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Exploiting 2A peptides to elicit potent neutralizing antibodies by a multi-subunit herpesvirus glycoprotein complex

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1. Introduction

Herpesviruses include a number of ubiquitous and highly-adapted human pathogens that can cause severe illnesses in individuals with impaired immune system such as transplant recipients or AIDS patients or in congenitally infected fetuses (Arvin et al., 2007). While licensed vaccines are available for varicella-zoster virus (VZV), vaccines candidates for other human herpesviruses including herpes-simplex viruses (HSV), human cytomegalovirus (HCMV), Epstein-Barr virus (EBV), and Kaposi sarcoma-associated herpesvirus (KSHV) remain elusive (Arvin et al., 2007; Stratton et al., 2000). Considerable efforts to develop herpesvirus vaccines have been made using approaches based on selected humoral or cellular immunodominant antigens as safer and more cost-effective alternatives to live-attenuated, inactivated, or replication-defective herpesvirus vaccines. These subunit vaccine approaches using plasmids, viral vectors, purified protein, or virus-like particles showed promising results in different animal models and clinical trials (Cohen, 2015; Dasgupta et al., 2009; Johnston et al., 2016; Pass et al., 2009; Schleiss, 2008; Schleiss, 2016).

Eliciting neutralizing antibodies (NAb) that interfere with glycoprotein complex-mediated virus entry into host cells is thought to be important for a vaccine formulation to prevent or control herpesvirus infection (Nelson et al., 2017; Plotkin, 2013). While the essential and highly-conserved herpesvirus envelope glycoprotein complexes composed of gB and gH/gL are principal targets for NAb, additional glycoprotein complexes or accessory glycoproteins associated with gH/gL can represent critical immune targets that contribute to the stimulation of herpesvirus neutralizing activity (Heldwein, 2016; Macagno et al., 2010; Sathiyaamoorthy et al., 2017; Vanarsdall and Johnson, 2012). Over the past years it has been discovered for HCMV that NAb blocking infection of many biologically relevant host cells recognize in majority three accessory glycoproteins called UL128, UL130, and UL131A that form a pentamer complex (PC) with gH/gL. (Fouts et al., 2012; Macagno et al., 2010; Wang and Shenk, 2005). While this complex is dispensable for HCMV infection of fibroblasts (FB), it is required for efficient infection of epithelial cells (EC), endothelial cells, and other cells thought to be important for HCMV dissemination and transmission (Hahn et al., 2004; Sinzger et al., 2008; Vanarsdall and Johnson, 2012; Wang and

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Neutralizing antibodies (NAb) interfering with glycoprotein complex-mediated virus entry into host cells are thought to contribute to the protection against herpesvirus infection. However, using herpesvirus glycoprotein complexes as vaccine antigens can be complicated by the necessity of expressing multiple subunits simultaneously to allow efficient complex assembly and formation of conformational NAb epitopes. By using a novel bacterial artificial chromosome (BAC) clone of the clinically deployable Modified Vaccinia Ankara (MVA) vector and exploiting ribosomal skipping mediated by 2A peptides, MVA vectors were generated that expressed self-processing subunits of the human cytomegalovirus (HCMV) pentamer complex (PC) composed of gH, gL, UL128, UL130, and UL131A. These MVA vectors expressed 2A-linked HCMV PC subunits that were efficiently cleaved and transported to the cell surface as protein complexes forming conformational neutralizing epitopes. In addition, vaccination of mice by only two immunizations with these MVA vectors resulted in potent HCMV NAb responses that remained stable over a period of at least six months. This method of eliciting NAb by 2A-linked, self-processing HCMV PC subunits could contribute to develop a HCMV vaccine candidate and may serve as a template to facilitate the development of subunit vaccine strategies against other herpesviruses.

1. Introduction
Shenk, 2005). In contrast to NAb interfering with the essential entry function of gB and gH/gL complexes, NAb interfering with PC-mediated entry are unable to block FB infection, though they are substantially more potent than NAb targeting gB or gH/gL epitopes in preventing infection of non-FB cell types such as EC (Chiuppesi et al., 2015; Macagno et al., 2010). These results suggest that the PC could be an important vaccine component to prevent or control HCMV infection.

