Bioinformatic analysis and characteristics of glycoprotein C encoded by the newly identified UL44 gene of duck plague virus

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ABSTRACT. Glycoprotein C is one of the duck plague virus (DPV) glycoproteins and is encoded by the DPV UL44 gene. DPV glycoprotein C (DPV-gC) comprises 431 amino acids with a putative molecular mass of 47.35 kDa. Sequence analysis indicated that the protein possesses typical characteristics of type-I membrane glycoproteins, containing an N-terminal signal sequence, an external domain, a C-terminal membrane anchor region, and a short cytoplasmic domain. Comparisons of 22 alphaherpesvirus-gC protein sequences revealed eight conservative Cys-residue sites, which may play a crucial role in the biological functions and structural stabilization of the DPV-gC protein. Estimates of potential antigenic epitopes and secondary structure identified four B cell dominant epitopes, which are located at amino acids 68-71, 87-91, 369-352, and 372-374. A model for the structure of DPV-gC was derived by associating its predicted secondary and three-dimensional
structures. In conclusion, these results will provide a basis for further functional studies of DPV-gC, establishing novel clinical diagnoses of DPV, and in the development of a new DPV vaccine.

Key words: Bioinformatics; Structure; Function; Glycoprotein C; Duck plague virus

INTRODUCTION

Duck plague virus (DPV), which is the causative agent of duck plague has been classified in the subfamily Alphaherpesvirinae of Herpesviridae based on the report of the ninth International Committee on Taxonomy of Viruses (ICTV), but has not yet been grouped into any genus (Davison et al., 2009; King et al., 2011). DPV has a typical herpes virion, consisting of a linear double-stranded DNA core, an icosahedral capsid, a tegument, and an envelope containing viral glycoprotein spikes on its surface (Pellett and Roizman, 2007; Sandhu and Shawky, 2008). In duck breeding areas of the world where the disease has been reported, DPV has caused significant economic losses in both domestic and wild waterfowl due to high mortality and low egg production rates (Sandhu and Shawky, 2008).

As a multifunctional glycoprotein in Alphaherpesvirinae, glycoprotein C is involved in viral attachment, release, virulence, and immune evasion, among other functions (Schreurs et al., 1988; Friedman et al., 1996; Chowdhury et al., 2000; Jarosinski and Osterrieder, 2010). Being situated on the envelope surface of mature virus particles, glycoprotein C contains many antigen determinants, and can adequately induce the immune response (Ben-Porat et al., 1986; Guo et al., 1989; Tewari et al., 1995). In the DPV genome, the UL44 gene, which is conserved with the UL44 gene of other alphaherpesviruses (Allen and Coogle, 1988; Fitzpatrick et al., 1989; Nicolson and Onions, 1990), was identified and sequenced in our laboratory. The complete nucleotide sequence of the UL44 gene from the DPV CHv strain has been submitted in the GenBank database and was assigned the accession number EU076811. We named the corresponding gene as DPV-UL44 and the protein as DPV-gC.

The objective of the present study was to provide information about the structure and functions of DPV-gC. This study may provide insight for further research related to DPV-gC, and provide a fundamental basis for establishing new types of clinical diagnoses of DPV and for the development of a new DPV vaccine.

MATERIAL AND METHODS

Prediction of the physicochemical properties of DPV-gC

The DNAStar 7.1 program was used to analyze components of the DPV-gC sequence. The deduced amino acid sequence was analyzed with online prediction tools. The signal peptide, phosphorylation site, transmembrane region, motifs, hydrophobicity, subcellular localization, and protein-protein interaction sites were analyzed with the programs SignalP 4.0 (http://www.cbs.dtu.dk/services/SignalP; Petersen et al., 2011), NetPhos 2.0 (http://www.cbs.dtu.dk/services/NetPhos/; Blom et al., 1999), TMHMM (http://genome.cbs.dtu.dk/services/TMHMM/), PSORT (http://www.psort.org/psortb/results.pl; Yu et al., 2010), and PredictPro-
Estimation of potential antigenic epitopes

To estimate the potential antigenic epitopes of DPV-gC, the amino acid sequence of the DPV-gC was analyzed using online prediction tools and commensurate software. Antigenic peptide prediction, hydrophilicity estimation, surface accessibility, and flexibility assumption were performed using prediction tools. An online web server (http://immunax.dfci.harvard.edu/Tools/antigenic.pl) provided a pathway to predict sequences of peptides within a protein that were likely to be antigenic by eliciting an antibody response. The SIB Bioinformatics Resource Portal (http://us.expasy.org/tools/protscale.html) was used to predict the hydrophilicity of DPV-gC, which was based on the method of Kyte & Doolittle (Kyte and Doolittle, 1982). The PredictProtein program (http://www.predictprotein.org/submit.php) was used to predict the surface accessibility of DPV-gC. The flexibility was DPV-gC was determined with the Protean procedure in DNAStar 7.1, based on methods of Karplus-Schulz (Karplus and Schulz, 1985).

