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Adjuvant-like Effect of Vaccinia Virus 14K Protein: A Case Study with Malaria Vaccine Based on the Circumsporozoite Protein

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Development of subunit vaccines for malaria that elicit a strong, long-term memory response is an intensive area of research, with the focus on improving the immunogenicity of a circumsporozoite (CS) protein-based vaccine. In this study, we found that a chimeric protein, formed by fusing vaccinia virus protein 14K (A27) to the CS of Plasmodium yoelii, induces strong effector memory CD8+ T cells responses in addition to high-affinity Abs when used as a priming agent in the absence of any adjuvant, followed by an attenuated vaccinia virus boost expressing CS in murine models. Moreover, priming with the chimeric protein improved the magnitude and polyfunctionality of cytokine-secreting CD8+ T cells. This fusion protein formed oligomers/aggregates that led to activation of STAT-1 and IFN regulatory factor-3 in human macrophages, indicating a type I IFN response, resulting in NO, IL-12, and IL-6 induction. Furthermore, this vaccination regimen inhibited the liver stage development of the parasite, resulting in sterile protection. In summary, we propose a novel approach in designing CS based pre-erythrocytic vaccines against Plasmodium using the adjuvant-like effect of the immunogenic vaccinia virus protein 14K.

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Malaria continues to present a major public health challenge and burden on economic development in many countries. Plasmodium falciparum, the main causative agent of malaria in humans, is known to cause ~225 million cases and ~781,000 deaths annually (1). Development of a cost-effective vaccine continues to be a daunting task because of the confounding ability of the parasite to manipulate the host immune system. The most advanced malaria vaccines have focused on the use of protein-in-adjuvant formulations, and a phase III clinical trial with RTS,S/AS01E (a liposome-based adjuvant system) is under way (2). First results of the phase II trial of RTS, S/AS01E in Africa revealed vaccine-reduced clinical episodes of malaria and severe malaria by about half during 12 mo after vaccination in children 5–7 mo age (3). The rationale behind the development of RTS,S is its ability to form virus-like particles, which enhances the immunogenicity of monomeric circumsporozoite (CS) protein. The ability of protein aggregates to stimulate strong immune responses compared with their monomeric counterparts has been studied for many decades and is supported by a number of recent studies (4–6).

CS protein is a monomeric protein of ~40–60 kDa covering the surface of infective sporozoites (7). However, the native CS from sporozoites is also known to form some aggregates that may facilitate the binding of CS to hepatocytes (8). The CS protein is GPI-anchored and consists of a central portion of immunodominant repeat regions of B and T cell epitopes (9). Reports of successful vaccination against malaria in humans have been attributed to the development of CS-specific humoral and cell-mediated immune responses (10). Therefore, the strong immunogenic nature of the CS protein and its presence during pre-erythrocytic and liver stages (11) make it a promising candidate for the development of a malaria vaccine.

Control of infection for most pathogens requires different strategies of intervention, including humoral or cell-mediated immune responses at different times in their life cycle. Therefore, an effective vaccination regimen should encompass these aspects to develop sterile immunity. Viral vaccine vectors expressing CS protein and other Plasmodium Ags as promising vaccine candidates have been previously described (12, 13). Many of these studies have focused on using DNA as priming agent, which met with success rates as high as 70–100% in mouse models, even though it was short-lived (14, 15). However, DNA-based vaccines have recently come under renewed scrutiny for a number safety factors such as integration of the DNA into host genome (16) and also by their induction of tolerance against the Ag (17). Another disadvantage is the reduced capacity of DNA vaccines to induce a strong Ab response. The quality of immune responses generated by DNA vaccines administered alone is comparatively weaker to the traditional vaccines such as subunit vaccines or attenuated organisms (18). Currently, most licensed vaccines are based on attenuated whole pathogens or subunit vaccines or attenuated organisms (18).
particles. Therefore, as an alternative to DNA vectors, the use of proteins as priming agents is gaining acceptance due to their better efficacy and safety.

In this study, we describe the development of a novel CS-based chimeric protein, which when combined in a heterologous prime−boost vaccine regimen with an attenuated vaccinia virus vector-induced enhanced immune responses. Additionally, this vaccine regimen gave sterile protection in mice when challenged with sporozoites. Our chimeric CS protein (CS14K) makes use of the oligomerization domain of the A27L gene of vaccinia virus. Our previous studies helped in the understanding of the structural organization of the 14-kDa protein encoded by the A27L gene (19). Considering the drawbacks of using a DNA priming strategy and CS protein encoding its C-terminal GPI sequence, we sought to determine the possible differences in the quality and quantity of humoral and T cell responses induced by different vaccine constructs and define how this might influence protection.

We also show how the differences in the induction of CD8+ T cells and Abs could influence the protective capacity of malaria vaccines. In agreement with previous studies with fusion protein (4, 20), we found that priming with CS14K resulted in the development of high avidity and balanced IgG Ab production. An important aspect of this work is the extensive analysis of quantity and quality of CD8+ T cells during primary and memory stages. Comparing our vaccine regimen to a protective vaccination regimen based on DNA priming followed by poxvirus boost, we showed that priming with CS14K followed by a modified vaccinia virus Ankara (MVA)-CS boost triggered strong memory poly-functional effector T cell responses, an important component for providing long-term protection (21). These results demonstrate that by fusing the vaccinia virus 14K (A27 protein) to CS we achieved a significant improvement in the protection of mice against liver stage development of (A27 protein) to CS we achieved a significant improvement in the provision of long-term protection (21).