Since NAb targeting the HCMV PC recognize mainly quaternary conformational epitopes that are formed effectively only upon assembly of UL128/130/131A with gH/gL, vaccine-mediated induction of HCMV NAb is promoted by simultaneous expression of all five PC subunits (Chandramouli et al., 2017; Chiuppesi et al., 2017a; Chiuppesi et al., 2017b; Ciferri et al., 2015a; Ciferri et al., 2015b; Macagno et al., 2010; Wussow et al., 2014; Wussow et al., 2013). Ribosomal skipping mediated by 2A peptides of picornaviruses is a widely used mechanism to express multiple proteins via a single transcript because of the relative small size of the 2A peptides (18–22 amino acids) and the potential stoichiometric expression of the co-expressed proteins (de Felipe, 2004; Kim et al., 2011; Szymczak et al., 2004). By using a novel bacterial artificial chromosome (BAC) clone of the well-characterized and clinically-deployable Modified Vaccinia Ankara (MVA) vector (Cottingham et al., 2008; Gilbert, 2013; Verheus et al., 2012), the use of 2A-mediated ribosomal skipping was exploited to stimulate NAb by self-processing subunits of the HCMV PC. The results indicate that recombinant MVA vectors either expressing 2A-linked polyproteins composed of all five PC subunits or co-expressing 2A-linked polyproteins composed of UL128/130/131A and gH/gL stimulate robust HCMV NAb responses in mice. This method of eliciting HCMV NAb by 2A-linked polyribonucleotide expression constructs could contribute to develop a HCMV vaccine candidate and may serve as template to facilitate the development of vaccine approaches based on multi-subunit glycoprotein complexes of other herpesviruses.

2. Materials and methods

2.1. Cells and viruses

Baby hamster kidney (BHK-21) cells, ARPE-19, MRC-5 (American Type Culture Collection (ATCC)) and chicken embryo fibroblasts (CEF: Charles River) were maintained by standard procedures. MVA was propagated in BHK-21 and CEF cells as described previously (Wussow et al., 2014). HCMV and MVA virus stocks were prepared following virus propagation in ARPE-19 or BHK-21 cells, respectively, and titrated as described (Chiuppesi et al., 2017a; Chiuppesi et al., 2015; Wussow et al., 2014). HCMV TB40/E expressing GFP was derived from TB40/Ewt-GFP BAC DNA (O’Connor and Murphy, 2012). MVA 1974/NIH clone 1 was kindly provided by Dr. Bernard Moss (NIAID) (Mayr TB40/Ewt-GFP BAC DNA (O’Connor and Murphy, 2012). MVA 1974/ NIH clone 1 was kindly provided by Dr. Bernard Moss (NIAID) (Mayr and Malicki, 1966). The construction of MVA-PC has been described previously (Wussow et al., 2014).

2.2. Plasmids

Transfer plasmids for generating a novel MVA BAC, termed MVABAC-TK (Fig. 1), or MVABAC-TK-derived recombinants with 2A-linked HCMV PC subunits were constructed by standard molecular biology cloning techniques. For generating MVABAC-TK, a transfer vector was generated in which pBlObac11 sequences (New England Biolabs) and a GFP expression cassette with vaccinia P11 promoter were inserted into the Thymidine kinase (TK) gene locus (Fig. 1). MVA TK homology flanks were derived by PCR from MVA 1974/NIH clone 1 and corresponded to base pairs 69313–70000 and 70001–70703 (MVA Acambis, Accession Nr. AF603355.1). The GFP marker with P11 promoter was derived by PCR from plasmid pL7W3 (Wyatt et al., 2009). A unique AvrII restriction site was introduced between the ends of the TK homology flanks to allow linearization of the entire transfer construct (Fig. 1). Transfer constructs for inserting 2A-linked HCMV PC subunits into MVABAC-TK by En passant mutagenesis were generated by cloning codon-optimized and P2A-linked UL128/130/131A or gH/gL subunit subset gene sequences between the vaccinia modified H5 early/late promoter (mH5) and vaccinia transcription termination signal (TTTTAT) of pGEM-T-mH5 (Wussow et al., 2013). A kanamycin expression cassette with adjacent I-SceI homing endonuclease restriction sites and flanking 50 bp gene duplication was subsequently introduced into the pGEM-T-mH5-cloned HCMV PC subunit genes sequences (Tischer et al., 2010; Wussow et al., 2013). All HCMV PC subunit gene sequences were based on HCMV strain TB40/E (TB40/E-BAC; Accession Nr. EF999921). Codon-optimized and P2A-linked PC gene sequences of HCMV TB40/E were synthesized by Genescrypt. Optimization of the HCMV TB40/E gene sequences for Vaccinia codon usage was performed using the Codon Optimization Tool from Integrated DNA Technologies. Runs of more than three nucleotides of the same type in a row within the codon-optimized P2A-linked HCMV gene sequences were silently mutated to enhance the stability of HCMV genes within MVA (Wyatt et al., 2009). Gene internal Kanamycin/I-SceI cassettes flanked by 50 bp gene
duplications were derived by PCR from plasmid pEPkan-S2 using primers that provided sequences for the 50 bp gene duplications. Detailed sequence maps generated by Vector NTI (Invitrogen) for all plasmids are available upon request. All cloned transfer constructs were confirmed by sequencing.