Comparison of the DPV-gC sequence with 21 published gC protein sequences and phylogenetic tree construction

We obtained information of the 21 published gC amino acid sequences of other alphaherpesviruses from the National Center for Biotechnology Information (NCBI) database (Table 1). Comparison and analysis of the conserved regions of DPV-gC with the 21 published alphaherpesvirus-gC amino acid sequences was performed using ClustalX 1.83 or DNAStar 7.1. The phylogenetic tree was constructed using the neighbor-joining method in PHYLIP 3.69. The reliability of each tree branch was tested by bootstrap analysis of 100 data samples and the trees were constructed using TreeView 1.66.

Table 1. Information about the amino acid sequences of the DPV-gC and the 21 referenced gC.

<table>
<thead>
<tr>
<th>Genus</th>
<th>Virus name (abbreviation)</th>
<th>GenBank Accession No.</th>
<th>CDS Sequence length (bp)</th>
<th>Protein ID</th>
<th>Protein length (aa)</th>
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Secondary and tertiary structure prediction

The presumed alpha helix, extended strand, and random coil of DPV-gC were obtained from the Network Protein Sequence Alignment (NPSA) (http://npsa-pbil.ibcp.fr/cgi-bin/secpre2_mlr.pl; Combet, et al., 2000). The prediction of the 3D structure of the DPV-gC protein was performed through the SWISS-MODEL server (http://www.cbs.dtu.dk/services/CPHmodels/; Kelley and Sternberg, 2009).

RESULTS

Physico-chemical properties of DPV-gC

The DPV UL44 gene was originally obtained by sequencing the DPV genomic library that was constructed in our laboratory. The predicted open reading frame of the DPV UL44 gene was expected to encode a protein comprising 431 amino acids with a putative molecular mass of 47.35 kDa and an isoelectric point (pI) of 5.29. DNAStar7.1 analysis revealed that the polypeptide encoded by the DPV UL44 gene was composed of 33.64% hydrophobic amino acids, 29.47% hydrophilic amino acids, 8.12% strongly basic amino acids, and 10.44% strongly acidic amino acids.

Furthermore, using different online web servers, we obtained more detailed information about the structure and function of DPV-gC. First, the N-terminus of the putative protein containing four potential N-linked glycosylation sites (at aa residues 52, 58, 163, and 279) and three N-myristoylation sites (at aa residues 51, 90, and 406) could be clearly identified. Second, one cAMP- and cGMP-dependent protein kinase phosphorylation site (at aa residue 166), nine protein kinase C phosphorylation sites (at aa residues 32, 162, 165, 176, 213, 230, 236, 264, and 365), and five casein kinase II phosphorylation site (at aa residues 77, 105, 142, 239, and 254) were identified in DPV-gC. Furthermore, amino acid sequence analysis revealed that DPV-gC is a type-I membrane glycoprotein that can be divided into three kinds of distinct functional domains: the extracellular domain, the hydrophobic transmembrane domain, and the cytoplasmic domain.

Estimated potential antigenic epitopes

The prediction tools of online web servers indicated that the DPV-gC amino acid sequence contains 20 main antigenic determinants, which were situated in amino acids 26-39, 59-78, 85-91, 94-101, 132-141, 148-158, 171-197, 199-207, 216-222, 227-233, 242-254, 262-267, 279-296, 298-316, 332-343, 351-357, 359-370, 372-386, 395-406, and 408-423. Hydrophilicity prediction revealed that the highest points of DPV-gC’s hydrophilicity were between amino acids 20-28, 29-59, 66-71, 80-97, 104-133, 143-149, 157-171, 194-203, 209-215, 217-219, 221-226, 231-243, 257-263, 268-278, 300-302, 321-331, 341-344, 346-377, 382-393, and 421-427. Due to their exposure on the virion or cell membrane surface, these parts of the protein are likely to contain the major antigenic epitopes of DPV-gC. Results obtained from the surface accessibility website indicated that the DPV-gC amino acid sequence contains 21 main surface accessibility domains, which are situated in amino acids 29-34, 50-60, 68-82, 86-93, 104-124, 127-133, 142-146, 155-166, 170-174, 181-185, 206-213, 220-232, 234-243, 256-262, 270-280,
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Comparing the 22 alphaherpesvirus-gC protein sequences and constructing the phylogenetic tree