Materials and Methods

Cells and viruses

Construction of recombinant MVA virus expressing circumsporozoite protein of Plasmodium yoelii 17XNL strain (MVA-CS) has been described previously (13). The virus was grown and titrated in primary chicken embryo fibroblast cells and in DF-1 cells (22). Murine J774 and human THP-1 macrophages used for the in vitro cell culture studies were maintained at appropriate conditions as specified by the American Type Culture Collection.

Plasmid construction

pCl-Neo-CS. The pCSP gene was amplified MVA-CS using the primers CS-XhoI-F (5′-CTGGAGCTGGAGACTGAGATGTGTTACAATGAAGAAAATG-3′) and CS-NotI-R (5′-ATTTCGCGCCGGCTTAAATTATATCTTGGAC-3′) to yield a 972-bp fragment lacking the N-terminal signal sequence and C-terminal GPI sequence. The gene was inserted into a mammalian expression vector, pCl-Neo, that had been previously digested with XhoI and NotI followed by ligation. The fusion gene was amplified by digesting with NotI followed by ligation. The fusion gene fragment was then inserted into pGEX-6p-1 plasmid to produce pGEX-CS14K plasmid.

Recombinant protein constructs and protein isolation

The recombinant proteins were purified from Escherichia coli strain DH5α. The starter culture was diluted 1:100 in fresh Luria-Bertani media and allowed to grow at 37°C until the OD600 reached 0.7, following which isopropyl-β-D-thiogalactoside was added to a final concentration of 1 mM. The culture was then incubated in an orbital shaker at 18°C and 200 rpm for 24 h. After the incubation, the cells were harvested and the pellet was suspended in extraction buffer (50 mM Tris-HCl [pH 7.5], 250 mM NaCl, 1 mM EDTA and protease inhibitor tablets; Roche). The cells were then lysed with lysozyme, added to a final concentration of 1 mg/ml, and incubated on ice for 20 min. Following lysozyme treatment, 1% Sarkosyl detergent and 1% Triton X-100 was added and incubated at 37°C for 10 min. Clarified supernatant from the lysate was incubated with glutathione-Sepharose 4B beads at 4°C overnight. The beads were then washed using three washes with wash buffer I (extraction buffer with 0.5% Triton X-114) and three with wash buffer II (extraction buffer with 0.1% Triton X-114) followed by two washes with extraction buffer. The purified protein was then eluted with 20 mM reduced glutathione. The protein was desalted using an Amicon centrifugal concentrator and the protein concentration was determined using Bradford reagent. The GST tag from the protein was cleaved using PreScission protease (GE Healthcare) according to the manufacturer’s protocol. The proteins were tested for LPS contamination using a chromogenic Limulus amebocyte lysate kit (QCL-1000; Lonza), which was maintained at <2 endotoxin units per microgram of protein.

Neutralization assay

Sera from immunized rabbit were inactivated at 56°C for 30 min, following which 2-fold serial dilutions of the serum were made and incubated with 200 PFU MVA at 37°C for 1 h. Afterwards, confluent DF-1 cells were infected in triplicate and were visualized by immunostaining with anti-WR serum after 48 h. As a control, nonimmune serum from nonimmunized animals was used. The number of plaques obtained from each serum dilution was normalized to this control value.

Nitrite measurement

NO synthesis was measured by estimating the levels of nitrite present in the supernatant. Briefly, J774 cells were stimulated with proteins for 24 h. In vivo nitrite analysis, 103 splenocytes from vaccinated animals, sacrificed 53 d after boost, were treated with 5 μg/ml CS protein. Nitrite accumulation was measured by treating 50 μl supernatant with 50 μl Griess reagent I (1% sulfanilamide solution in 2.4 N HCl) for 10 min in dark followed by the addition of 50 μl Griess reagent II (0.1% naphthalylenediamine in 2.4 N HCl) for 10 min. The assay was read by a spectrophotometer at 540 nm.

RNA extraction and RT-PCR

Total RNA was extracted from cells treated with respective proteins for 24 h, using a RNaseasy Mini kit according to the manufacturer’s instructions. Analysis of RNA was carried out using RT-PCR as described in the literature (23). cDNA was used as a template with primers specific for inducible NO synthase, IL-12p40, and GAPDH (23). All the experiments were done in triplicates and the bands of gel electrophoresis were quantified using Adobe Photoshop CS4.

Confoical microscopy

Immunostaining was carried out as described previously (24). Briefly, after fixation (30 min; 4% formaldehyde in PBS, 37°C), permeabilization in 0.1% Triton X-100 (Sigma-Aldrich), and blocking with 10% FCS in PBS, cells were incubated with primary Ab (anti-CS Ab, 1:500; CS-anti-14K, 1:400; NF-Bp65, 1:500) along with DNA staining dye, DAPI (1:200), for 1 h at room temperature. Following extensive washing with PBS secondary Abs (Alexa 546 goat anti-mouse and Alexa 488 goat anti-rabbit, 1:500) were applied for 1 h at room temperature. The slides were washed three
times with PBS and mounted in ProLong antifade medium and analyzed with a Bio-Rad Radiance 2100 confocal laser microscope.

**Animals and immunizations**

All animal procedures were approved by the Ethical Committee of Animal Experimentation of Centro Nacional de Biotecnologia (Consejo Superior de Investigaciones Científicas). Female BALB/C mice (H-2b), 6–8 wk old, were obtained from Harlan U.K. A standard immunization protocol based on a heterologous prime–boost approach designed in the laboratory was followed (25). In short, animals were primed with DNA (100 μg; DNA-CS or empty DNA-d) or protein (20 μg; CS or CS-14K) via intradermal route and were boosted after 2 wk with 2 × 10^6 PFU respective sucrose-purified viruses (MVA or MVA-CS) through i.p. injection. All the preparations were made in endotoxin-free PBS.

