHA and other members of the Ig superfamily. Our observation, together with a recent finding that human CMV possibly encodes a protein similar to the MHC class I antigens (13), provides evidence supporting the fact that the viral capture of cellular Ig-related genes is more common than expected in vaccinia and other viruses, and that the usage of an Ig-like domain as recognition signals might be extended from higher animals to animal viruses.

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