Low doses of killed parasite in CpG elicit vigorous CD4+ T cell responses against blood-stage malaria in mice

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Development of a vaccine that targets blood-stage malaria parasites is imperative if we are to sustainably reduce the morbidity and mortality caused by this infection. Such a vaccine should elicit long-lasting immune responses against conserved determinants in the parasite population. Most blood-stage vaccines, however, induce protective antibodies against surface antigens, which tend to be polymorphic. Cell-mediated responses, on the other hand, offer the theoretical advantage of targeting internal antigens that are more likely to be conserved. Nonetheless, few of the current blood-stage vaccine candidates are able to harness vigorous T cell immunity. Here, we present what we believe to be a novel blood-stage whole-organism vaccine that, by combining low doses of killed parasite with CpG-oligodeoxynucleotide (CpG-ODN) adjuvant, was able to elicit strong and cross-reactive T cell responses in mice. Our data demonstrate that immunization of mice with 1,000 killed parasites in CpG-ODN engendered durable and cross-strain protection by inducing a vigorous response that was dependent on CD4+ T cells, IFN-γ, and nitric oxide. If applicable to humans, this approach should facilitate the generation of robust, cross-reactive T cell responses against malaria as well as antigen availability for vaccine manufacture.

Introduction
Vaccines are among the most cost-effective strategies for preventing infectious disease. However, a highly effective vaccine against malaria remains elusive (1). Historically, vaccines have been most successful against diseases in which natural infection leads to long-lasting immunity, and the best formulations have usually been the ones closely mimicking natural infection (2, 3). However, in the case of malaria, natural immunity is slow to develop and incomplete even after years of continuous exposure (4). While some protection from severe disease can be achieved, “immune” individuals can rapidly become reinfected following cure (5). These observations have indicated that a vaccine-mimicking natural immunity is unlikely to be effective, and alternative paradigms have been investigated.

RTS,S, the most advanced malaria vaccine, for instance, relies on administration of a complex construct containing a large part of the circumsporozoite protein (CSP) fused to the HBV surface antigen (HBV-S Ag) and free HBV-S Ag, formulated in a 3-component adjuvant. This preerythrocytic vaccine induces an approximately 30%–50% reduction in the risk of clinical malaria through antibody- and cell-mediated responses, thus positioning it close to licensure trials (6, 7). In the case of blood-stage malaria, however, success has been limited. The main targets have been proteins on the surface of the merozoite or the infected red cell; and thus, antibodies have been recognized as the main mediators of protection (8). Therefore, several vaccine candidates have been selected using sera from immune individuals (9, 10), and many of the current candidates aim to induce antibody responses of the same type as those induced through natural infection (11). Antigenic variation and polymorphism will probably limit their protective efficacy (12, 13).

In contrast to antibodies, T cells mainly recognize antigen after it has been processed and presented in the context of MHC molecules. This process allows T cells to target internal and possibly more conserved epitopes and/or antigens. In rodent malaria, for example, CD4 T cells are able to recognize cryptic epitopes of polymorphic antigens such as apical membrane antigen 1 (AMA-1) or CSP (14, 15). Similarly, rodent CD4 T cells can recognize invariant regions on surface antigens such as CSP (16) or internal and highly conserved proteins such as hypoxanthine-xanthine-guanine phosphoribosyl transferase (HGXPRT) (17). More importantly, rodent and human CD4 T cells are able to target invariant epitopes of polymorphic antigens such as CSP in the context of different MHC-II molecules (16, 18, 19), suggesting that malaria-specific CD4 T cells can be broadly reactive. This phenomenon has already been documented in B cell–deficient mice in which cross-protection against heterologous parasites can be readily induced in the absence of humoral responses (20). Therefore, it would be expected that a malaria vaccine able to induce vigorous T cell responses against multiple antigens would engender some degree of cross-protection. However, most blood-stage candidates induce suboptimal T cell responses and only against 1 or 2 antigens. To maximize the spectrum of antigens, whole-organism immunization has been proposed and tested against liver or blood stages (1, 21). The available data suggest that (a) immunization of humans with irradiated sporozoites or (b) mice with genetically attenuated sporozoites (22, 23) or (c) humans with blood-stage parasites curtailed by antimalarials (24, 25) can achieve sterile immunity or delayed parasitemia through the induction of vigorous T cell responses.

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Here, we set out to develop a whole-organism vaccine able to stimulate vigorous and broadly reactive T cell responses against malaria. Four considerations were essential prior to development. First was the need to keep a broad spectrum of target antigens. To circumvent the limitations of using live parasites yet retain the maximum spectrum of antigens, a formulation based on killed parasites was ideal. Second was the selection of a potent T cell adjuvant. Here, the selection of an adjuvant such as CpG-oligodeoxynucleotides (Cpg-ODN), used extensively in human trials, easy to admix with parasites and conducive of vigorous Th1 differentiation, was suitable (26). Third and more importantly was the determination of the right antigenic dose. Here, administration of low doses of parasite was desired in order to elicit better T cell responses. This was based on extensive evidence demonstrating that low doses of live parasite induce vigorous T cell responses (24, 25), while higher parasite burdens result in deletion of effector cells (27). Finally, a high-dose whole-parasite vaccine would not be possible due to logistical constraints.