**P. yoelii sporozoite challenge study**

Challenge experiments were performed as previously described (26). Briefly, sporozoites were obtained from the salivary glands of *Anopheles stephensi* mosquitoes. Two weeks after boost, mice were challenged with 2 × 10^6 sporozoites via i.v. route through the tail vein. After 42 h, animals were sacrificed and levels of *P. yoelii* 18S rRNA levels were assessed by quantitative RT-PCR. To determine sterile protection 2 wk after immunization, mice were challenged i.v. with 300 *P. yoelii* sporozoites. Parasitemia was monitored by performing daily blood smears from days 3 to 21.

**ELISA and Ab avidity**

Abs present in the serum of immunized animals were determined using ELISA as previously described (25). Purified CS protein was coated to the 96-well Nunc MaxiSorp plates at a concentration of 2 μg/ml in coating buffer (NaHCO₃/Na₂CO₃) at 4°C overnight. Bound Abs were detected using 1:2000 dilution of alkaline phosphate-conjugated goat anti-mouse Ab total IgG or IgG1 or IgG2a (SouthernBiotech, Birmingham, AL). Plates were developed by adding TMB substrate (Sigma-Aldrich) and stopping the reaction with 1 M H₂SO₄. OD was read at 450 nm. Endpoint titer values were determined as the last positive dilution of serum giving an absorbance value 3-fold higher than the serum. For analyzing the avidity of Abs, an initial serum dilution that gave an absorbance of 2.7 ELISA units was selected. Following incubation, with serum, the Ag/Ab interaction was disrupted using a range of dilutions from 0 to 5 M urea (Invitrogen) in Tris-HCl (pH 8.0) for 15 min before the addition of secondary Ab. EC₅₀ was then calculated by linear regression (R² = 0.99) between 1 and 5 M urea concentration (log of 50% reduction, 1.699).

**IFN-γ ELISPOT analysis**

Fresh IFN-γ ELISPOT analysis was carried out as previously described (27).

**Multiparameter flow cytometry**

Flow cytometry and intracellular cytokine staining analysis were performed to detect the different phenotype of lymphocyte secreting cytokines, as reported previously (25). Briefly, 4 × 10^5 splenocytes were stimulated with 1 μg/ml CDS1 peptide, PyCS260-288 (SYVPSAEQI), with GolgiPlug (BD Biosciences) for 6 h in a 96-well plate. The cells were then washed and Fc receptors were blocked using anti-CD16/CD32, following which the cells were stained with surface-specific mouse Abs, namely CD4-Alexa 700, CD8-PerCP, CD3-FITC, and CD62L-PE for memory studies. Cells were permeabilized using the BD Cytofix/Cytoperm kit (BD Biosciences) and were stained for intracellular cytokines IFN-γ/alkaline phosphocyanin, IL-2/PE, and TNF-α/PECy7 for adaptive response or with IFN-γ/PECy7, IL-2/allophycocyanin, and TNF-α/PE for memory response. A million cells were then passed through an LSRII flow cytometer (BD Biosciences) and the data were analyzed with FlowJo (Tree Star) and Spicpe (version 5.0). Appropriate controls were used and the values from unstimulated samples were subtracted.

**Statistical Analysis**

Statistical analysis was performed using Minitab for Windows. For ELISA, to determine the differences between groups we performed a linear regression of the logarithmic absorbance versus the logarithmic dilution, removing those samples clearly not following a linear trend (R² > 0.98). We then computed the logarithmic dilution at which this regression line was crossing the threshold given by twice the absorbance of the control values for that mouse. Doing this for the four mice we have four estimates of the logarithmic dilution beyond which the absorbance is smaller than twice the absorbance given by the control. We will refer to this value as the critical logarithmic dilution. Finally, we compared with a Student t test the hypothesis that the critical logarithmic dilution between the groups. For intracellular cytokine staining and ELISPOT, statistical analysis was done based on a previously described method (25). Briefly, we developed a novel method for analysis by correcting the control values (RPMI 1640) for determining standard deviations and p values for the samples.

**Results**

**14K fusion aids CS protein aggregation**

To investigate the effects of 14K conjugation with CS, we determined the structural organization of CS protein and its reactivity against Abs generated from *P. yoelii* sporozoites. With the aim of improving the immune response against CS, we removed the GPI motif (28) and signal sequence in the CS and fused it with the 14K protein after deleting the first 28 aa from A27, which is responsible for producing neutralizing Abs against vaccinia virus (Supplemental Fig. 1A). Abs generated against CS14K did not neutralize the infectivity of MVA as seen by neutralization assay (30–36 spots/well) (Supplemental Fig. 1B). Using SDS-PAGE analysis under reducing and nonreducing conditions, we evaluated the oligomerization/aggregation status of recombinant proteins and their reactivity with a CS-specific mAb obtained after immunization with sporozoites (Fig. 1A, 1B). Under reducing conditions CS14K has a molecular mass of ~60 kDa, compared with ~50 kDa for CS, whereas under nonreducing conditions CS14K has a size apparently >250 kDa, in contrast to 50 kDa for CS. Additional experiments by analytical ultracentrifugation revealed that CS14K protein has a main sedimenting species with a standard s value of 4.3 ± 0.1, which is compatible with a moderately elongated protein dimer (calculated frictional ratio f/f₀ = 1.8). The CS14K has high tendency to form oligomers/aggregates (data not shown). To determine the kinetics of CS expression by the vector MVA-CS, we performed time-course analysis of cells infected with the MVA recombinant expressing CS (Fig. 1C). We observed that after infection with MVA-CS, CS was detectable 2 h postinfection, and these expression levels remained elevated up to 6 h postinfection. Next, to determine whether CS expressed by MVA undergoes posttranslational modifications, we incubated infected cells in the presence of tunicamycin and observed that most of the CS expressed in the virus-infected cells was glycosylated, as indicated by the reduced intensity of the protein bands (Fig. 1D). We also investigated whether there was any difference in the localization of proteins in macrophages expressed by virus or upon transient transfection. To study the subcellular localization of full-length CS during infection with MVA-CS, J774 cells were infected at a multiplicity of infection of 5 PFU/cell for 18 h. We observed that MVA-CS expresses CS as punctuated spots with complete cytoplasmic spread (Supplemental Fig. 1C). In contrast, when macrophages were transfected with DNA encoding CS and CS14K, we observed that CS and CS14K were strongly localized with the nuclear envelope of the cell.

