M2e-Displaying Virus-Like Particles with Associated RNA Promote T Helper 1 Type Adaptive Immunity against Influenza A

Lorena Itatí Ibanéz1,2, Kenny Roose1,2, Marina De Filette1,2, Michael Schotsaert1,2, Jessica De Sloovere3, Stefan Roels3, Charlotte Pollard2,4, Bert Schepens1,2, Johan Grooten2, Walter Fiers1,2, Xavier Saelens1,2*

1 Department for Molecular Biomedical Research, VIB, Ghent, Belgium, 2 Department of Biomedical Molecular Biology, Ghent University, Ghent, Belgium, 3 Unit of Surveillance, Orientation and Veterinary Support, Operational Direction Interactions and Surveillance, Veterinary and Agrochemical Research Centre (CODA/CERVA), Brussels, Belgium, 4 Virology Unit, Department of Biomedical Sciences, Institute of Tropical Medicine, Antwerp, Belgium

Abstract

The ectodomain of influenza A matrix protein 2 (M2e) is a candidate for a universal influenza A vaccine. We used recombinant Hepatitis B core antigen to produce virus-like particles presenting M2e (M2e-VLPs). We produced the VLPs with and without entrapped nucleic acids and compared their immunogenicity and protective efficacy. Immunization of BALB/c mice with M2e-VLPs containing nucleic acids induced a stronger, Th1-biased antibody response compared to particles lacking nucleic acids. The former also induced a stronger M2e-specific CD4+ T cell response, as determined by ELISPOT. Mice vaccinated with alum-adjuvanted M2e-VLPs containing the nucleic acid-binding domain were better protected against influenza A virus challenge than mice vaccinated with similar particles lacking this domain, as deduced from the loss in body weight following challenge with X47 (H3N2) or PR/8 virus. Challenge of mice that had been immunized with M2e-VLPs with or without nucleic acids displayed significantly lower mortality, morbidity, and lung virus titers than control-immunized groups. We conclude that nucleic acids present in M2e-VLPs correlate with improved immune protection.

Introduction

Human influenza is a highly contagious respiratory disease caused by influenza A and B viruses, which undergo frequent antigenic changes. Few infections are as detrimental as influenza in terms of school and work absenteeism, medical consultation load, hospitalization, and death toll [1]. Seasonal influenza is currently counteracted by vaccination with a tri- or quadrivalent vaccine based on the haemagglutinin antigen, which is highly variable. Therefore, the composition of the influenza vaccine is adapted nearly every year so that it corresponds as much as possible to the antigenicity of the strains expected to circulate during the upcoming season [2]. The World Health Organization conducts continuous global surveillance of new influenza virus strains and guides the annual updating of the vaccines [3].

Human influenza pandemics are associated with the introduction of a novel haemagglutinin subtype or a haemagglutinin that is antigenically very different from the circulating epidemic strains. Pandemic outbreaks are unpredictable, and the Mexican flu pandemic in 2009 caused by an H1N1 virus of swine origin certainly took the world by surprise [4]. The impact of H1N1pdm2009 on human health during its first year of circulation was somewhat milder than that caused by most seasonal H3N2 influenza viruses. However, most fatal cases associated with the H1N1pdm2009 virus occurred in young people, which is unusual for seasonal flu [5,6]. Seasonal influenza vaccines offer little if any protection against a pandemic virus. In 2009, it took six months before a monovalent H1N1pdm2009 vaccine could be distributed: in other words, it became available only after the first wave of the pandemic virus attack [7]. Different approaches are being followed to develop influenza vaccines with broadened immune protection in order to control pandemic influenza outbreaks more efficiently. For example, attempts to introduce antibody-based immunity directed against conserved parts of the haemagglutinin have met this goal with some success [8,9,10]. Other universal influenza vaccine candidates are based on the induction of broadly reactive T-cell responses [11,12,13]. We and others have focused on the development of a recombinant protein vaccine based on the conserved, extracellular domain of the influenza A matrix protein 2 (M2e) linked to a carrier [14,15,16,17,18].
M2 is an integral membrane protein of 97 amino acid residues. It is scarce on virus particles but abundant on virus-infected cells [19,20]. It self-assembles as a homo-tetramer into a proton-selective ion channel [21]. During budding, M2 also changes the cell membrane curvature and promotes membrane scission and hence release of newly formed virions [22]. M2e is the amino-terminal extracellular part of M2 and consists of 23 amino acid residues. It is minimally immunogenic during infection and following conventional vaccination [23]. This might partially explain its remarkable sequence conservation across all human influenza A strains [24]. In addition, the coding information for M2e overlaps with the open reading frame of M1. This imposes a genetic constraint that limits the tolerance for mutations in M2e [25].

We fused M2e to the Hepatitis B virus core capsomer (HBc) to produce recombinant virus-like particles (VLPs). These VLPs, which display M2e on their surface at high density, are highly immunogenic and induce protection against the death and morbidity caused by challenge with influenza A virus [16,26]. Induction of protection against multiple HA-subtypes of influenza by M2e vaccines has been confirmed by using various carrier conjugates fused chemically or genetically and in both mouse and ferret models [15,27,28,29,30]. Protection induced by immunization with M2e-fusion constructs against experimental challenge is largely dependent on antibodies directed against M2e, although a contribution of an MHC class-II-restricted CD4+ T-cell epitope in M2e is likely also involved [16,31]. Antibodies directed against M2e lack virus-neutralizing activity. Instead, protection by M2e-specific IgG antibodies relies on Fc receptors and innate immune cells such as macrophages and natural killer cells [32,33,34]. Interestingly, and in line with the infection-permissive nature of M2e-based vaccines, exposure of M2e-immune mice to influenza A virus infection is compatible with the induction of cross-reactive T cells [35].

HBc-fusions are frequently used to present weakly immunogenic determinants to the immune system. HBc is a powerful immunogen that functions as both a T-cell-dependent and a T-cell-independent antigen [36] and can induce strong humoral [37], T helper (Th) [38] and even cytotoxic T cell (CTL) responses [39]. The HBc gene can be expressed efficiently in E. coli, RNA from the bacterial host is also displayed as a dense, highly immunogenic constellation [41]. The HBc capsomer contains a carboxy-terminal nucleic acid binding domain that is important for the encapsidation of the HB virus genome [42]. During recombinant expression and VLP assembly of HBc capsomers in E. coli, RNA from the bacterial host is also incorporated within VLPs. This incorporated prokaryotic RNA can act as an adjuvant to prime Th1 immunity directed against the HBc antigen [43].