Fig. 2. Expression of P2A-linked HCMV PC subunits using MVA. MVA-PC2A1 (A) and MVA-PC2A2 (B) were passaged five times on BHK cells and whole cell lysates of infected cells of the different virus passages (1–5) were investigated by Immunoblot using rabbit polyclonal antisera and MAb specific for the individual HCMV PC subunits. MVA-PC, uninfected BHK cells, and BHK cells infected with MVA expressing HCMV gB were investigated for controls. Vaccinia protein BR5 was detected as a loading control. kDa = kilo Dalton. Arrows indicate expected molecular weights of the PC subunits.
MVA recombinants were isolated following six rounds of plaque purification using the MVA transfer vector (Fig. 1) using Fugene HD transfection reagent (Roche) according to the manufacturer’s instructions. AvrII-linearized BAC transfer vector (Fig. 1) using Fugene HD transfection reagent (Roche) according to the manufacturer’s instructions. The HCMV gB were investigated for controls. Histogram axes in A and B represent fluorescence intensity (X-axis) and cell count (Y-axis).

2.3. BAC construction

MVABAC-TK was generated by a procedure similar to that described previously by Domi and Moss (Domi and Moss, 2002). Briefly, 70–90% confluent CEF cells were infected with MVA 1974/NIH clone 1 at 0.01 multiplicity of infection (MOI) and 2 h later transfected with 2 μg of AvrII-linearized BAC transfer vector (Fig. 1) using Fugene HD transfection reagent (Roche) according to the manufacturer’s instructions. MVA recombinants were isolated following six rounds of plaque purification using GFP expression of the inserted BAC vector as a marker (Fig. 1). CEF cells were then infected at 5 MOI with the isolated MVA and grown in presence of 45 μM Isatin-β-thiosemicarbazone (IβT) to inhibit viral hairpin resolution and to promote heat-to-tail genome concatemerization and circularization (Domi and Moss, 2002). After 5 h of incubation in presence of IβT, DNA was isolated from the infected CEF cells using the DNAeasy Blood and Tissue genome isolation Kit (Qiagen) according to the manufacturer’s instructions. Numerous BAC clones were investigated by PCR and restriction fragment length analysis. One BAC clone (#1–81) was ultimately sequenced by Illumina shotgun sequencing at the City of Hope Core Facility, and the sequence of the cloned MVA genome (excluding the large terminal repeat sequences) was found to be identical to the genome sequence of MVA strain Acambis (AccessionNr. AY603355.1).