Taken together, these data show that the 14K (A27L) protein of vaccinia virus when fused to CS protein of *Plasmodium* is able to form oligomers/aggregates displaying an apparent molecular mass of >250 kDa compared with its monomeric CS counterpart of 50 kDa. A moderately elongated protein forming oligomers/aggregates was observed when CS14K protein was run under nonreducing conditions by analytical ultracentrifugation. The lower band that we see in the immunoblot under reducing condition is the cleaved CS14K protein. We therefore sought to investigate
a possible biological effect on the immunogenicity of CS protein when fused with A27.

**CS14K protein modulates innate immune responses in macrophages**

Fusion of Ags to immunomodulatory fragments such as TLRs (5) or to complement proteins (29), which is aimed at improving immunogenicity by aggregation of proteins or by exploiting the innate immune signaling of macrophages, is an intense area of research. Because fusion of 14K protein to CS facilitates its aggregation, we sought to investigate whether this could also modulate the innate immune responses in macrophages.

Ability of macrophages to induce NO to inhibit the growth of parasite has been well documented (30); therefore, we sought to evaluate the ability of the fusion protein to induce NO production. When comparing the NO levels in supernatant of J774 cells treated with fusion protein or normal protein, it was apparent that only CS14K resulted in NO production (Fig. 2A). However, when the cells were stimulated with proteins along with recombinant IFN-γ, elevated levels of NO were found in all samples, regardless of the presence of additional proteins (data not shown). Further analysis revealed a significant increase in NO (p = 0.044) and IL-12p40 (p = 0.05) mRNA levels in CS14K-treated macrophages compared with CS protein treatment (Fig. 2B). We next investigated the effects of proteins on IL-6 and TNF-α secretion, given their possible role in controlling pre-erythrocytic stages of malaria. Chimeric protein was able to induce a significant increase in IL-6 in macrophages (Fig. 2C). Although not statistically significant, we also observed increased TNF-α production at 24 h with CS14K treatment.

Given the significance of TLR activation and the induction of type I IFN, experiments focused on the analysis of the downstream molecules involved in IFN signaling such as IFN regulatory factor (IRF)-3 and STAT-1 were performed. To determine whether there are differences between CS and CS14K, human macrophage THP-1 cells were stimulated and harvested at different time intervals and probed with phospho-Abs against STAT-1 and IRF-3 (Fig. 2D). It is notable that CS14K was particularly effective in activating STAT-1 and IRF-3. To ensure that the activation of STAT-1 and IRF-3 is not influenced by any dsRNA contamination, the cells were stimulated with proteins previously digested with RNase III (Supplemental Fig. 2).

NF-κB, an important transcription factor responsible for activating several genes involved in innate immune signaling in macrophages, has been shown to be blocked by CS protein (11). Therefore, we analyzed whether CS14K could overcome this constraint. Although both CS and CS14K proteins led to early degradation of IkB (Fig. 2E), the movement of p65 into the nucleus was inhibited (Fig. 2F).

In summary, these data establish that the fusion of A27 protein to CS could alter the innate immune responses in macrophages by STAT-1 and IRF-3 activation without the involvement of NF-κB, which possibly resulted in increased levels of NO, IL-6, and IL-12 production, suggesting signaling via the TLR4 pathway. CS14K was also able to activate higher levels of CD54, a marker for macrophage activation (data not shown). We therefore sought to investigate whether such a response could alter the immunogenicity of the protein in vivo.

**CS14K fusion protein priming arms the humoral responses in vivo**

It is well established that a strong Ab response against CS correlates with higher protection in many animal and human models (31, 32). The magnitude, isotype, and systemic availability of the Ab produced are important parameters that govern the control of parasites and help to resolve disease (33, 34).

Two weeks after the final phase of the prime–boost vaccination protocol described in Materials and Methods, quantitative analysis...
of the Abs produced showed that priming with CS14K chimeric protein followed by MVA-CS indicated significantly higher titers of Abs than mice primed with DNA-CS or CS protein alone (Fig. 3A). The Ab titers of CS14K-primed group was 2-fold greater than the CS-primed group (p = 0.008) and 8-fold more than in the DNA-CS–primed group (p = 0.002). Even 53 d after boost, the Ab levels in all of the immunized groups were maintained at similar levels except for DNA-CS, which showed a reduction by 1 log (data not shown).