Here, we compared the immunogenicity and protective efficacy of M2e-VLPs containing or lacking part of the nucleic acid binding domain of HBc. We demonstrate that in the presence of this domain, VLPs with or without the amino-terminal M2e fusion contain nucleic acids derived from the E. coli host. In addition, immunization of BALB/c mice with M2e-VLPs containing nucleic acids resulted in a higher M2e-specific serum IgG2a response as well as higher IFNγ responses in splenocytes. Although immunization of mice with M2e-VLPs (whether lacking or containing nucleic acids) protected against a lethal challenge with mouse-adapted X47 (H5N2) virus or PR/8 virus, protection against weight loss was significantly improved by the presence of entrapped RNA. We conclude that nucleic acids entrapped within M2e-VLPs improve protection against the morbidity caused by influenza A virus challenge.

Materials and Methods

2.1. Preparation and Analysis of Recombinant HBc VLPs

All VLPs were produced in and purified from E. coli. The VLPs used in this study are shown schematically in Figure 1A. VLP-1123 (M2e+/RNA−) and VLP-1818 (M2e+/RNA+) have been described [44]. VLP-1123 was derived from a truncated HBc gene coding for amino acids 1–149 followed by a cysteine residue. VLP-1818 contains three tandem copies of M2e fused to the N-terminus of VLP-1123. The N-terminal M2e copy in VLP-1818 retains the initiator methionine. In the first and second copy of M2e, the cysteine at positions 17 and 19 are mutated to serines, whereas the third M2e copy retains its cysteines. VLP-1632 (M2e−/RNA+) and VLP-1965 (M2e+/RNA+) are similar to VLP-1123 and VLP-1818, respectively, except that the HBc carrier consists of amino acids 1–163 followed by a cysteine residue. Therefore, VLP-1632 and VLP-1965 contain part of the arginine-rich carboxy-terminal domain of HBc.

All VLPs were expressed from a modified pKK223-3 plasmid (Amersham Pharmacia, UK) [44] and purified as described previously [45]. Briefly, overnight cultures of E. coli TB1 transformed with the respective expression vectors were harvested by centrifugation and lysed in Tris-EDTA buffer (50 mM Tris-HCl, 10 mM EDTA, pH 8.0) using a French press (Thermo, Cambridge, United Kingdom). Proteins in the cleared lysate were precipitated with ammonium sulfate and the precipitate was resuspended in Tris-EDTA buffer (50 mM Tris-HCl, 10 mM EDTA, pH 8.0). Recombinant VLPs were purified from the dialyzed ammonium sulfate fraction by gel filtration chromatography on a Sepharose CL-4B column (Pharmacia, Piscataway, NJ), followed by hydroxyapatite chromatography (Clarkson, South Williamsport, PA) and anion-exchange on a Mono-Q (Pharmacia). Purified VLPs were dialyzed against 20 mM sodium phosphate, pH 6.8, and the protein concentration was determined using a standard bichinonic acid assay (Pierce). The efficiency of particle formation and the presence of particle-bound nucleic acids were analyzed by agarose gel electrophoresis. VLPs (10 µg) were loaded on a 2% agarose gel and run for 2 h at 80 V in TAE buffer. The gel was stained with ethidium bromide to reveal nucleic acids and with Coomassie brilliant blue to reveal proteins. In addition, 2 µg of each VLP was boiled for 5 min in reducing Laemmli sample buffer, separated by SDS-PAGE, and stained with Coomassie brilliant blue.

Nucleic acids from 40 µg VLP-1632 were isolated using the NucleoSpin RNA Virus kit (Macherey-Nagel) according to manufacturers recommendations, including the optional proteinase K treatment but without addition of carrier RNA. Total cellular RNA was isolated from HEK293T cells using by the use of RNAeasy kit (Qiagen, Hilden, Germany). One µg of purified VLP-1632 nucleic acids, cellular RNA, miniprep plasmid DNA, as well as control plasmid mini prep DNA that was additionally submitted to the procedures used for the isolation of the VLP nucleic acids were treated with either PBS, 5 µg RNAse from bovine pancreas (Sigma-Aldrich, St. Louis, USA) or 1.4 Kunitz of DNase I (Qiagen) for 30 minutes at 25°C. The samples were then analysed by agarose gel electrophoresis followed by SYBR Safe staining.

The LPS-content of different VLP preparations was determined using the Toxinsensor Chromogenic LAL Endotoxin Assay Kit (GenScript, NJ, USA) according to the manufacturer’s instruc-
Briefly, a standard curve was generated from a two-fold dilution series of *E. coli* endotoxin standard in LAL reagent water (1 EU – 0.125 EU). Two-fold dilution series of VLPs were made in LAL reagent water and assayed along with the standard and a blank control. After 10 min incubation at 37°C of samples and controls with LAL (Limulus Amebocyte Lysate), and at the end of the reactions absorption was measured spectrophotometrically at 545 nm, using LAL reagent water as a blank. Endotoxin contents in VLP preparation were calculated using the standard curve and expressed as EU/mL or EU/µg of protein.

