A VLP-based vaccine targeting domain III of the West Nile virus E protein protects from lethal infection in mice

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Abstract

Background: Since its first appearance in the USA in 1999, West Nile virus (WNV) has spread in the Western hemisphere and continues to represent an important public health concern. In the absence of effective treatment, there is a medical need for the development of a safe and efficient vaccine. Live attenuated WNV vaccines have shown promise in preclinical and clinical studies but might carry inherent risks due to the possibility of reversion to more virulent forms. Subunit vaccines based on the large envelope (E) glycoprotein of WNV have therefore been explored as an alternative approach. Although these vaccines were shown to protect from disease in animal models, multiple injections and/or strong adjuvants were required to reach efficacy, underscoring the need for more immunogenic, yet safe DIII-based vaccines.

Results: We produced a conjugate vaccine against WNV consisting of recombinantly expressed domain III (DIII) of the E glycoprotein chemically cross-linked to virus-like particles derived from the recently discovered bacteriophage AP205. In contrast to isolated DIII protein, which required three administrations to induce detectable antibody titers in mice, high titers of DIII-specific antibodies were induced after a single injection of the conjugate vaccine. These antibodies were able to neutralize the virus in vitro and provided partial protection from a challenge with a lethal dose of WNV. Three injections of the vaccine induced high titers of virus-neutralizing antibodies, and completely protected mice from WNV infection.

Conclusions: The immunogenicity of DIII can be strongly enhanced by conjugation to virus-like particles of the bacteriophage AP205. The superior immunogenicity of the conjugate vaccine with respect to other DIII-based subunit vaccines, its anticipated favourable safety profile and low production costs highlight its potential as an efficacious and cost-effective prophylaxis against WNV.

Background

West Nile virus (WNV) is a positive-stranded RNA flavivirus grouped within the Japanese encephalitis virus serocomplex. Transmitted primarily between birds via Culex mosquitoes, it occasionally infects humans, where it usually remains asymptomatic or causes a mild undifferentiated febrile illness called West Nile fever. Under certain conditions, mainly in immunocompromised or elderly individuals, and in individuals deficient in expression of the chemokine receptor CCR5, WNV infection can develop into severe, potentially life-threatening encephalitis [1-4]. In 2002, WNV was responsible for the largest outbreak of arthropod-borne encephalitis recorded in the USA, accounting for 2946 diagnosed cases and 284 deaths [5]. Since then the virus has been spreading throughout the USA, as well as Canada, Mexico and the Caribbean basin [6]. Isolated clinical cases have also been reported in recent years in Mediterranean countries, suggesting emergence of the virus in Western Europe [7,8]. In the absence of an effective treatment, there is a medical need for the development of a safe and efficient prophylactic vaccine against WNV.

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A chimeric virus incorporating the envelope proteins of WNV into the infectious backbone of a yellow fever vaccine strain is currently being developed as a live-attenuated vaccine [9-11]. While immunogenic in humans, such a vaccine carries the inherent risk of reversion to a more virulent form, requiring stringent monitoring of the production process and careful safety assessment during clinical development. Alternative vaccination strategies are therefore focusing on recombinant subunit vaccines based on the large envelope glycoprotein (E) of WNV. The E protein is crucial for virus attachment and entry into host cells and is also the major antigen eliciting neutralizing antibody responses [12]. In particular a structurally distinct domain of the E protein (DIII) has been proposed as the receptor-binding domain [13]. Antibodies recognizing epitopes in this domain have been shown to neutralize the virus in vitro [14-19] and passive transfer of DIII-specific antibodies has been shown to protect mice from WNV challenge [19]. Subunit vaccines based on recombinantly expressed DIII have been tested in animal models and have proven effective in protecting from WNV infection [20-24]. However, multiple injections and/or strong adjuvants were needed to induce neutralizing antibody responses, indicating that isolated DIII is poorly immunogenic.

We have previously shown that by displaying antigens in a repetitive and highly ordered fashion on the surface of virus-like particles (VLPs) derived from the bacteriophage Qβ, specific B cells can be efficiently activated and rapid and robust antibody responses can be induced [25-28]. Here we describe the production of a conjugate vaccine based on recombinant DIII covalently linked to VLPs derived from the recently discovered bacteriophage AP205. A single injection of the conjugate vaccine was sufficient to induce virus-neutralizing antibodies and provide significant protection from WNV challenge, demonstrating its superior immunogenicity over previously described DIII-based vaccines and highlighting its potential as an efficient and safe prophylaxis against WNV.