Further analysis of IgG isotype switching showed that fusion protein induced a higher IgG1 response whereas the DNA-vaccinated group showed a predominance of IgG2a response. The levels of IgG1 (p = 0.012) and IgG2a (p = 0.030) induced by the fusion protein-primed group were almost double those of the CS-primed group (Fig. 3B). Finally, we sought to determine whether the vaccination with chimeric protein affected the avidity of the CS-specific Abs. The Abs from mice primed with CS14K showed higher avidity (EC50 = 4.2 μM urea) whereas those primed with DNA-CS (EC50 = 2.3 μM urea) or CS (EC50 = 2.5 μM urea) protein had lower affinity (Fig. 3C). Taken together, these data indicate that priming with proteins, as compared with DNA alone, enhances levels of Abs against CS. Additionally, priming with CS14K not only increases the overall production of Abs but appears to induce a more balanced production of high-affinity IgG1 and IgG2a Abs.

Having established that CS14K can mediate the induction of NO in cultured macrophages, we sought to assess the ability of splenocytes from vaccinated mice to respond to purified CS protein by evaluating NO production. Consequently, splenocytes harvested from animals receiving a different prime–boost vaccine regimen 53 d after boost were stimulated with 5 μg/ml purified CS protein for 48 h. Mice that were primed with DNA-CS showed no detectable NO. Alternatively, mice that received proteins as priming agents were able to induce NO production (Fig. 3D). Whereas CS protein was able to increase NO by low levels, CS14K increased NO production by 4-fold (p = 0.001). These data indicate a reduced capability of CS-based vaccination to induce NO compared with CS14K. The unique nature of CS-based malaria vaccines to induce NO has not been reported before. Thus, the fusion of oligomerization domain of 14K protein to CS improves the humoral arm of immune responses against the Ag along with increased production of NO.

**CS14K generates durable and polyfunctional CS-specific CD8⁺ T cells in mice**

Given the critical role of 14K fusion in aggregating CS protein and the enhancement of innate immune responses, we examined the influence of these factors may have on the development and maintenance of CD8⁺ T cells in vivo.

After mice were immunized, they were sacrificed on day 14 to study the adaptive immune response and on day 53 for memory analysis. Efficacy of the vaccination regimen was analyzed using an IFN-γ-based ELISPOT assay and multiparameter flow cytometry. The fusion protein significantly improved the CS-specific IFN-γ-secreting cells (p < 0.05) compared with other vaccine regimes (Fig. 4A, 4B). We characterized the immune responses in terms of polyfunctionality, CD8⁺ T cells specific for the peptide 280–288 of *P. yoelii* CS protein, secreting IFN-γ, TNF-α, IL-2, or any combination among these three cytokines. A clear dominance in CD8⁺ T cell-secreting cytokines was seen in CS14K-primed group of animals over CS protein (2.2-fold)- or DNA-CS (4.3-fold)-primed groups (Fig. 4B) at day 14 after boost. Clearly, cells that secreted IL-2 were associated with triple cytokine-producing cells. Vaccine regimen involving DNA-CS priming resulted in single- or...
double-positive cells and did not induce any triple-positive population. In contrast, animals receiving protein prime were able to induce IFN-$\gamma$+TNF-$\alpha$+IL-2+–secreting cells with CS14K bringing an $\sim$3.6-fold increase over CS. Additionally, CS14K significantly increased double-positive IFN-$\gamma$+TNF-$\alpha$+ over both CS protein and DNA-CS priming.

To study memory responses, the CD8$^+$ T cells were classified into central memory (CD44$^{\text{high}}$CD62L$^{\text{high}}$), effector cells (CD44$^{\text{high}}$CD62L$^{\text{low}}$), terminally differentiated memory effector memory cells (CD44$^{\text{low}}$CD62L$^{\text{low}}$), and naive cells (CD44$^{\text{low}}$CD62L$^{\text{high}}$) (35, 36). Upon analyzing memory responses on day 53 and also on day 120 (data not shown), we observed rapid proliferation of the peptide-specific effector CD8 T cell population. Interestingly, the frequency of cells secreting IFN-$\gamma$ was maintained even during the memory stage by CS14K priming compared with other groups (Fig. 4C). In terms of polyfunctionality, priming with fusion protein had a significant increase in the triple-positive (IFN-$\gamma$+TNF-$\alpha$+IL-2+) and double-positive (IFN-$\gamma$+TNF-$\alpha$+) population of cytokine-secreting cells over both CS protein and DNA-CS priming.

Most of the single-positive cells were dominant for TNF-$\alpha$. A 3.5-fold increase in IFN-$\gamma$+TNF-$\alpha$+IL-2+–secreting cells by the CS14K-primed group over the CS protein-primed group and a nearly 14-fold increase over the DNA-CS–primed groups was seen. A clear hierarchy in CS peptide-specific total CD8$^+$ T cells secreting cytokines was observed when priming was carried out with CS14K (Fig. 5A). Mice immunized with proteins showed significant increase in magnitude of cytokines released over DNA-CS priming. Indeed, priming with CS14K protein rather than CS protein led to a 2-fold increase in total IFN-$\gamma$ ($p < 0.05$) and TNF-$\alpha$ ($p < 0.0005$) in addition to the 3-fold increase in IL-2–secreting cells ($p < 0.05$). Interestingly, all of the positive CD8$^+$ T cells produced by the vaccine regimen had a phenotype resembling either T effector cells or terminally differentiated effector memory cells. To extend the immune analysis, we evaluated the amount of IFN-$\gamma$ or TNF-$\alpha$ secreted by the different populations based on the mean fluorescence intensity (MFI) calculation (Fig. 5B). We observed that, irrespective of priming agent, most of the IFN-$\gamma$ was produced by IFN-$\gamma$+TNF-$\alpha$+IL-2+ or IFN-$\gamma$+TNF-$\alpha$+ cells with a similar pattern for TNF-$\alpha$. The lack of CM CD8$^+$ T cells could be explained by the fact that upon exposure to peptide stimulus most of the CM cells rapidly acquired effector memory characteristics. These data indicate that proteins are able to prime to a higher extent an effective long-term CD8$^+$ T cell response than does DNA.