2.2. Mouse Vaccination and Challenge

All mouse experiments were conducted according to the national (Belgian Law 14/08/1986 and 22/12/2003, Belgian Royal Decree 06/04/2010) and European (EU Directives 2010/63/EU, 86/609/EEG) animal regulations. Animal protocols were

![Figure 1. The C-terminal arginine-rich domain of recombinant HBc-based VLPs binds nucleic acids.](image)

A. Schematic representation of the recombinant M2e-VLPs and control VLPs used in this study. Light grey boxes on the left in VLP-1818 and VLP-1965 represent three tandem M2e copies; white boxes represent HBc amino acid residues 1–149; dark grey boxes on the right in VLP-1965 and VLP-1632 represent part of the arginine-rich domain of HBc (residues 150–163). In all constructs, a C-terminal cysteine residue (cys) was engineered to improve particle stability. B. Native agarose gel analysis of VLPs, followed by staining with Coomassie Brilliant blue (left) or ethidium bromide (right). C. Purified recombinant VLPs (2 µg each) were separated by SDS-PAGE and stained with Coomassie Brilliant Blue. D. VLP-1632 contains RNA. One µg of nucleic acids purified from VLP-1632, similarly purified plasmid DNA, plasmid DNA and total mammalian cellular RNA were treated with either PBS, RNAse or DNAse and analyzed by agarose gel electrophoresis followed by SYBR Safe staining. Numbers on the left indicate sizes in base pairs of the DNA marker in the left hand lane.

doi:10.1371/journal.pone.0059081.g001
approved by the ethics committee of Ghent University (permit number LA1400091, approval ID 2010/001). All efforts were made to minimize the suffering of the animals. Specific-pathogen-free female BALB/c mice were obtained from Charles River (France) and immunized at the age of eight weeks. The animals were housed in a temperature-controlled room (biosafety level 2) with 14/10-h light/dark cycles and received food and water ad libitum.

Figure 2. VLPs containing nucleic acids promote a Th1-polarized humoral immune response. A. Three groups of 16 BALB/c mice were immunized twice intraperitoneally with 10 μg of VLP-1123, VLP-1818 or VLP-1965, without adjuvant. Priming and boosting injections were given at three week intervals. Two weeks after the boost, M2e-specific serum IgG1 and IgG2a titers were determined by M2e peptide ELISA. White and black bars: serum samples after priming and boosting, respectively. IgG1/IgG2a ratios were calculated by dividing the mean of the IgG1 endpoint titers by the mean of the IgG2a endpoint titers, and are shown on the right. (*: P<0.05, **: P<0.01 two-sample t-test). B. The particles were adjuvanted with Alhydrogel and used to immunize BALB/c mice (14 per group, 10 μg of VLP i.p. per animal). Two weeks after priming, M2e-specific (left) and HBc-specific (right) IgG1 and IgG2a titers were determined in sera pooled from the 14 mice. C. Groups of 12 BALB/c mice were immunized twice by i.p. injection of 10 μg of VLP-1123, VLP-1623, VLP-1818 or VLP-1965 formulated with Alhydrogel. Priming and boosting injections were three weeks apart. M2e-specific serum IgG1 and IgG2a titers are shown in the top panels and HBc-specific titers in the bottom panels. White and black bars: serum samples two weeks after priming and boosting, respectively. (*: P<0.05, **: P<0.01, ***: P<0.001, two-sample t-test). Bars in A and C represent averages and error bars represent standard error of the mean. #: titer lower than 100.

doi:10.1371/journal.pone.0059081.g002
M2e-VLPs or control VLPs (10 μg each) were used without adjuvant or adjuvanted with Alhydrogel (Brenntag Biosector A/S Fredriksund, Denmark) and injected intraperitoneally (i.p.). Particles were adsorbed using a 10% (v/v) solution of a 2.0% Alhydrogel stock solution prior to injection. Mice were primed and boosted at intervals of two or three weeks as indicated in the figure legends. To evaluate the protective efficacy of the vaccine, mice were challenged with 4 LD₅₀ of mouse-adapted X47 (H3N2) virus or with 4 LD₅₀ of PR/8 virus, as indicated in the figure legends. The mice were anesthetized by i.p. injection of a mixture of ketamine (12 mg/kg) and xylazine (60 mg/kg). Survival and body weight were monitored for 14 days after challenge.

2.3. Determination of Serum Antibody Levels and M2e-specific Cellular Responses

Two weeks after each immunization, blood samples were collected from the lateral tail vein. Blood was allowed to clot for 30 min at 37°C, and serum was obtained by combining the supernatants from two successive centrifugations. The titers of anti-M2e antibodies and of IgG subclasses in the serum were determined by ELISA. Serum samples. HBc-specific antibodies were determined using a sandwich ELISA as described [44].

M2e-specific T cell responses were evaluated by measuring interferon-γ (IFN-γ) by enzyme-linked immunospot (ELISPOT). This analysis was done on splenocytes from mice that had been immunized twice two weeks apart by i.p. injection of Alhydrogel-adjuvanted VLP-1123, VLP-1632, VLP-1818, VLP-1965 or PBS. ELISPOT plates were purchased from U-Cytech Biosciences (Utrecht, The Netherlands) and used according to the manufacturer’s protocol. Briefly, 96-well immunoplates were coated with monoclonal anti-IFN-γ antibodies and blocked with blocking buffer. After 14 days after the second immunization, and six days after challenge with 4 LD₅₀ of X47 virus, the spleens of five mice per group were isolated aseptically and single-cell suspensions were prepared. After lysis of red blood cells with ammonium buffer, 3 × 10⁵ splenocytes were plated in 100 μl of culture medium supplemented with 4 μg/ml of HPLC-purified peptide for restimulation. Splenocytes from each mouse were analyzed in triplicate. The M2e-specific peptide was SLLTVETPFR-NEWGCRCDSSD (M2e), and the negative control peptide was MNNAFTNVTNPNISHIR. For ELISPOT analysis of splenocytes isolated six days after challenge, nucleoprotein-specific responses were determined by restimulation with the H2d-restricted NP-derived TYQRTRALV peptide. After 16 h of peptide restimulation, plates were washed with ELISA wash buffer and IFN-γ trapped on the plates was detected by a biotinylated polyclonal anti-IFN-γ antiserum. Subsequent incubation with GABA-conjugated streptavidin was used to develop silver spots at places were immune cells secreted IFN-γ during peptide restimulation. Splenocytes from each mouse were analyzed in triplicate and the spots were counted using an inverted light microscope.