Results

Production of the AP205 VLP Carrier

The bacteriophage AP205 has first been isolated from Acinetobacter spp. and belongs to the Leviviridae family, a group of ssRNA phages which infects a wide range of Gram-negative bacteria. Its positive-stranded RNA genome comprises three large open reading frames, which encode the maturation, coat and replicase proteins [29]. The icosahedral T = 3 phage particles are 29 nm in size and contain the genomic RNA in a densely packaged fashion [30]. In analogy to the coat proteins of the related phages Qβ [31] and MS2 [32], the AP205 coat protein self-assembles into virus-like particles upon recombinant expression in E. coli [33]. We purified AP205 VLPs from lysates of E. coli overexpressing the coat protein by sequential anion exchange and mixed-mode cation/anion exchange chromatography. Figure 1 shows that AP205 VLPs purified in this way were 25-30 nm in size and strongly resembled in their appearance the wild type particles [29]. Spectrophotometric analysis and agarose gel electrophoresis indicated that AP205 VLPs contained approximately 25-30 μg of host cell RNA per 100 μg of coat protein (not shown).

Production of the DIII-C-AP205 Conjugate Vaccine

Domain III of the WNV E protein was engineered to comprise a hexahistidine tag and a cysteine-containing linker at its C-terminus (DIII-C). Upon expression in E. coli the protein formed insoluble inclusion bodies. DIII-C was solubilized in urea, purified under denaturing conditions by affinity chromatography and refolded into its native form by stepwise dialysis. A heterobifunctional cross-linker was used to covalently conjugate DIII-C via the introduced C-terminal cysteine residue to lysine residues on the surface of AP205 VLPs. The isolated vaccine components and the conjugate vaccine (DIII-C-AP205) were analysed by SDS-PAGE and Western Blot, as well as size exclusion chromatography, with results shown in Figure 2. Recombinant DIII-C migrated as a distinct 13.5 kDa band in reducing, denaturing SDS-PAGE (Figure 2A, left panel), consistent with its predicted molecular weight. Size exclusion analysis of the protein showed a single peak with an elution volume corresponding to an

Figure 1 Electron micrograph of purified AP205 VLPs

Purified AP205 VLPs were adsorbed onto carbon-Formvar-coated grids, stained with 2% phosphotungstic acid and subjected to transmission electron microscopy.
**Figure 2** Production and characterization of the DIII-C-AP205 conjugate vaccine. 

**A** Derivatized AP205 (dAP205), DIII-C, and the dialysed conjugate vaccine (DIII-C-AP205) were analyzed by reducing, denaturing SDS-PAGE (left panel). Corresponding amounts of derivatized AP205, DIII-C, and of the non-dialysed (nd) conjugate vaccine were also analysed by non-reducing, non-denaturing SDS-PAGE (right panel). For identification of the coupling bands, proteins were separated on reducing, denaturing SDS-PAGE, blotted on nitrocellulose and detected with AP205- and His-tag-specific antibodies (middle panels). Bands corresponding to AP205-crosslinked DIII-C are indicated by arrows. The 27 kDa band which is visible in the SDS-PAGE and in the His-tag-specific Western Blot corresponds to dimeric DIII-C (*). 