The enhanced CD8$^+$ T cell responses in mice immunized with CS14K rather than CS is due to the adjuvant-like effect of the unique A27 element.

**CS14K fusion protein abrogates the liver stage development of parasites when challenged with sporozoites**

Given that polyfunctional CD8$^+$ T cell responses in addition to high-avidity Abs are crucial in enhancing protection against parasite infection, and also that CS14K improved both adaptive and memory immune responses, compared with CS or DNA-CS
priming, we sought to investigate the efficacy of immunization against sporozoite challenge. Protection was evaluated by measuring levels of \textit{P. yoelii} 18S rRNA in liver and also by determining parasitemia in sporozoite-challenged animals. To investigate the protective capacity of this subunit vaccine, groups of mice were immunized in various prime–boost protocols (Fig. 6). We show that a heterologous protein prime/vaccinia virus boost regimen was found to be more effective than a homologous protein prime/protein boost regimen, and a 32% reduction in the liver stage burden of parasites was observed in mice receiving CS14K (CS14K/CS14K) compared with CS (CS/CS) protein. Furthermore, a MV A-CS boost significantly lowered the parasite levels in the liver. Moreover, the CS14K protein priming followed by MV A-CS resulted in a near complete inhibition (∼99.9%; \( p, 0.005 \)) of parasite development in liver compared with CS protein and DNA-CS (Fig. 6).

To evaluate whether protection from liver-stage parasites induced by CS14K/MVA prime–boost can prevent the development of blood stages, mice were challenged with 300 live \textit{P. yoelii} sporozoites and monitored until patency. Importantly, none of the vaccinated animals developed blood stages from days 3 to 21 of daily follow-up after challenge with sporozoites (Table I).

Thus, priming with CS14K protein in combination with a boost by MVA-CS is a highly effective vaccine regimen against murine malaria liver stage parasites. A proposed mechanism is shown in Fig. 7.

**Discussion**

In this study, we report a novel approach in designing effective protein vaccines based on the fusion of 14K (A27) protein of vaccinia virus with a model Ag, CS of \textit{P. yoelii}, for developing an effective vaccination regimen based on protein prime/vaccinia virus boost against murine malaria. An obvious advantage of this vaccine strategy is the concomitant increase in both CS-specific humoral and CD8 + T cells up to 4 mo. By comparing CS14K protein with other priming agents such as CS- and DNA-CS–based vaccines, we showed the correlation between the immune responses generated and protective efficacy generated in the murine malaria model.

In this study, we showed that the fusion of 14K protein with CS results in a protein with high tendency to form oligomers/aggregates of CS14K, which, in turn, enhanced the immune response profile leading to protection against murine malaria. Previous studies have demonstrated the differences in CS-specific T cell proliferation using recombinant soluble CS protein, viable sporozoites, or heat-killed sporozoites (37). This could explain the enhancement of polyfunctional T cells and Abs when animals are primed with chimeric protein rather than CS, which contains lower amounts of CS aggregates. An obvious advantage of this chimeric protein is the incorporation of near full-length CS protein containing both B and T cell epitopes whose presentation would be enhanced as a result of aggregation. Additionally, the use of viral vaccinia vectors for boosting could further enhance the development of long-term effector CS-specific T cells. In Fig. 6, we show how the vaccine regimen based on CS14K prime/MVA-CS boost inhibits the liver stage development of the parasite in mice when challenged with a high dose of sporozoites. When we calculated the C\(_T\) value of individual mice in each group we found that for the CS14K/MVA-CS group the values were well >35, the reference value in naive mice (indicated as dotted lines in Fig.
This suggests that the mice in this group did not have any parasite in the liver. Moreover, given that sterile protection is the target of pre-erythrocytic vaccines, we showed that CS14K/MVA-CS prime-boost prevented the development of blood stage parasites (Table I). The maintenance of both cellular and humoral responses even after 4 mo indicated the strong nature of the vaccine regimen used in this study.

Besides protein aggregation, another mechanism that could explain the enhanced immunogenicity of CS14K is its ability to activate type I IFN signaling. Given that an early activation of innate immune responses curbs the pre-erythrocytic development of parasites (38, 39), activation of type I IFN signaling by fusion protein further validates its use as a priming agent. Indeed, activation of type I IFN signaling by the chimeric protein did result in elevated levels of cytokines belonging to this family such as IL-6, TNF-α, and IL-12. It is thought that the parasiticidal action of TNF-α in hepatocytes is mediated through the synthesis of IL-6 (40). Furthermore, engaging TLRs is critical in developing anti-malarial immunity, which could be explained by the poor immunogenicity of RTS,S without a strong adjuvant such as AS01E, a TLR4 agonist (41). A recent study also shows the importance of TLR signaling for effective development of protective Abs to Plasmodium (42). This is in agreement with our study, which shows how enhancement in STAT-1 and IRF-3 by CS14K suggested a role in TLR4 activation (Fig. 7), resulting in improved immunogenicity and inhibiting the liver stage development of the parasite (43, 44).