2.4. Bone-marrow Derived Dendritic Cell (BMDC) Isolation, Stimulation, and RT-qPCR Analysis

Bone-marrow was isolated by flushing the tibia and the femur of WT, MyD88⁻/⁻ and TRIF⁻/⁻ mice (all in the B6 background) with ice-cold PBS. After lysing red blood cells with ACK red blood cell lysis buffer (BioWhittaker, Wakersville, MD, USA), 3.5 × 10⁶ bone marrow cells were seeded in petri dishes and cultured for 8 days in 10 ml RPMI 1640 (Invitrogen, Merelbeke, Belgium) supplemented with 10% fetal calf serum and 20 ng/ml GM-CSF (Peprotech, London, UK). Medium was replenished every 3 days. On day 8, 1 x 10⁶ immature dendritic cells were stimulated with 20 μg of VLP-1123, VLP-1632, VLP-1818, VLP-1965 or 1 μg ODN 1826 (Invivogen, San Diego, CA) for 20 h. RNA was isolated using the RNeasy Plus Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocol. cDNA was synthesized using a Superscript II Reverse Transcription Reagent Kit (Invitrogen, Merelbeke, Belgium). Real-time quantitative PCR (qPCR) was performed on a Lightcycler 480 using a qPCR kit for SYBR Green I (both Roche Diagnostics, Mannheim, Germany). Real-time qPCR amplification was performed in triplicate reactions. mRPL13a mRNA was used as a reference housekeeping gene for normalization. All primers were purchased from Invitrogen. The PCR conditions were as follows: preincubation at 95°C for 5 min; 50 cycles at 95°C for 10 s and at 60°C for 30 s. The following forward and reverse primers were used: 5’-TAGTCCTTCTTGACCATTTAGCG-3’ and 5’-TTGTTGCTTTAGCCACTTCCTTG-3’ (murine il1β); 5’-CAGCTC- CACAAAGCAGAGCATCAG-3’ and 5’- GCATTAGAAG GATGCAGACCTAC-3’ (murine il10); 5’- CCTGCTGTCCTCAAGGTGTTT-3’ and 5’- TGGCCTGTCGACTGCTGGAAGTACT-3’ (murine il13a).
2.5. Determination of Lung Virus Titers

Three to five mice from each group were killed by cervical dislocation four or six days after challenge. The lungs were removed aseptically and homogenized in 1 ml of sterile, ice-cold PBS. The extracts were transferred to centrifuge tubes and cell debris was pelleted for 10 min at 400 g and 4°C. The titers of infectious virus were determined in triplicate by titration of the cleared lung extracts on MDCK cells. Briefly, monolayers of cells were infected for 1 h with 50 μl of serial 1:10 dilutions of the lung homogenates in a 96-well plate in serum-free DMEM medium containing penicillin and streptomycin (Invitrogen). Following infection, the medium was replaced by medium containing 2 μg/ml of TPCK-treated trypsin (Sigma-Aldrich, Germany). Endpoint virus titers were determined after four days, as described by Reed and Muench, by interpolating the dilution that infected 50% of the wells, as determined by the presence of chicken red blood cell hemagglutinating activity in the supernatants.

Figure 4. VLP-associated nucleic acids signal through MYD88 and TRIF. RT-qPCR was used to analyze mRNA expression levels of the pro-inflammatory cytokines IL-1β and IL-6 20 hours after stimulation of WT, MyD88−/− or TRIF−/− DCs with 20 μg of VLP-1123, VLP-1632, VLP-1818, VLP-1965 or 1 μg of ODN 1826. Results represent mean n-fold induction levels compared to unstimulated control cells ± SD from triplicate PCR reactions.

doi:10.1371/journal.pone.0059081.g004
confirming their uniform particulate nature [46] (Figure 1B, left). In native agarose gel electrophoresis, SDS-PAGE analysis demonstrated that the purified particles were composed of subunits of the expected molecular weight (Figure 1C). In native agarose gel electrophoresis, SDS-PAGE analysis demonstrated that the purified particles were composed of subunits of the expected molecular weight (Figure 1C). In native agarose gel electrophoresis, SDS-PAGE analysis demonstrated that the purified particles were composed of subunits of the expected molecular weight (Figure 1C).

### Results

#### 2.6. Histopathology

Four days after challenge, mice were euthanized by cervical dislocation and samples of the apical, median and caudal regions of the lungs (bilateral) were removed, fixed in 4% paraformaldehyde, embedded in paraffin wax, sectioned at 5 μm, and stained with hematoxylin-eosin. Slides were examined in a blinded fashion using an Olympus BX 50 microscope and micrographs were made.

#### 2.7. Statistical Analysis

Comparison of antibody titers between two groups was done with the two-tailed Student’s t-test. Survival rates were plotted as Kaplan-Meier curves and analyzed with the log-rank test. Morbidity parameters after infection of more than two groups were compared using the Holm-Sidak method. Statistical analyses were performed with Graphpad Prism version 4.00 for Windows (GraphPad Software, San Diego California, www.graphpad.com) and with the R language and environment for statistical computing, R Development Core Team, 2009 (R Foundation for Statistical Computing, Vienna, Austria (ISBN 3-900051-07-0, www.R-project.org).

**Results**

**In vitro Characterization of M2e-VLPs Containing or Lacking Bound Nucleic Acids**

Previously, we reported on the characterization and immunoprotective efficacy of M2e-VLPs produced by the HBc presentation technology [16,44]. In these studies we used HBc VLPs that either contained or lacked the carboxy-terminal arginine-rich domain of HBc. This domain is responsible for nucleic acid binding and was shown to prime the mammalian immune response towards a Th1 response because it facilitates the incorporation of RNA within the HBc-based VLP [43]. Such a Th1 response is favorable for anti-M2e mediated protection [32,44]. To investigate if HBc VLPs containing the carboxy-terminal arginine-rich domain can induce a more favorable M2e immune response than HBc VLPs lacking this domain, we constructed M2e-presenting HBc VLPs and corresponding control VLPs. The carboxy-terminal arginine-rich domain of HBc was either included [VLP-1632 (M2e+/RNA+) and VLP-1632 (M2e+/RNA−)] or excluded [VLP-1818 (M2e+/RNA+) and VLP-1123 (M2e+/RNA−)] (Figure 1A). SDS-PAGE analysis demonstrated that the purified particles were composed of subunits of the expected molecular weight (Figure 1C). In native agarose gel electrophoresis, all VLPs migrated predominantly as single bands, confirming their uniform particulate nature [46] (Figure 1B, left panel). As expected, co-migrating nucleic acids, revealed by ethidium bromide staining, were clearly detectable only in VLP-1632 particles by proteinase K treatment followed by silica membrane-based purification. These purified nucleic acids derived from VLP-1632 were susceptible to RNAse and resistant to DNase treatment (Figure 1D).