**B** Size exclusion chromatography. A superdex 75 column was calibrated with a molecular weight (MW) calibration kit and then loaded sequentially with the indicated proteins. AP205 VLPs and the conjugate vaccine DIII-C-AP205 elute in the void volume of the column ($V_0$, 40 ml) while purified DIII-C elutes at 72.4 ml.
apparent molecular weight of approximately 16 kDa, indicating that DIII-C was indeed a folded monomer in solution (Figure 2B). Analysis of the conjugate vaccine DIII-C-AP205 showed the presence of several high molecular weight bands in reducing, denaturing SDS-PAGE, which reacted with both AP205- and His-tag specific antisera, demonstrating the successful crosslinking of DIII-C to the AP205 VLPs (Figure 2A, middle panels). Similar to AP205 VLPs, the DIII-C-AP205 conjugate vaccine eluted in the void volume of the size exclusion column, confirming its virus-like particle assembly state (Figure 2B). In order to estimate the number of DIII-C molecules displayed by each VLP in the conjugate vaccine, the corresponding amounts of DIII-C used in the coupling reaction and of the non-dialysed DIII-C-AP205 conjugate vaccine were loaded side by side on a non-reducing, non-denaturing SDS polyacrylamide gel (Figure 2A, right panel). Under these conditions, DIII-C still migrates as a 13.5 kDa monomer while the conjugate vaccine migrates as a high molecular weight complex. Densitometric analysis showed a 55% decrease in the amount of free DIII-C after coupling to AP205. Assuming that the decrease in free DIII-C is due to coupling to AP205 subunits and taking into account the molar ratio of VLP subunits and DIII-C molecules used in the coupling reaction it could be calculated that an average 27% of AP205 subunits had been cross-linked to DIII-C molecules. As each AP205 VLP is composed of 180 subunits, it can be estimated that an average of 50 DIII-C molecules are displayed per vaccine particle.

**Immunogenicity of DIII-C-AP205 in Mice**

Groups of mice were immunized subcutaneously either with the DIII-C-AP205 conjugate vaccine or a mixture of the corresponding amounts of non-conjugated AP205 carrier and free DIII-C protein in the absence of any additional adjuvant. Figure 3 shows that one or two injections of the non-conjugated AP205/DIII-C mixture did not result in any detectable DIII-specific IgG antibodies, confirming the notion that isolated DIII is poorly immunogenic. A measurable IgG antibody response could be induced only after a third injection (ELISA titer of 15,300 on day 42). In contrast, DIII-specific IgG titers were measured after only a single administration of the DIII-C-AP205 conjugate vaccine (6,600 on day 14). A second vaccine injection boosted the specific antibodies to a titer of 106,900 (day 28), while a third injection did not lead to a further increase. In the absence of additional injections, antibody titers slowly declined with an approximate half life of two months.

**Vaccination with DIII-C-AP205 Induces Neutralizing Antibodies and Protects from Lethal WNV Infection**

Groups of mice were immunized with DIII-C-AP205 three times (days 0, 14, 28) either in the absence or in the presence of Alum as adjuvant. One group was immunized only once (day 28) in the presence of Alum. The neutralizing capacity of the induced antibodies was determined two weeks after the last injection (day 42). Figure 4A shows that three injections of DIII-C-AP205 in the absence or presence of adjuvant resulted in average neutralizing titers of 360 and 760, respectively. Neutralizing ability was also detected in sera from 6 of the 8 mice that had received a single injection of vaccine. All vaccinated mice and a group of control mice, that had been immunized with the AP205 carrier alone, were then challenged with a lethal dose of West Nile virus strain NY99. Figure 4B shows that all mice that had received three injections of DIII-C-AP205 either in the presence or absence of Alum survived the viral challenge. In contrast, all control mice immunized with AP205 succumbed to the infection. Interestingly, 5 out of 8 mice that had received a single injection of DIII-C-AP205 in Alum also survived the challenge with WNV. This survival rate is in the range of the ones achieved after single doses of different live attenuated WNV vaccines in mice [9,34,35]. Vaccination with DIII-C-AP205 resulted in a marked reduction of viremia as measured by real time PCR in blood 3 and 7 days after infection (Figures 4C and 4D, respectively). All mice that had received three injections of DIII-C-AP205, as well as 6 out of 8 mice that had received a single injection had cleared the virus by day 7; the 2 remaining mice of this group had reduced viral titers as compared to the average viral load of AP205-immunized animals at this time point.