2.4. BAC recombinants

MVA vectors expressing P2A-linked HCMV PC subunits were generated by En passant mutagenesis in GS1783 bacteria cells as described previously (Tischer et al., 2010; Wussow et al., 2013). Briefly, transfer constructs of the P2A-linked UL128/130/131A or gH/gL subunits were amplified from pGEM-T-mH5 cloning vectors with primers containing 50 bp extensions homologous to the target site, and inserted into the MVA genome via an initial Red recombination. Subsequently, the kanamycin marker within the gene sequences was seamlessly removed by a second Red recombination utilizing the engineered 50 bp gene duplications flanking the marker sequences (Tischer et al., 2006; Wussow et al., 2013). For generating MVA with P2A-linked HCMV PC subunits inserted into two separate MVA insertion sites (MVA-PC2A2, Fig. 1), the UL128/130/131A and gH/gL subunit subsets were inserted by two successive En passant mutagenesis reactions into the MVA Deletion 2 site (Del2) and the intergenic region between MVA genes 69 and 70 (IGR69/70; Accession Nr. U94848), respectively. For inserting the five HCMV PC subunits all together into only one insertion site (MVA-PC2A1, Fig. 1), the UL128/130/131A subunits were inserted into the IGR69/70 by a first En passant mutagenesis reaction, and the gH/gL subunits were subsequently inserted into the UL128/130/131A-containing IGR69/70 site by a second En passant recombination reaction. All inserted HCMV PC gene were confirmed by PCR, restriction fragment length analysis, and sequencing. Detailed BAC sequence maps are available upon request. Virus reconstitution from recombinant BAC DNA was performed in BHK cells as described (Cuttingham et al., 2008; Domi and Moss, 2002; Wussow et al., 2014).

2.5. Immunoblot

Western blot analysis to detect HCMV PC subunits in whole cell lysates of MVA infected BHK cells was performed using standard procedures (Wussow et al., 2014). HCMV gl and UL131A were detected with peptide-specific rabbit polyclonal antisera (Wussow et al., 2014). UL128 was detected using MAb Z9G11 (Gerna et al., 2008), a kind gift from Dr. Giuseppe Gerna (Pavia University, Italy). gH was detected using MAb AP86 (Simpson et al., 1993), kindly provided by Dr. William Britt (University of Alabama at Birmingham). UL130 was detected using MAb 3C5 (Wang and Shenk, 2005), a kind gift from Dr. Thomas Shenk (Princeton University). Vaccinia virus BR5 was detected using MAb 19C2 (Schmelz et al., 1994).

2.6. Flow cytometry

Cell surface flow cytometry staining to detect HCMV PC subunits on MVA-infected cells by monoclonal NAb was performed as described previously (Chiuppesi et al., 2015). Briefly, BHK cells (70–90% confluent) were infected with the MVA vectors at MOI 5. At 4 h post infection, infected cells were collected, washed in phosphate buffered saline (PBS), and incubated for 1 h at 4 °C with 10 μg/ml N Ab. After washing with PBS, the cells were incubated with Alexa Fluor 647 goat anti-mouse IgG (Life Technologies) at a dilution of 1:2000. The cells were washed again and resuspended in PBS with 0.1% bovine serum albumin (BSA). Fifteen thousand events were collected using a Gallios flow cytometer (Beckman Coulter) and analyzed with FlowJo software (Tree Star).

2.7. Mouse immunization

The Institutional Animal Care and Use Committee (IACUC) of the Beckman Research Institute of City of Hope approved protocol 98004 assigned for this study. All study procedures were carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals and the Public Health Service Policy on the Humane Care and Use of Laboratory Animals. BALB/c mice (Jackson Laboratory) were vaccinated via intraperitoneal (i.p.) route with 5 × 10^5 PFU of MVA. Blood samples were collected by retro-orbital bleeding.
2.8. Neutralization assay

HCMV microneutralization assay was performed similar to published reports (Chiuppesi et al., 2015; Wussow et al., 2014). Heat-inactivated sera were serially two-fold diluted in 100 μl volumes using complete growth medium for ARPE-19 EC or MRC-5 FB depending on the cell type used in the assay. Dilutions ranged from 1:25 to 1:102400. Diluted serum was mixed with 100 μl of complete growth medium containing approximately 2400 PFU of HCMV TB40/Ewt-GFP-derived virus. After 2 h incubation, virus/sera mixtures were added in triplicate (50 μl) to ARPE-19 or MRC-5 cells seeded the day before at 1.5 × 10^4 cells/well in a clear bottom polystyrene 96-well plate (Corning) that contained 50 μl per well of complete growth medium. Cells were grown for 48 h and fixed in methanol/acetone. Infected cells were identified by immunostaining using mouse anti-HCMV IE1 Ab (p63-27 (Andreoni et al., 1989); kindly provided by William Britt (University of Alabama at Birmingham)) and the Vectastain ABC kit (VectorLabs). The substrate was 3, 3′-diaminobenzidine (DAB, VectorLabs). Plates were analyzed by an automated system using the Axio Observer Z1 inverted microscope equipped with a linear motorized stage (Carl Zeiss). IE1 positive nuclei per field of view using a 5X objective were counted using ImagePro Premier (Media Cybernetics). For each dilution the average number of positive nuclei in triplicate was calculated. The percent neutralization titer (NT) for each dilution was calculated as follows: NT = [1 – (positive nuclei number with immune sera)/(positive nuclei number with pre-immune sera)] × 100. The titers that gave 50% neutralization (NT50) were calculated by determining the linear slope of the graph plotting NT versus plasma dilution by using the next higher and lower NT values that were closest to 50% neutralization.