#### Table 1. Characteristics of virus-like particles used in this study.

| VLP     | Make-up       | EU/ml*       | EU/μg of protein* |
|---------|---------------|--------------|-------------------|
| VLP-1123| M2e−/RNA−     | 2485         | 2.16              |
| VLP-1632| M2e−/RNA+     | 50880        | 46.26             |
| VLP-1818| M2e+/RNA−     | 250022       | 320.54            |
| VLP-1965| M2e+/RNA+     | 7840         | 18.67             |

*LPS-content per ml of VLP solution or per μg of protein, expressed as international units and as determined by the LAL-assay.

doi:10.1371/journal.pone.0059081.t001

Nucleic Acids Bound to M2e-VLPs Correlates with a Th1-polarized Humoral Immune Response

We next compared the immunogenicity of the VLP types. BALB/c mice (n = 16 per group) were immunized by i.p. injection of 10 μg of unsubstituted control VLPs, VLP-1818 or VLP-1965. In the absence of added adjuvant, the M2e-specific IgG1 titers in serum were significantly lower in mice that had been immunized with VLP-1965 than in VLP-1818 immunized mice. Conversely, M2e-specific IgG2a titers were significantly higher in mice immunized with VLP-1965 compared to VLP-1818 vaccine recipients, and this was true after both priming and boosting (Figure 2A).

Adjuvants promote the immunogenicity of M2e-fusion constructs [16]. The alum-based adjuvant Alhydrogel is suitable for human use and we previously reported that this adjuvant enhances the immunogenicity of M2e-VLPs carrying a single M2e epitope [44]. Therefore, we also analyzed the effect on immunogenicity of the particle-associated nucleic acids when these particles were formulated with Alhydrogel. To be able to compare our results with those of Riedl et al., who used a single immunization without adjuvant [43], we analyzed M2e- and HBc-specific antibodies two weeks after priming in groups of 14 BALB/c mice. In this experiment, the IgG1 titers were lower for alum-adjuvanted particles containing bound nucleic acids (VLP-1632 and -1965) than for those lacking it (VLP-1123 and -1818). In contrast, serum IgG2a titers against M2e and against the carrier were clearly higher in recipients of alum-adjuvanted VLPs with associated nucleic acids (Figure 2B).

We next performed an independent prime-boost experiment with alum-adjuvanted particles. M2e-specific serum IgG2a responses were higher following vaccination with VLP-1965 than following VLP-1818 (Fig. 2C). Likewise, significantly lower IgG1 and higher serum IgG2a titers directed against the HBc carrier were apparent when nucleic acid-containing VLPs were used, which was observed for the unsubstituted particles (VLP-1123 compared to VLP-1632) after prime and boost and only after the boost immunization for the M2e-displaying particles VLP-1818 and VLP-1965 (Fig. 2C).

M2e contains an MHC-II-restricted T cell epitope that can contribute to immune protection in the BALB/c mouse model [31]. Therefore, we used IFN-γ ELISPOT to compare the magnitude of the M2e-specific T helper cell responses. Mice that had been immunized with VLP-1965 displayed a significantly stronger splenic Th1 response upon in vitro restimulation with M2e peptide than mice immunized with VLP-1818 recipient mice (Fig. 3).

From these results we conclude that extending VLPs with part of the carboxy-terminal nucleic-acid-binding domain of the HBc

---

1965 and in the corresponding control particle VLP-1632, both of which harbour the nucleic acid binding domain (Figure 1B, right panel). Incubation of the particles at 25 °C for one hour with RNase or DNase failed to remove the nucleic acids, suggesting that the nucleic acids were trapped within the VLPs and hence protected from exogenous nucleases under these conditions (data not shown). Therefore, we isolated the nucleic acids from VLP-1632 particles by proteinase K treatment followed by silica membrane-based purification. These purified nucleic acids derived from VLP-1632 were susceptible to RNAse and resistant to DNase treatment (Figure 1D). We conclude that extending the HBc capsid protein at its amino-terminal end with multiple copies of M2e affects neither VLP formation nor the nucleic acid (most likely RNA) binding capacity of these VLPs.
Figure 5. Immunization with VLP-1965 provides better protection against morbidity caused by influenza A virus challenge than VLP-1818. A. Groups of 12 BALB/c mice were immunized twice with an interval of three weeks with 10 µg of the indicated VLPs, adjuvanted with Alhydrogel. Three weeks after the second immunization, mice were challenged with 4 LD₅₀ of mouse-adapted X47 virus. Survival rate (left; P<0.001, Kaplan Meier test) and body weight (right; P<0.05, two-sample t-test on days 6 and 7 after challenge) were monitored for two weeks starting from the day of challenge. B. Groups of 8 (PBS), 12 (VLP-1818) or 13 (VLP-1965, -1632 and -1123) BALB/c mice were immunized twice with a two-week interval with 10 µg of the indicated VLPs, adjuvanted with Alhydrogel. One group of mice received PBS with Alhydrogel (PBS). Two weeks after the second immunization, mice were challenged with 4 LD₅₀ of mouse-adapted X47 virus. Survival rate (left; P<0.001, Kaplan Meier test) and body weight (right; P<0.05, two-sample t-test on days 6 after challenge) C. Groups of 14 BALB/c mice were immunized three times with three-week intervals by i.p. injection of 10 µg of VLP-1123, VLP-1818 or VLP-1965, adjuvanted with Alhydrogel. Three weeks after the second boost, mice were challenged with 4 LD₅₀ of PR/8 virus. Survival rate (left; P<0.0001) and body weight (right; P<0.05 on days 8, 10 and 12 after challenge).

doi:10.1371/journal.pone.0059081.g005
deficiency evoked a strong decrease in the expression level of IL-1β agonist ODN 1826 which signals through MyD88. MyD88 VLP-1965. As a control, BMDCs were incubated with the TLR-9 stimulated with 20 ng of X47 virus. Six days after challenge, four (PBS) or five (all other groups) mice from each group were sacrificed to determine lung virus titers. Virus titers are expressed as mean log_{10} TCID_{50}/ml and error bars depict the standard error of the mean.
doi:10.1371/journal.pone.0059081.g006

capsomer correlates with a Th1 type of immune response in BALB/c mice.