**Discussion**

Domain III of the envelope protein of WNV is a major target of virus-neutralizing antibody responses and has been identified as a promising candidate antigen for the development of recombinant subunit vaccines [36]. In this study we produced a highly immunogenic and efficacious WNV vaccine consisting of recombinant domain III chemically cross-linked to virus-like particles of the bacteriophage AP205. The conjugate vaccine produced high DIII-specific antibody titers in mice, which were able to efficiently inhibit viral replication both in vitro and in vivo. In contrast to other experimental vaccines based on recombinant DIII and comprising different adjuvants [20-23], a single injection in the presence of Alum was sufficient to induce neutralizing antibody responses and confer partial protection from WNV challenge (Figure 4). Interestingly, the neutralizing titers observed (approximately 30-40) were in the range of those induced by a single injection of live WNV vaccines such as a chimeric attenuated flavivirus vaccine [9,35] or a recombinant attenuated influenza strain expressing the WNV E protein [34]. Multiple injections either in the presence or absence of Alum as adjuvant yielded sustained high titers of DIII-specific antibodies, which efficiently neutralized
the virus. The reason for the superior immunogenicity of the DIII-C-AP205 conjugate vaccine most likely resides in its virus-like connotations. It has been shown that highly ordered and repetitive antigen arrays can cause an efficient cross-linking of BCRs on specific B cells and induce a rapid and sustained antibody response [37]. As domain III is presented to the immune system in an oriented and densely packaged fashion on the surface of the AP205 VLP, it is likely that DIII-specific B cells are promptly and efficiently activated to produce specific IgG antibodies. The particulate nature of the VLP vaccine furthermore ensures a preferential uptake by antigen-presenting cells such as dendritic cells, and thereby an efficient presentation of DIII- as well as AP205-derived epitopes on MHC class II for the priming of specific TH cells. Activation of antigen-presenting cells is also enhanced by bacterial RNA, which is spontaneously packaged into the VLP carrier during the recombinant expression and assembly process. Upon uptake by B cells and APCs, the RNA is co-delivered with the AP205 particle to the endosomal compartment, where it can activate TLR3 or TLR7/8 (for review see [38,39]).

In addition to its good immunogenicity, the DIII-C-AP205 vaccine is also expected to be safe and well tolerated. In contrast to live vaccines based on attenuated viruses, which inevitably carry the risk of genetic recombination and mutation into a more virulent form, DIII-C-AP205 is based on non-replicating virus-like particles derived from a bacteriophage, which are unable to infect mammalian cells. Moreover, both the VLP carrier and the antigen components of the vaccine can be produced in large amounts in bacterial expression systems and purified with relatively simple biochemical methods, suggesting that large scale production of the conjugate vaccine can be achieved in a cost-effective manner. A highly immunogenic, yet safe and affordable WNV vaccine would be attractive for veterinary prophylaxis and might also be used in elderly or immunocompromised individuals in high-risk areas. The high immunogenicity of the VLP vaccine might also offer the potential of inducing cross-protection against related flaviviruses such as Japanese encephalitis virus or dengue virus. That cross-protection may occur in principle has been shown by immunization of mice with recombinant domain III of the WNV E protein [21]. The increased immunogenicity of the domain III by conjugation to the VLP carrier may therefore be sufficient to confer cross-protective immunity.

Conclusions

In the present study we show that the immunogenicity of DIII of the WNV E protein can be strongly enhanced by conjugation to virus-like particles of the bacteriophage AP205. In contrast to other vaccination approaches based on recombinant DIII, which require multiple injections and/or strong adjuvants for the induction of neutralizing antibodies, a single injection of the conjugate DIII-C-AP205 vaccine in Alum was sufficient to induce a signifi-
Figure 4: Immunization with DIII-C-AP205 protects from lethal WNV infection. 

A. Induction of neutralizing antibodies. Groups of female C57BL/6 mice (n = 8) were immunized subcutaneously on days 0, 14 and 28 with 50 μg DIII-C-AP205 either in the absence or presence of Alum as adjuvant or with 50 μg AP205 VLPs in the absence of adjuvant. A fourth group was immunized once with 50 μg DIII-C-AP205 in Alum on day 28. Virus-neutralizing titers of individual sera were measured on day 42. The dashed line indicates the detection limit. Shown are individual titers and group means.

B. Protection from WNV challenge. Two weeks after the last vaccine injection mice were challenged with a lethal dose of WNV (arrow). Statistical significance of differences in survival curves was calculated by log-rank test using GraphPad-Prism (***p < 0.001 vs. 3xAP205 control). p.i. = post infection.