Comparison of the 22 alphaherpesvirus-gC protein sequences by DNAStar7.1 indicated that there were eight conserved Cys-residue sites. Combined with the results of signal peptide and transmembrane domain prediction of the gC protein, statistical results of the interval among the conserved Cys-residue sites are shown in Table 2. Except for ILTV and PsHV-1 from the genus Iltovirus, the biggest difference in sequence length was observed in the interval between the signal peptide and the conserved Cys-residue site 1, followed by the length of the cytoplasmic domain and the interval between the conserved Cys-residue sites 2 and 3.

<table>
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<tr>
<th>Virus name (abbreviation)</th>
<th>Signal peptides</th>
<th>C1-C2</th>
<th>C2-C3</th>
<th>C3-C4</th>
<th>C4-C5</th>
<th>C5-C6</th>
<th>C6-C7</th>
<th>C7-C8</th>
<th>Transmembrane regions</th>
<th>Intracellular region</th>
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<td>3</td>
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<td>160</td>
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<td>3</td>
<td>27</td>
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<td>16</td>
<td>145</td>
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The amino acid sequence alignment based on truncated gC protein sequences (for which the residues before the conserved Cys-residue site 1 were deleted) showed that the Cys-
residue sites 1, 2, 5, and 8, along with three Pro-residue, three Trp-residue, two Tyr-residue,
and one Gly-residue were all completely conserved in alphaherpesviruses. In particular, there
was a conservative module (Y*C***G) around the conserved Cys-residue site 8 in alphaher-
pesviruses. Moreover, there were also conservative modules found around Cys-residue con-
servative sites 3, 4, 5, 6, and 7, except for ILTV and PsHV-1 from the genus Iltovirus.

The phylogenetic tree was constructed by comparing the truncated DPV-gC amino
acid sequence with 21 truncated gC amino acid reference sequences of the other alphaher-
pesviruses. As shown in Figure 1, the 22 gC amino acid sequences were divided into four
branches, which were respectively termed Mardivirus (which contains HVT, MDV-1, and
MDV-2), Iltovirus (which contains ILTV and PsHV-1), Simplex virus (which contains CeHV-
1, CeHV-2, HVP-2, HSV-1, HSV-2, and SaHV-1), and Varicello virus (which contains BoHV-
1, BoHV-5, CeHV-9, EHV-1, EHV-4, EHV-8, EHV-9, FHV-1, PRV, and VZV). Further obser-
vation of the phylogenetic branching showed that DPV-gC was closely related to sequences in
the Mardivirus genus. Notably, however, the DPV sequence clustered within a monophyletic
clade, which was independent of the others.

Figure 1. Phylogenetic analysis of the truncated DPV-gC amino acid sequence and 21 truncated gC amino acid
reference sequences of the other alphaherpesviruses was aligned using ClustalX 1.83. Phylogenetic tree was
constructed by using Phylip 3.69 and drawn using TreeView 1.66.

Secondary and tertiary structure prediction

The prediction of DPV-gC’s secondary structure is shown in Figure 2. The result in-
dicated that 9.51, 32.95, and 57.54% of the DPV-gC protein consists of an alpha helix (H), an
extended strand (E), and a random coil (C), respectively. The prediction of the 3D structure of the DPV-gC protein was determined with the SWISS-MODEL database (Figure 3). Three hundred sixty-three residues (84% of DPV-gC) were modeled with 99.8% confidence by the single highest scoring template. These 363 residues clearly represented the extracellular domain truncated signal peptide of DPV-gC. By associating the predicted secondary and 3D structures and the disulfide bond formed by the eight Cys-residues, we were able to predict a model for the structure of DPV-gC (Figure 4).

Figure 2. Secondary structure presumption of the DPV-gC. Blue represents alpha helix, red represents extended strand, and purple represents random coil.

Figure 3. Prediction of DPV-gC’s tertiary structure.
DISCUSSION

The aim of the present study was to provide predicted information about the structure and functions of DPV-gC, which is encoded by the newly identified DPV UL44 gene, using bioinformatic software and tools.

Results of DPV-gC sequence analysis revealed that DPV-gC has the standard membrane protein arrangement of an N-terminal signal sequence, an external domain, a C-terminal membrane anchor region, and a short cytoplasmic domain, and that the protein may show good immunogenicity and likely undergoes chemical modifications such as glycosylation and phosphorylation. Therefore, these results indicate that DPV-gC is a very relevant protein for the development of new antibodies and immunoassays in the clinical diagnosis of DPV.