2.9. Statistics

GraphPad Prism software version 5.0 (GraphPad) was used to compare NAb titers in the different vaccine groups by statistical analysis using Wilcoxon matched-pairs test.

3. Results

3.1. Construction of MVA expressing 2A-linked HCMV PC subunits

Using a novel MVA BAC, termed MVABAC-TK, with genome sequence based on the MVA isolate of the NIH (Fig. 1), two different recombinant MVA vectors were generated that expressed self-processing HCMV PC subunits. Codon-optimized and 2A-linked gene sequences of the five HCMV PC subunits were inserted either as a single polycistronic expression construct into only one MVA insertion site to generate MVA-PC2A1, or as UL128/130/131A and gH/gL expression constructs into two separate MVA insertion sites to generate MVA-PC2A2 (Fig. 1). All 2A peptide sequences were based on the 2A peptide of porcine teschovirus-1 (P2A), which in combination with an N-terminal GSG-linker has recently been shown to mediate highly efficient polyprotein processing in human cells and animal models (Kim et al., 2011). DNA sequences with alternate codon usage were used to encode for P2A peptides between different HCMV PC subunits to avoid instability by homologous recombination of the multiple P2A sequence signals (Fig. 1). MVA-PC2A1 and MVA-PC2A2 were reconstituted and propagated in BHK cells, and purified virus stocks of these MVA recombinants were characterized in vitro and in vivo.
3.2. Expression of 2A-linked HCMV PC subunits using MVA

In order to characterize the expression of the P2A-linked HCMV PC subunits of MVA-PC2A1 and MVA-PC2A2, whole cell-lysates of MVA-infected BHK cells were investigated via Immunoblot analysis following five virus passages. As control, a previously generated MVA vector, termed MVA-PC, was included, which co-expressed all five HCMV PC subunits using expression constructs inserted into five separate MVA insertion sites (Wussow et al., 2014; Wussow et al., 2013). As shown in Fig. 2, both MVA-PC2A1 and MVA-PC2A2 showed comparable expression levels for the individual HCMV PC subunits during the five virus passages on BHK cells, suggesting that the PC subunits are stably maintained within these MVA vectors. In addition, the detectable protein sizes of the individual HCMV PC subunits expressed from MVA-PC2A1 and MVA-PC2A2 were consistent with anticipated molecular weight values for the HCMV PC subunits as a consequence of efficient polyprotein processing. Most of the HCMV PC subunits expressed from MVA-PC2A1 and MVA-PC2A2 showed slightly higher molecular weight values compared to their counterparts expressed from MVA-PC, which was consistent with an expected increase in protein size (∼ 2 kD) due to remaining C-terminal peptides remnants following P2A-mediated cleavage. This was in particular evident for UL128 and UL131A of MVA-PC2A1 and UL128 of MVA-PC2A2. The two protein bands that were observed for UL128 expressed either from MVA-PC2A1 or MVA-PC2A2 appeared to be specific for these MVA vectors as they were not detected in MVA-PC-infected cells and have not been described for HCMV. The C-terminal P2A peptide remnant may have rendered the UL128 protein unstable, leading to specific UL128 degradation products associated with MVA-PC2A1 or MVA-PC2A2. Compared to MVA-PC, MVA-PC2A2 appeared to express overall slightly lower amounts of the HCMV PC subunits, and lowest expression levels of the HCMV PC subunits were observed for MVA-PC2A1. In particularly UL131A expressed from MVA-PC2A1 appeared to be expressed in only very low amounts (Fig. 2). These results indicated that MVA-PC2A1 and MVA-PC2A2 stably expressed all five HCMV PC subunits that were efficiently cleaved in MVA-infected cells.