C and D. Viral titers were determined 3 and 7 days post infection (p.i.) in blood of infected animals. Shown are individual titers and group means. The Mann-Whitney test was used to assess statistical significance (**p < 0.01, ***p < 0.001 vs. 3xAP205 control).
significant amount of virus-neutralizing antibodies in mice. Three injections of the vaccine completely protected mice from a lethal WNV challenge, even when given in the absence of any adjuvant. The relatively low production costs of the DIII-C-AP205 vaccine, its superior immunogenicity with respect to other DIII-based approaches and its anticipated good safety profile make it an attractive candidate for WNV prophylaxis both in humans and in veterinary applications.

**Methods**

**Expression and Purification of AP205 Virus-like Particles**

Cleared bacterial lysates containing the recombinantly expressed coat protein of AP205 were dialysed against AEX loading buffer (20 mM NaH₂PO₄, pH 7.2) and loaded on a Fractogel™ TMAE column (Merck). After removal of host cell proteins by a salt wash with 333 mM NaCl, viral capsids were eluted with 600 mM NaCl and dialyzed against HAp loading buffer (5 mM NaH₂PO₄, 100 mM NaCl, pH 6.8). Capsids were bound to a hydroxyapatite column (Macro-prep ceramic hydroxyapatite type II, Bio-rad) and eluted with 60 mM NaH₂PO₄, 220 mM NaCl, pH 6.8, resulting in depletion of bacterial LPS.

**Expression and Purification of Recombinant Domain III of the E Glycoprotein of WNV**

A DNA fragment encoding domain III of the glycoprotein E of WNV NY99 was amplified from plasmid pTRHis2A-WNV-E [21] with the oligonucleotide pair WNV1/ WNV2 (5'-ATATATCATATGAAATTCAGTTGAAGG-3'; 5'-ATATATCTCGAGTTGCAATGCTG C TTCCAG-3'; Ndel and Xhol restriction sites are in bold) and cloned into the expression vector pET42T [40]. The resulting plasmid encoded a fusion protein consisting of domain III of the WNV E protein (corresponding to amino acids 582-696 of the WNV polyprotein precursor), a hexahistidine tag, and a short C-terminal, cysteine containing linker (DIII-C). E. coli BL21 DE3 cells were transformed with this plasmid, and protein expression was induced in a logarithmic phase culture by addition of isopropyl-β-D-thiogalactopyranoside to a final concentration of 1 mM. After overnight growth bacteria were harvested by centrifugation, resuspended in 50 mM NaH₂PO₄, 150 mM NaCl, 0.25% Triton X-100, pH 7.2, and lysed by sonication. Nucleic acids were digested by 1 h incubation at room temperature with 1500 U Benzonase (Sigma-Aldrich), and inclusion bodies were solubilized in 8 M urea, 100 mM Tris-Cl, 100 mM DTT, pH 8.0, and loaded on a Ni-NTA column (Qiagen), which had been previously equilibrated with 8 M Urea, 100 mM NaH₂PO₄, 10 mM Tris-Cl, 2 mM β-mercaptoethanol, pH 8.0. Bound DIII-C was eluted with 8 M urea, 100 mM NaH₂PO₄, 10 mM Tris, 2 mM β-Mercaptoethanol pH 4.5, and dialysed against 2 M urea, 50 mM NaH₂PO₄, 0.5 M arginine, 0.5 mM oxidized glutathione, 5 mM reduced glutathione, 10% glycerol, pH 8.5. DIII-C was then refolded by stepwise dialysis against 50 mM NaH₂PO₄, 0.5 M arginine, 0.5 mM oxidized glutathione, 5 mM reduced glutathione, 10% glycerol, pH 8.5, and against 50 mM NaH₂PO₄, 10% glycerol, pH 8.5.