It is known how parasites such as Leishmania and Plasmodium downregulate synthesis of NO (45). Our study shows the inherent capacity of chimeric protein to enhance the production of NO both in vitro and in vivo. Because neither CS nor 14K protein alone could induce NO, it seems reasonable to conclude that the CS14K

**FIGURE 5.** Polyfunctionality related to increased cytokine production by CS14K induced memory CD8+ T cells: characterization of memory CD8+ T cells. (A) Representation of total CS peptide specific CD8 cells were gated on CD44 and CD62L cell surface markers represented on left-hand side of the diagram. Total CD8 responses for each group are represented inside the box. Different populations of memory CD8 cells secreting different cytokines are represented on the right-hand side with their respective total responses as depicted inside the box. (B) MFI of IFN-γ and TNF-α produced by different polyfunctional population of effector memory CD8+ T cells (n = 4 mice/group).
The presence of strong adjuvants such as Freund’s adjuvant (50) enhances the ability of fusion proteins to induce strong humoral responses (47). Previous studies have also shown the importance of fusion proteins to induce strong humoral responses based on their ability to form large molecular mass aggregates (20). Similar to the CS14K priming approach presented in this study, the malaria vaccine in the latest stages of clinical trials, RTS,S, is also dependent on a higher magnitude of Abs belonging to the IgG1 subclass (48). Vaccine studies based only on DNA-CS or protein alone suggest that differences exist in the processing of these Ags by APCs. Additionally, early production of NO has been proposed to be required for proliferation of CD8+ T cells against the parasite (46). In conclusion, fusion of 14K to CS leads to a more favorable immunogenic milieu to enhance the efficacy of the vaccine.

Protective immunity against malaria associates with circulating IgG Abs against CS (47). Previous studies have also shown the ability of fusion proteins to induce strong humoral responses based on their ability to form large molecular mass aggregates (20). Similar to the CS14K priming approach presented in this study, the malaria vaccine in the latest stages of clinical trials, RTS,S, is also dependent on a higher magnitude of Abs belonging to the IgG1 subclass (48). Vaccine studies based only on DNA-CS vaccination lacking the GPI anchor improve protection up to 95% based only on Ab titers (49). Also, the serum Ab titers obtained by fusion protein priming are much higher than those obtained in presence of strong adjuvants such as Freund’s adjuvant (50).

Two weeks after boost animals were challenged with 2 × 10^4 P. yoelii sporozoites through i.v. route via tail vein. After 42 h, the amounts of parasites in the liver were estimated by measuring the number of copies of 18S rRNA by RT-PCR. The dotted line represents the minimum detectable levels by the highly sensitive quantitative RT-PCR (26). Data are expressed as means ± SEM (n = 6 mice/group). *p < 0.05, **p < 0.005.

Table I. CS14K priming induces sterile protection

<table>
<thead>
<tr>
<th>Priming*</th>
<th>Boosting*</th>
<th>Challenge*</th>
<th>Protected/Challenged</th>
<th>Preemptant Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS14K</td>
<td>MVA-CS</td>
<td>300 sporozoites</td>
<td>10/10 (100)</td>
<td>—</td>
</tr>
<tr>
<td>PBS</td>
<td>PBS</td>
<td>14 d after boost</td>
<td>0/10 (0)</td>
<td>4 d</td>
</tr>
</tbody>
</table>

*Protein (20 µg) administered intradermally.

*Virus (2 × 10^7 PFU) administered i.p.

*Challenge via i.v. route.

However, unlike vaccination with DNA-CS or protein alone, vaccination with the CS14K fusion protein results in the concomitant induction of high levels of both IgG1 and IgG2a Abs, and this could be beneficial in providing protection against malaria. The chimeric protein proved to be superior in inducing high-avidity Abs against CS comparable to those achieved by using nanoparticle-based CS peptide vaccination (51). High-avidity Abs produced by the chimeric protein could be due to the unique folding pattern attained by CS14K protein resulting in exposure of other potent epitopes. These data have important implications in the choice of Ags designed to induce Ab responses against malaria.

CD8+ T cell responses against CS protein are known to be an important factor in the development of sterile immunity using irradiated sporozoites (52). A role of CD8+T cells in subunit vaccine-induced protection against malaria was initially established in a heterologus prime–boost approach with flu and vaccinia virus vectors expressing CS (53). Vaccine regimes based on DNA as well as protein prime followed by vaccinia boost have shown to induce CS peptide-specific CD8+ T cell responses (15, 54). The ability of the CS14K regimen to produce significantly elevated levels of IFN-γ and TNF-α single-positive—as well as double-positive—secreting CD8 T cells even after 53 d makes it a suitable vaccine candidate. We also report a shift in polyfunctionality of CD8 T cells secreting all three cytokines produced by chimeric protein, which may be effective in controlling the growth of parasites. Furthermore, the ability of CS14K vaccine regimen to induce a large amount of TNF-α has an added advantage because it is known to be an important cytokine for the maintenance of memory CD8 T cells in malaria (55). The concomitant increase in TNF-α by CS14K could also contribute to elevated NO and Ab responses (56, 57). The need for IL-12 in the development of effective adaptive responses is in agreement with our study, which shows the ability of CS14K protein to prime an enhanced T cell response (58). The data from MFI studies show that double-or triple-positive populations are better in producing elevated levels of NO and Abs.
of cytokines, both of which are elevated by CS14k priming. The low levels of CD8 responses against DNA-CS suggest that the protective capability of this regimen may be based on the Abs induced. The maintenance of such durable and multifunctional memory responses even after 4 mo of vaccination, taken together with sterile protection, justifies the potent nature of A27 protein fusion and its use for developing better malaria vaccines. In recent years, many experimental CS-based vaccines have been developed using novel adjuvants or by modifying the structure of CS protein (Supplemental Fig. 3). The assessment of protective capacity of vaccine regimen gave a good platform to study the efficacy of chimeric protein generated by 14k fusion. Chimeric protein prime–boost alone could inhibit the liver stage development better than those attained by DNA-CS prime and MVA-CS boost under a high challenge dosage. This could be explained by the potent nature of the attenuated vaccinia viral boost (59).