MYD88 and TRIF are Involved in Gene Induction by HBc-based VLPs

Toll-like receptors (TLRs) are vital components of the mammalian innate immune system. They recognize microbial molecular patterns such as double-stranded RNA, 5’ uncapped RNA, CpG DNA, and LPS. Engagement of TLRs by their cognate ligands results in the induction of type I IFN and inflammatory cytokine genes. This TLR-dependent gene activation depends on the adaptor proteins myeloid differentiation primary response protein 88 (MyD88) and TIR domain-containing adapter inducing IFNβ (TRIF) and may promote a subsequent adaptive immune response [47]. To obtain some mechanistic insights in the contribution of these adapter proteins to the immunogenic profile of the VLPs used in this study, we compared the expression levels of the pro-inflammatory genes IL-6 and IL-1β in BMDCs derived from TRIF−/−, MyD88−/− and WT mice stimulated with 20 μg of VLP-1123, VLP-1632, VLP-1818 or VLP-1965. As a control, BMDCs were incubated with the TLR-9 agonist ODN 1826 which signals through MyD88. MyD88 deficiency evoked a strong decrease in the expression level of IL-1β and completely blocked the induction of IL-6. In TRIF−/− DCs, expression of IL-6 was profoundly downregulated while the decrease in IL-1β expression was more modest (Fig. 4). Comparison of the LPS content of the E. coli-derived recombinant VLPs, revealed a variable level with the lowest amount found in VLP-1123 and VLP-1965 and more significant amounts in VLP-1632 and VLP-1818 (Table 1). Taken together, we found no clear correlation between LPS-content, RNA-content and MyD88- or Trif-dependent gene induction from these in vitro BMDC stimulation assays. It is possible that the contribution of the RNA-contained in VLP-1632 and VLP-1965 was masked by LPS or other microbial-derived TLR agonists.

Incorporation of Nucleic Acids in VLPs Improves Protection by M2e-VLPs Vaccination

We next compared the protective efficacy of VLP-1818 and VLP-1965 particles. M2e-immune mice survive a lethal influenza A virus challenge but they usually lose some weight. The extent of weight loss in M2e-immune mice depends on the adjuvant, route of immunization and antibody isotype [48]. In BALB/c mice, IgG2a antibodies directed against M2e protect better than their IgG1 counterpart [32]. In independent immunization experiments we consistently noticed significantly higher M2e-specific IgG2a antibody titers in serum of mice that had been primed and boosted with VLP-1965 particles containing nucleic acids than mice that received VLP-1818 (Fig. 2). Because we previously found that immunization with VLP-1818 protected against a potentially lethal challenge with influenza A virus [44], we anticipated that it would require a very large number of mice to obtain a significant difference in survival outcome when applying this parameter as a read out. Therefore, we focused on body weight drop to compare protective efficacy following challenge and we did so in independent experiments. This comparison of efficacy between VLP-1818 and VLP-1965 was important because its outcome would help to guide the choice of the lead M2e-VLP vaccine candidate for a Phase I clinical study.

First, we vaccinated groups of 12 BALB/c mice twice three weeks apart with 10 μg of the VLPs in the presence of Alhydrogel. Three weeks after the boost, mice were challenged with 4 LD_{50} of mouse-adapted X47 virus, and morbidity and mortality were monitored. All mice immunized with VLP-1818 or VLP-1965, except for one in the VLP-1818 group, survived the challenge. Morbidity after challenge was most severe in mice immunized with control VLP-1123 or VLP-1632 and all animals in these groups died after challenge (Figure 5A). Interestingly, during the peak days of morbidity following challenge, VLP-1965-immunized mice lost significantly less body weight than mice immunized with VLP-1818 (Figure 5A).

In an independent second experiment, we compared the protective efficacy in groups of BALB/c mice that had been immunized twice with 10 μg of control or M2e-substituted VLPs. Alhydrogel was used as adjuvant, a PBS-Alhydrogel group was also included and priming and booster immunizations were two weeks apart. Two weeks after the booster immunization, mice were challenged with 4 LD_{50} of mouse-adapted X47 virus and monitored for survival and morbidity. Following challenge, all mice in the control groups gradually lost weight and eventually died or had to be euthanized. In contrast, all VLP-1965 and all except one VLP-1818 immunized mice survived the challenge (Figure 5B). Body weight loss was again higher in the VLP-1818 group and differed significantly from that of the VLP-1965 mice on day 6 after challenge (Figure 5B).

Finally, we also compared the effect of an additional boost vaccination on control of morbidity, between Alhydrogel-adjuvanted VLP-1818 and VLP-1965 vaccines. Groups of 14 mice were used and challenge was with PR/8 virus, which is highly virulent to mice [49]. All VLP-1123 immunized mice died after infection whereas M2e-immunized survived. Again, VLP-1965-immunized mice experienced significantly less body weight loss than mice immunized with VLP-1818 (Figure 5C). In summary, mice vaccinated with VLP-1965 or VLP-1818 survived a potentially lethal influenza A virus challenge. In addition, VLP-1965 was more protective than VLP-1818 against morbidity caused by challenge.

Lung Virus Load is Reduced in M2e-immune Mice

In the aforementioned challenge experiments, we also determined the lung virus load. Three to five mice from each group were sacrificed on day 4 or 6 after challenge. We consistently observed a one to two log lower virus load in VLP-1818 and VLP-1965 immunized mice compared to control groups. As an example, Figure 6 depicts the lung virus loads in mice that had...
been immunized twice with 10 μg of Alhydrogel-adjuvanted control (PBS, VLP-1123 or VLP-1632) or M2e-substituted VLPs (VLP-1818 or VLP-1965) and challenged with 4 LD₅₀ of X47 virus. However, there was no significant difference in lung virus titer between mice vaccinated with VLP-1965 or with VLP-1818.

**Cellular Immune Responses Following Challenge**

We recently reported that M2e-immune mice that are challenged with a sublethal dose of X47 virus mount cellular immune responses against the viral nucleoprotein (NP). These NP-specific T-cell responses were similar in magnitude to those in challenged naive mice [35]. Therefore, we determined the M2e- and NP-specific T cell responses in splenocytes isolated six days after X47 challenge. An IFN-γ specific ELISPOT analysis of splenocytes stimulated in vitro with M2e peptide showed that after challenge, VLP-1965 immunized mice had significantly stronger cellular responses than VLP-1818 immunized mice (Fig. 7A).