**Chemical Cross-linking of Recombinant DIII-C to AP205 Virus-like Particles**

AP205 VLPs (in PBS, pH 7.2) were first reacted for 1 h at room temperature with a 2.5 fold molar excess of the heterobifunctional cross-linker succinimidyl-6-(β-maleimidopropionamido)hexanoate (Pierce). Free cross-linker was removed by dialysis against PBS, pH 7.2. Recombinant DIII-C was incubated for 1 h at room temperature with an equimolar amount of tri(2-carboxyethyl)phosphine-hydrochloride. Under these mildly reducing conditions the cysteine residue contained in the linker is reduced, while the internal disulfide bridge of DIII-C remains intact. The reduced protein was then mixed with the derivatized AP205 VLPs at a molar ratio of 1 DIII-C monomer per 2 AP205 monomers and incubated over night at 17°C to allow cross-linking. Free DIII-C was removed by extensive dialysis against PBS pH 7.2 using cellulose ester membranes with a cut-off of 100 kDa (Spectrum Laboratories). The conjugate vaccine was analyzed by SDS-PAGE followed by Coomassie Blue staining or by Western Blot using AP205- and His-tag- specific antisera. The molecular masses of the DIII-C and AP205 monomers are similar; 13.5 kDa and 14.0 kDa, respectively. The coupling product comprising one AP205 monomer covalently conjugated to one DIII-C monomer co-migrates with the AP205 dimer band. Hence the coupling efficiency could not simply be calculated by densitometry of protein bands on a reducing SDS-PAGE stained with Coomassie Blue. Instead the conjugate vaccine was loaded on a non-reducing non-denaturing SDS-PAGE side by side with the corresponding amount of free DIII-C, which had been used in the cross-linking reaction. By comparing the intensities of the DIII-C monomers before and after cross-linking to AP205 by densitometry, the amount of DIII-C coupled to the AP205 carrier could then be quantified.

**Analysis of DIII-C, AP205 and DIII-C-AP205 by Size Exclusion Chromatography**

A superdex 75 column (GE Healthcare) was calibrated with a mixture of Dextran Blue (~2000 kDa), BSA (67 kDa), Ovalbumin (43 kDa), Chymotrypsinogen (25 kDa), and RNase A (14 kDa). AP205 VLPs, purified DIII-C pro-
tein and the conjugate vaccine DIII-C-AP205 were then sequentially analysed on the same column. The apparent molecular weight of DIII-C was calculated from a standard curve obtained by plotting the logarithm of the molecular weights of the protein standards against their partition coefficients.

**Immunogenicity of DIII-C-AP205**

Female BALB/c mice (8 weeks of age) were purchased from Charles River Laboratories. DIII-C-AP205 vaccine or the mixture of the non-conjugated vaccine components AP205 and recombinant DIII-C were diluted in PBS to 200 μl and injected subcutaneously (100 μl on two ventral sites) in the absence of additional adjuvants. Sera from immunized mice were serially diluted in PBS containing 0.05% Tween-20, 2% BSA, and applied to ELISA plates (Nunc) that had been coated with 1 μg/ml recombinant DIII-C protein. Reactivity of serum antibodies with the target protein was determined using a HRP-conjugated goat anti-mouse IgG secondary antibody (Jackson ImmunoResearch Laboratories) at a dilution of 1:1000 in PBS/0.05% Tween-20/2% BSA. After development with 1,2-phenylenediamine dihydrochloride (0.4 mg/mL in 0.066 M Na2HPO4, 0.035 M citric acid, 0.01% H2O2, pH 5.0) the optical density at 450 nm (OD450 nm) was determined using an ELISA reader (Biorad). Titers were expressed as the reciprocal of those serum dilutions that lead to half-maximal OD450 nm (OD50%).

**WNV Challenge**

Female C57BL/6 mice (6 weeks of age) were purchased from Harlan and allowed to acclimate to the facility for one week before experiments were performed. Experiments were approved by the animal ethics committee of the Erasmus MC Rotterdam, The Netherlands. Mice were immunized as indicated in the legend of Figure 4 and one week before experiments were performed. Experiments on vaccine analytics, coordinated the study and drafted the manuscript. GTJ conceived and designed the project. IU performed immunization and ELISA experiments. MB purified recombinant DIII-C, and produced the conjugate vaccine. PP performed the electron microscopy experiments. ADMEO and MFB conceived and designed the project. All authors read and approved the final manuscript.

**Acknowledgements**

Part of this work has been supported by a grant of the European Community (contract LSHB-CT-2004-005246 “COMPUVAC”). The authors would like to thank Alexander Link for critical reading of the manuscript.

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**Published: 6 July 2010**

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