Developing an effective pre-erythrocytic vaccine against malaria is difficult because it requires high levels of both humoral and cell-mediated immune responses. The role of CSP Abs in providing protection is a much debated topic. However, the latest trial of RTS,S vaccine does show that the protection could be mediated by CSP Abs (60, 61). In this study, we describe an optimal prime–boost approach using modified CS protein fused to the 14k protein of vaccinia virus as priming and MVA-CS as a boost. This unique approach is able to improve the magnitude and polyfunctionality of both humoral as well as cell-mediated immune responses, resulting in complete protection of vaccinated animals compared with other experimental CS based vaccines (62–65). Additional studies in primates could be performed using the chimeric protein in prime–boost protocols. These data establish a ground for the studies in primates could be performed using the chimeric protein with other experimental CS based vaccines (62–65). Additional

Disclosures

The authors have no financial conflicts of interest.

References


43. Li, C., and J. Langhorne. 2000. Tumor necrosis factor α g55 receptor is important for development of memory responses to blood-stage malaria infection. Infect. Immun. 68: 5724–5730.


MCYHEEHDIKLYHVLHSKIGKIVHRHHLRLGDZLNGKEPE
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IHAETLRAAMISLAAKDVO/TGRRFYE
Fig S1. Related to Figure 1. (A) Design and construction of plasmid encoding CS14K. A schematic representation of gene encoding the CS protein of *P. yoelii* and that of 14K of vaccinia virus is shown. For generating the CS fragment of chimeric protein, the signal sequence represented by 37 amino acids of amino terminal and GPI motif involving the 19 amino acids at the carboxy terminal of CS gene was removed by PCR amplification. The C terminal of amplified CS fragment was then fused in frame to the N terminal of gene, A27L of vaccinia, lacking first 28 amino acids. All PCR amplifications were carried out using appropriate primers carrying restriction enzyme sites. This fusion construct was then fused in frame with the GST tag present within the pGEX-6P-1 vector. (B) CS14K fusion protein does not induce neutralizing antibodies against MVA. Serum from rabbits injected with CS14K protein was not able to neutralize the infectivity of MVA virus *in-vitro*. 200 PFU of MVA virus were incubated with two fold dilutions of serum from rabbit injected with CS-14K protein. Following infection of DF-1 cells with the serum treated virus, immunostaining was carried out using anti-vaccinia antibody used at a dilution of 1:1000. Virus incubated with rabbit serum was used as mock control. The numbers of plaques formed by the viruses were counted to determine the extent of infection. (C) Localization of proteins in macrophages: Confocal microscopy analysis of CS and CS14K localization in macrophages using NYS1 antibody at a dilution of 1:500. Pattern of CS expression by MVA-CS. Confluent J774 cells were infected with MVA-CS at an MOI of 5 PFU/cell for 18 hours. Protein localisation in transfected macrophages. CS and CS14K encoding plasmids, pCINeo-CS (10 µg) and pCDNA-CS14K (10 µg), both lacking the signal and GPI motifs were transfected into confluent mono layers of J774 cells for 24 hours. (D) Amino acid sequence of CS14K: Amino acid composition of CS14K protein is represented with the 14K sequence under lined.
Fig S2. Related to Figure 2. IRF-3 and Stat-1 activation by CS14K is independent of ds-RNA: Proteins (5 µg/ml) and LPS (1 µg/ml) were treated with 2 µl of Shortcut RNase III for 30 minutes at 37ºC. Following incubation the proteins were mixed with RPMI medium containing 10% serum to inhibit RNase III and incubated with 2×10⁶ cells of THP-1 for defined time intervals. The levels of phospho IRF-3 and Stat-1 were detected as mentioned in Materials and Methods.
<table>
<thead>
<tr>
<th>Priming Agent</th>
<th>62 CS (DNA)</th>
<th>49 CS (DNA)</th>
<th>15 CS+CM-CSF (DNA)</th>
<th>CS14K (Protein)</th>
<th>63 CS gp64 (Protein)</th>
<th>64 Sb824 pST-TB (Protein) (3)</th>
<th>65 ADPyCS + 7DW8-5</th>
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<td>Boosting Agent (Number of Boosts)</td>
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</table>

a CS gp64 : Recombinant Baculovirus expressing CS fused with baculovirus envelope protein gp64.
b Sb824 pST-TB : Salmonella expressing fusion protein between YoPE and CS peptide.
c ACT-CS : Bordetella pertussis adenylate cyclase toxoid fused with CS peptide.
d ADPyCS + 7DW8-5 : Adenovirus expressing CS mixed with a synthetic analogue of Alpha-Galactosylceramide.

+ Low Levels (Elispot: Less than 400 spots.) (CD8 T-cells: Less than 1%.) (Antibody Levels: < 1:50,000)
++ Moderate Levels (Elispot: 450-850 spots.) (CD8 T-cells: 1% to 3%). (Antibody Levels: 1:50,000 to 100,000)
+++ High Levels (Elispot: 900-1200 spots.) (CD8 T-cells: 4% to 6%). (Antibody Levels: 1:200,000 to 1:500,000)
++++ Very High Levels (Elispot: 1500-2000 spots.) (CD8 T-cells: 10% and above.) (Antibody Levels: > 1:800,000)

**Fig S3.** Comparative analysis of immunogenicity of various CS based vaccine regime. A detailed comparison of various murine plasmodial CS based vaccine regime against CS14K protein prime and MVA-CS boost. Experiments were appropriate studies were not carried out are represented as ND.