These responses were presumable due to the activation of CD4⁺ T-cells [31]. However, this result should be interpreted with care because these M2e-specific T cell responses include those induced by vaccination in the VLP-1818 and VLP-1965 immunized mice as well as those induced by the influenza A virus challenge. IFN-γ specific ELISPOT analysis revealed comparable NP-specific responses in splenocytes from all mice, even though M2e-immune mice were protected from challenge whereas control mice died (Figure 7A).

**Discussion**

We describe the use of VLP-1965, an improved M2e-based universal influenza A vaccine candidate, and compare its properties and efficacy to the previously reported VLP-1818 [44]. The carboxy-terminus of VLP-1965 comprises part of the arginine-rich domain of HBc (amino acid residues 150–163), allowing these particles to trap nucleic acids. Immunization with...
either of these M2e-displaying VLPs protected mice from death from a potentially lethal influenza A virus challenge. Furthermore, after challenge, M2e-VLP immunized mice displayed less histopathological abnormalities than control-VLP-immunized mice: the alveolar structure was less affected, and alveolar pathology was more focal [Figure 7B]. However, immunization with M2e-displaying VLPs harboring the RNA-binding domain (VLP-1965) protected better against weight loss following challenge than immunization with VLPs lacking this domain (VLP-1818). In contrast, immunization with the control VLPs did not protect and did not result in a difference in morbidity in mice immunized with either control VLP. This indicates that the reduction in morbidity was due to induction of an adaptive immune response against the M2e-antigen and not to the induction of inflammation. Lung virus titers were lower in M2e-immunized mice compared to the controls whereas viral loads determined on day six after challenge were comparable in the VLP-1818 and VLP-1965 immunized mice. This may be explained by the lack of in vitro neutralizing activity of M2e-specific IgG [35]. In addition, a difference in morbidity does not necessarily correlate with a difference in lung virus load in the mouse model for influenza [50].

M2e-based protection against influenza can be mediated by M2e-specific antibodies but might also depend on M2e-specific CD4 T cells [31]. M2e-specific IgG protects mice from influenza infections by a mechanism that involves both Fcγ receptors and alveolar macrophages [32]. As M2e-specific antibodies probably cannot readily bind to influenza particles, it is likely that M2e-specific antibodies protect against infection by eliminating the infected cells via ADCC or phagocytosis. We previously demonstrated that M2e-specific IgG2a antibodies can protect against influenza infections more efficiently than M2e-specific IgG1 antibodies. In addition, M2e-specific IgG1 failed to protect Fcγ receptor III knockout mice, but M2e-specific IgG2a could rescue protection in these animals [32]. Vaccination with M2e-VLPs harbouring the RNA-binding domain induces higher levels of M2e-specific IgG2a antibodies. In this way, the enhanced protection triggered by immunization with M2e-VLPs harbouring an RNA-binding domain might be explained by the enhanced induction of IgG2a antibodies. So by promoting the induction of M2e-specific IgG2a antibodies, VLP-1965 could promote Fcγ-receptor-mediated elimination of infected cells, thereby further limiting inflammation and morbidity.

Following immunization, nucleic acids bound within VLP-1965 are presumably taken up by antigen-presenting cells. The vertebrate innate immune system is armed with both extracellular and intracellular molecular sensors that recognize pathogen-associated molecular patterns such as unmethylated CpG DNA and single- and double-stranded RNA [51]. Triggering of these sensors, e.g. the Toll-like receptors TLR-3, -7, -8, and -9 by nucleic acids induces a signaling cascade that leads to the expression of several cytokines. These in turn activate the innate immune system, which then programs a specific immune response. Riedl et al demonstrated that deleting the Argyriev C-terminus of HBc results in particles of 149-aa that retain less than 2% of the RNA binding capacity of full-length HBc [43]. This reduced RNA binding correlated with an IgG-isotype switch from the induction of a predominantly Th1 response with high IgG2a titers (upon vaccination with full-length HBc) towards a Th2 response with high HBc-specific IgG1 antibody titers (upon vaccination with truncated HBc-149 lacking nucleic acids) [43]. VLP-1965 combines three tandem M2e repeats presented at high density on its surface and entraps nucleic acids. Most likely, these nucleic acids (presumably bacterial RNA) bias the M2e-specific immune response towards a Th1-type, as evidenced by the enhancement of the serum titer of anti-M2e IgG2a. A major Th1-type cytokine is IFN-γ. Indeed, based on our ELISPOT data, M2e-specific T cell responses resulting in IFN-γ production were increased in the spleen of VLP-1965 immunized mice. In addition, we compared IL-6 and IL-1β mRNA levels induced by in vitro stimulation of differentiated dendritic cells derived from TRIF−/−, MyD88−/− and WT mice with VLP-1632 or VLP-1965. In the absence of MyD88 and to a lesser extent of TRIF, the induction of these pro-inflammatory genes was strongly diminished, suggesting the involvement of Toll-like receptors in costimulation. Interestingly, bacterial mRNA co-administered with bacterial antigens, acts as a pathogen-associated molecular pattern that promotes humoral responses, including switching of antibody isotype classes [52].

M2e-specific IgG2a antibodies play an important role in the protection induced by the M2e-vaccine against influenza [32,46]. These findings are corroborated by the improved protection obtained with VLP-1965 relative to VLP-1818. In addition, we recently demonstrated that M2e-specific IgG1 failed to protect Fcγ-receptor III knockout mice, and that M2e-specific IgG2a could rescue protection in these animals [32]. An important role for IgG2a antibodies in protection against influenza has also been demonstrated in SCID mice by passive immunization with anti-M2e IgG monoclonal antibodies [32,53].

Conventional influenza vaccines hamper the development of heterosubtypic immunity endowed by T-cells [35]. Such T cell responses are induced by natural infection with influenza viruses and many of these responses are cross-protective because they are mainly directed against conserved internal structural proteins. Following infection with influenza A virus, M2e-immune mice mount cytotoxic T cell responses directed against NP and HA that are comparable to those in sham-vaccinated mice, as we recently reported in a study involving the VLP-1018 vaccine antigen [35]. It was beyond the scope of the current work to extensively compare such T cell responses in challenged mice that had either been vaccinated with either VLP-1818 or the Th1-biasing VLP-1965 vaccine, but nevertheless we found that the latter particles did not compromise the T cell responses directed against NP (Figure 7).