Detailed molecular understanding plays a critical role in the improvement of vaccines. It is therefore important to determine which surface features on the DPV are responsible for inducing protective or enhancing immune responses. In this study, we focused on characterizing the B cell epitopes of DPV-gC by adopting bioinformatic approaches. The sequences at amino acids 68-71, 87-91, 369-352, and 372-374 of DPV-gC were predicted as the more prevalent dominant antigen epitopes based on multiple parameters and independent analyses in different software. The predicted dominant antigen epitopes can be used for the development of a vaccine against DPV. Further experimental studies will be performed to determine the immunogenicity and protection effect of DPV-gC peptides and to identify specific vaccine candidates.

Figure 4. Model for the structure of the DPV-gC.
Comparison of the 22 alphaherpesvirus-gC protein sequences revealed that amino-terminal sequences have undergone more extensive evolutionary divergence than those at the C-terminal half of the molecule. This appears to be a universal phenomenon for gC-like glycoproteins among all alphaherpesviruses (Allen and Coogle, 1988; Fitzpatrick et al., 1989; Nicolson and Onions, 1990). Notably, eight Cys-residue sites of the DPV-gC were found to be conserved, suggesting that these amino acids may be disulfide bond-linked and that they are likely to be important for maintaining the proper structure and functions of the protein. A structural model of HSV-1 gC demonstrated that the disulfide bond arrangement is cysteine 1 to cysteine 2, cysteine 3 to cysteine 4, cysteine 5 to cysteine 8, and cysteine 6 to cysteine 7 (Rux et al., 1996). The first disulfide bonds and basic and hydrophobic residues of HSV-1 gC constitute a major heparan sulfate-binding domain (Mardberg et al., 2001); the first two disulfide bonds produce loops between adjacent cysteines within the C3b binding domain (Hung et al., 1992; Rux et al., 1996), and the last two disulfide bonds result in an extended loop which may stabilize the carboxy terminus (Fitzpatrick et al., 1989; Rux et al., 1996).

The evolutionary relationship among the alphaherpesvirus-gC protein sequences was analyzed by constructing a phylogenetic tree based on the truncated DPV-gC and the truncated gC of 21 referenced sequences of the other alphaherpesviruses. The results revealed that DPV was clustered within a monophyletic clade, despite its close evolutionary relationship with the genus *Mardivirus*, which contains HVT, MDV-1, and MDV-2. These results should be useful for DPV classification studies and indicate that DPV should be placed in a separate, single genus within the Alphaherpesvirinae.

Extended strand and random coils are relatively more non-cohesive flexibility structures, which always contain B cell dominant epitopes. The alpha-helix and beta-fold of the protein have higher chemical bonding energies, which can firmly maintain proteinic higher structures, but are difficult to bind antibody, and because they are usually located at the protein interior, they seldom become B cell epitopes. In contrast, the extended strand and random coil of the protein are more loose in texture, which results in easy retortion, and can therefore stretch out of the proteinic surface and bind antibody, enabling them to become B cell epitopes (Barlow et al., 1986; Apostolopoulos et al., 2002). Therefore, secondary structure predictions of proteins are generally applied to definite B cell epitopes as an assistance medium. Combining the results of antigenic epitopes estimation and secondary structure prediction, four B cell dominant epitopes could be initially defined in DPV-gC, which were located at amino acids 68-71, 87-91, 369-352, and 372-374.

The function of a protein depends on its tertiary structure. If this is disrupted, the protein is denatured, and it loses its activity. For many proteins and peptides, disulfide bridges are a prerequisite for their proper biological function. The eight conservative Cys-residue sites in DPV-gC, which were clustered around several other conserved residues, and the general pattern of relatively low N-terminal conservation and higher C-terminal conservation of the gC-like glycoproteins of all alphaherpesviruses, indicated that these glycoproteins have retained a common carboxy-terminal, disulfide bond-based framework in spite of their amino-terminal heterogeneity (Fitzpatrick et al., 1989).

In conclusion, in order to characterize the role of glycoprotein C encoded by the newly identified UL44 gene of duck plague virus, bioinformatic and molecular approaches were used to predict and analyze its structure and functions. Together, these results provide necessary data for further DPV-gC research, the establishment of new types of clinical diagnoses of DPV, and for the development of a new DPV vaccine.
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Bioinformatic analysis and characteristics of DPV-gC