3.3. Complex assembly and formation of NAb epitope by 2A-linked HCMV PC subunits

To evaluate whether the P2A-linked HCMV PC subunits expressed from MVA-PC2A1 and MVA-PC2A2 assembled into complexes that form NAb epitopes, cell surface flow cytometry staining analysis of MVA-infected BHK cells was performed using MAb targeting PC- and gH-specific neutralizing epitopes. These MAb were previously isolated from mice immunized with MVA-PC and recognized quaternary conformational neutralizing epitopes formed by the UL128/130/131A subunits, a continuous neutralizing epitope within UL128, or a neutralizing epitope within gH (Chiuppesi et al., 2017a; Chiuppesi et al., 2015). In addition, these vaccine-derived NAb afforded similar neutralizing potency as PC- and gH-specific NAb induced by HCMV during natural infection (Chiuppesi et al., 2015). MVA-PC was used as a control in the analysis. As shown in Fig. 3, BHK cells infected with either MVA-PC2A1 or MVA-PC2A2, or control vector MVA-PC were stained efficiently and with similar intensity with all different NAb. Only slightly higher staining intensity was observed for MVA-PC2A2 compared to MVA-PC2A1 and control vector MVA-PC. Overall lowest staining intensity was observed for MVA-PC2A1. These results indicated that the HCMV PC subunits expressed from both MVA-PC2A1 and MVA-PC2A2 assembled efficiently and were transported to the cell surface as protein complexes that displayed NAb epitopes.

3.4. NAb induction by 2A-linked HCMV PC subunits in mice

To determine the immunogenicity of the P2A-linked HCMV PC subunits expressed from MVA-PC2A1 and MVA-PC2A2 to stimulate HCMV NAb, HCMV NAb induction by these MVA vectors in mice using a two-dose immunization schedule was evaluated. As control MVA-PC was used, which has previously been shown to elicit high-titer and durable HCMV NAb in mice and rhesus macaques that significantly exceeded those induced by MVA expressing only PC subunits or subunit subsets or only gB (Wussow et al., 2014). In addition, MVA-PC elicited NAb that exceeded those induced by HCMV during natural infection (Chiuppesi et al., 2015). Hence, NAb induced by MVA-PC may indirectly serve as a benchmark to elicit NAb of greater potency to those induced by HCMV in infected individuals. Balb/c mice were two times vaccinated by intraperitoneal route with the MVA vectors, and HCMV NAb in mouse sera were measured against HCMV strain TB40/E on ARPE-19 EC and MRC-5 FB by microneutralization assay over a period of 6 months. Both MVA-PC2A1 and MVA-PC2A2 stimulated robust HCMV NAb responses that were similar or even slightly elevated compared to those stimulated by control vector MVA-PC (Fig. 4). Potent HCMV NAb responses were induced by all MVA vectors after only one immunization, and these responses were efficiently boosted in all vaccine groups after the second immunization. In addition, NAb remained relatively stable in all vaccine groups until the end of the experiment. Consistent with previous observation for MVA-PC and other vaccine approaches employing the PC (Kabanova et al., 2014; Wen et al., 2014; Wussow et al., 2014), MVA-PC2A1 and MVA-PC2A2 elicited high-titer NAb that blocked EC infection and less potent NAb blocking FB infection. This is likely due to immunological properties of the HCMV PC to stimulate NAb targeting UL128/130/131A epitopes that potently and specifically block EC infection, and NAb of lower potency to gH epitopes that have the capacity to block both EC and FB infection (Chiuppesi et al., 2015; Kabanova et al., 2014). While EC NAb titers were generally comparable across all vaccine groups, FB NAb titers were significantly higher in MVA-PC2A2 immunized animals at week 7 and 24 and in MVA-PC2A1 immunized animals at week 7 compared to the control group (Fig. 4). These results indicated that the P2A-linked HCMV PC subunits expressed from MVA-PC2A1 and MVA-PC2A2 were highly immunogenic to stimulate HCMV NAb responses in mice using only two immunizations.