VLP-1965 was recently evaluated successfully in a Phase I clinical study to document its safety and immunogenicity in humans (ClinicalTrials.gov, Identifier NCT00819013). The M2e-vaccine remains a valid panlemic influenza vaccine candidate and also holds promise as a vaccine that could be used in young children to control disease without compromising T cell immunity upon exposure to seasonal or pandemic influenza [33]. Alternatively, the M2e-vaccine could be used as an adjunct of the currently licensed influenza vaccines to broaden their protective efficacy [34]. VLP-1965 is a promising candidate for such strategies.

Acknowledgments

We thank Dr. Amin Bredan for editing the manuscript. We thank Anouk Smet, Tine Ysenbaert and Liesbeth Vande Ginste for excellent technical support.

Author Contributions

Conceived and designed the experiments: BS JG WF XS. Performed the experiments: LJJ KR MDF MS JDS SR CP. Analyzed the data: LJJ KR MDF MS JDS SR CP BS JG WF XS. Contributed reagents/materials/analysis tools: JG WF XS. Wrote the paper: XS.
References

1. Thomp WW, Conner L, Shao DK (2006) Epidemiology of seasonal influenza: use of surveillance data and statistical models to estimate the burden of disease. The Journal of infectious diseases 194 Suppl 2: S82–91.

2. Wood JS, Robertson JS (2007) Reference viruses for seasonal and pandemic influenza vaccine preparation. Influenza Other Respi Viruses 1: 5–9.

3. Nichol KL, Trenam JG (2006) Vaccines for seasonal and pandemic influenza. J Infect Dis 194 Suppl 2: S111–118.

4. Garten RJ, Davis CT, Russell CA, Shu B, Lindstrom S, et al. (2009) Antigenic and genetic characteristics of swine-origin 2009 A(H1N1) influenza viruses circulating in humans. Science 325: 197–201.

5. Dawood FS, Iuliano AD, Reed C, Meltzer MI, Shay DK, et al. (2012) Estimated global mortality associated with the first 8 months of 2009 pandemic influenza A (H1N1) virus circulation: a modelling study. The Lancet infectious diseases. 12: 1570–1576.

6. Wu YP, Tang SS, Tse CM, Yuen KY (2006) Detection of prokaryotic mRNA signifies microbial viability and promotes recognition by the innate immune system. Annual review of immunology 29: 185–214.

7. Hillare ML, Österhaus AD, Rimmelzwaan GF (2011) Induction of virus-specific cytotoxic T lymphocytes as a basis for the development of broadly protective influenza vaccines. Journal of virology & virology. 2011: 099060.

8. Price GE, Sobolev SA, Johnson R, Major D, et al. (2011) The nucleocapsid of hepatitis B virus as carrier for foreign epitopes. Adv Virus Res 50: 141–182.

9. Mozdzanowska K, Furchner M, Gerhard W (1999) Treatment of hepatitis B virus infection. Drugs 59: 617–630.

10. Jegelkær J, Schmitz N, Hinton JJ, Schwarz K, Jegelkær A, et al. (2008) Efficient induction of mucosal and systemic immune responses in virus-like particles administered intranasally: implications for vaccine design. European journal of immunology 38: 1141–1146.

11. Proctor P, Stecher D, Oehninger C, Melber K, Reimann J, et al. (2002) Priming Th1 immunity to viral core particles is facilitated by trace amounts of RNA bound to its arginine-rich domain. J Immunol 168: 4951–4959.

12. Ferrari C, Bertolotti A, Penna A, Cavalli A, Valli A, et al. (1991) Identification of immunodominant T cell epitopes of the hepatitis B virus nucleocapsid antigen. J Virol 65: 5232–5241.

13. Robertson JS, Nicolson C, Harvey R, Johnson R, Major D, et al. (2011) The active oligomeric state of the hepatitis B virus core particle contains multiple epitopes of the Plasmodium falciparum circumsporozoite protein provides a highly immunogenic malaria vaccine in preclinical analyses in rodents and primate hosts. Infect Immun 70: 6680–6670.

14. Tseng SY, Wang TT, Tan GS, Hai R, Pica N, et al. (2010) Vaccination with a synthetic peptide from the influenza virus hemagglutinin provides protection against distinct viral subtypes. Proc Natl Acad Sci U S A 107: 18097–18098.

15. De Filette M, Min Jou W, Birkett A, Lyons K, Schultz B, et al. (2005) Universal vaccine based on the extracellular domain of M2: weak protection mediated via antibody-dependent NK cell activity. Journal of virology 79: 5258–5265.

16. Kawai T, Akira S (2006) TLR signaling. Cell death and differentiation 13: 816–826.

17. Ferrari C, Bertolotti A, Penna A, Cavalli A, Valli A, et al. (1991) Identification of immunodominant T cell epitopes of the hepatitis B virus nucleocapsid antigen. J Virol 65: 5232–5241.

18. Mozdzanowska K, Zharikova D, Cudic M, Otvos L, Gerhard W (2007) Roles of functional domains in core antigen and e antigen from hepatitis B virus. J Virol 81: 66: 5232–5241.

19. Brown DM, Shafer HC, Brooke AG, Reading PC (2011) Responses of mouse airway epithelial cells and alveolar macrophages to viral and avirulent strains of influenza A virus. Virology 410: 331–338.

20. Kawai T, Akira S (2006) TLR signaling. Cell death and differentiation 13: 816–826.

21. Mozdzanowska K, Zharikova D, Cudic M, Otvos L, Gerhard W (2007) Antibody restriction of virus growth and detection of M2 in virions. Journal of virology 81: 66: 5232–5241.

22. Ferrari C, Bertolotti A, Penna A, Cavalli A, Valli A, et al. (1991) Identification of immunodominant T cell epitopes of the hepatitis B virus nucleocapsid antigen. J Virol 65: 5232–5241.

23. Mozdzanowska K, Zharikova D, Cudic M, Otvos L, Gerhard W (2007) Antibody restriction of virus growth and detection of M2 in virions. Journal of virology 81: 66: 5232–5241.