4. Discussion

The results of this study show in mice that the ribosomal skipping mechanism mediated by 2A peptides of picornavirus can be utilized to stimulate robust NAb responses by self-processing subunits of the HCMV PC. Using a novel MVA BAC with genome sequence identical to the MVA isolate of the NIH (excluding the terminal repeats), recombinant MVA vectors were generated that simultaneously expressed all five HCMV PC subunits either via only a single polycistronic expression construct (MVA-PC2A1) or via two separate transcripts encoding UL128/130/131A and gH/gL (MVA-PC2A2). While these MVA vectors appeared to express different amounts of the HCMV PC subunits, all five PC subunits expressed from both of these MVA vectors were efficiently cleaved and transported to the cell surface of MVA-infected cells where they formed protein complexes and displayed conformational and linear neutralizing epitopes. In addition, both of these MVA vectors were at least as potent in eliciting HCMV NAb as the previously generated MVA vector co-expressing all five PC subunits using separate insertion site, suggesting that the polycistronic MVA vectors have the potency to elicit NAb that exceed those induced by HCMV during natural infection (Wussow et al., 2014). Whether the HCMV PC subunits are stoichiometrically produced from MVA-PC2A1 and MVA-PC2A2 remains unclear. Yet, the observations that all five PC subunits were expressed in lower levels from MVA-PC2A1 compared MVA-PC2A2 may suggest that the 2A linkage tightly controls the expression levels of the individual HCMV PC subunits. While the 2A technology has been used successfully to produce soluble and highly immunogenic HCMV PC protein (Kabanova et al., 2014), this report describes the first example for the use of the 2A system to elicit robust
NAb responses by self-processing subunits of a multi-subunit herpesvirus glycoprotein complex using a clinically-deployable viral vector. Similar to the PC of HCMV, the gH/gL/gO complex of HCMV or multi-subunit envelope glycoprotein complexes of other herpesviruses such as gH/gL/gp42 of EBV play a critical role in receptor-mediated entry into host cells (Heldwein, 2016; Kabanova et al., 2016; Sathiyamoorthy et al., 2017). Considering the findings for the HCMV PC (Kabanova et al., 2014; Macagno et al., 2010; Wussow et al., 2014), it is likely that efficient vaccine-mediated induction of NAbs by NAB by multiple subunit herpesvirus glycoprotein complexes will require to express all individual complex subunits simultaneously. Because the procedure presented in this study for eliciting NAB by 2A-mediated polycistronic expression avoids the need for multiple promoter elements or insertion sites, this method could serve as a general approach for subunit vaccine development to stimulate robust NAB by multi-subunit herpesvirus glycoprotein complexes using virtually any kind of expression or delivery system. In addition, due the small size of the 2A signal sequences compared to internal ribosomal entry sites (IRES) (Kim et al., 2011; Szymczak et al., 2004), the use of 2A peptides considerably reduces the size of the element needed to combine multiple glycoprotein subunits into a single polycistronic expression construct. While the results provide proof-of-concept that the 2A cleavage system can be used to express the HCMV PC in the context of MVA, they do not a priori predict that other herpesvirus glycoprotein complexes can be expressed in a similar way using the 2A technology. The processing efficiency of other herpesviruses glycoprotein complex subunits by 2A-mediated cleavage may be significantly lower. In addition, other herpesvirus glycoprotein complex subunits may be more sensitive to negative effects of the residual C-terminal 2A peptides on subunit folding, complex assembly, stability, or translocation to the cell surface.

Because the developed procedure avoids or reduces repeated manipulation of the MVA genome to insert multiple antigens into a single MVA vector as it would be the case when using different insertion sites, this methodology may generally serve as a valuable alternative to construct multi-antigenic MVA vectors. Hence, the procedure may also be utilized for a subunit vaccine formulation to facilitate simultaneous co-expression of multiple subunits of different envelope glycoprotein complexes such as the PC and gB of HCMV to stimulate NAbs that interfere with multiple steps of herpesvirus entry including virus attachment, fusion, and receptor binding. Moreover, the system could be utilized to easily combine multiple humoral and cellular immunodominant antigens into a subunit vaccine formulation to elicit potentially protective immune responses that cover both arms of adaptive immunity. Based on the large insertion capacity and numerous insertion sites of MVA and the relative small size of the 2A peptides, MVA in combination with 2A-mediated ribosomal skipping could represent an ideal system to develop multivalent subunit vaccines to prevent the infection of herpesviruses or other viruses that utilize a multi-glycoprotein complex entry machinery for the infection of host cells.

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