By addressing these issues, we developed a 3-prong vaccine approach combining (a) low doses of (b) whole-killed parasite (c) adjuvanted in CpG-ODN. By applying this strategy, we show that rodent immunization with only 1,000 killed parasites in CpG-ODN induces durable protection against homologous and heterologous challenge. Strong, durable, and cross-reactive CD4 T cells were easily induced by this type of vaccine, thus providing useful insight for the development of better T cell vaccines against malaria.

**Results**

**Definition of a low-dose whole-parasite vaccine.** To define an effective vaccine containing low doses of *Plasmodium* antigen, A/J mice were immunized with decreasing doses of *P. c. chabaudi* AS antigen equivalent to 10³, 10¹, or 10⁰ parasitized rbc (prbc) in CpG-ODN or control ODN. Two weeks after immunization, mice were given a lethal parasite challenge. Parasitemia (Figure 1A) and disease severity (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI39222DS1) were monitored. Similar to naive mice, animals immunized with adjuvant alone (0 prbc), ultra-low (10⁰ prbc), or low (10¹ prbc) doses of parasite in control ODN experienced high mortality and severe disease. Mice immunized with high doses of parasite (10⁴ prbc) in control ODN survived challenge but exhibited high parasitemia and severe disease. In contrast, high doses of parasite (10⁴ prbc) in CpG-ODN achieved 100% protection, markedly reducing parasitemia and disease severity. Of greater interest, however, low doses of parasite (10¹ prbc) in CpG-ODN induced complete protection with limited disease severity and low parasitemia. Ultra-low doses (10⁰ prbc) in CpG-ODN still resulted in 60% protection.

To investigate the induction of cellular and humoral immunity following vaccination, parasite-specific proliferation (Figure 1B) and antibody titers (Figure 1C) were assessed in all groups. PBS-control mice showed no parasite-specific proliferation (<5 x 10³ cpm) or antibody responses (IgG titers: <1:500). In accordance with their susceptibility to challenge, animals immunized with adjuvant alone (Cpg-ODN or control ODN), ultra-low (10² prbc), or low (10¹ prbc) doses of parasite in control ODN also failed to exhibit proliferative or antibody responses (<5 x 10³ cpm and IgG titers: <1:500). Animals immunized with high doses of parasite in control ODN showed modest T cellular (10 ± 3 x 10³ cpm) and humoral responses (IgG titer: 1:1,000–2,000) while the same doses of parasite (10⁴ prbc) in CpG-ODN induced robust proliferation (30.0 ± 3 x 10⁵ cpm) and higher antibody levels (IgG titer: 1:9,000–12,000). Interestingly, animals immunized with low doses (10¹ prbc) of antigen in CpG-ODN exhibited strong cellular responses (20 ± 4 x 10³ cpm) yet only modest antibody levels (IgG titer: 1:3,000–5,000). Ultra-low doses (10⁰ prbc) of parasite in CpG-ODN induced limited cellular (12.0 ± 2 x 10³ cpm) and no measurable antibody responses (IgG titer: <1:500). Thus, lower doses of antigen (10¹–10² prbc) induced protection by the elicitation of strong cellular and modest antibody responses.

**Mechanisms of protection.** Because our strategy was to investigate the most effective low-dose vaccine, we focused on the formulation containing the lowest dose of parasite able to elicit a stable and protective response (i.e., 10⁰ prbc in CpG-ODN). To define the protective mechanisms, first, we assessed the pattern of serum cytokine secretion during immunization (priming and boost injections) and challenge. During priming, administration of parasite in CpG-ODN, but not parasite in control ODN, induced rapid secretion of IFN-γ, IL-12, TNF-α, and IL-10 (Figure 2A). Following boost injections (parasite alone), minimal cytokine release was observed. Upon lethal challenge, however, mice immunized with parasite in CpG rapidly produced IFN-γ, IL-12, TNF-α, and IL-10 (Figure 2A). Mice immunized with parasite in control ODN also produced significant amounts of TNF-α and IL-10 but succumbed to infection. This finding suggested that neither TNF-α nor IL-10 were required for protection. To ascertain this hypothesis, mice immunized with parasite in CpG were depleted of each cytokine, IFN-γ, IL-12, TNF-α, or IL-10, and then challenged (Figure 2B). In accordance with serum cytokine data, depletion of IFN-γ and IL-12 abrogated protection, while blockade of TNF-α or IL-10 had no effect on survival. Because plasmacytoid DC (pDC), conventional DC (cDC), NK, and T cells were shown to be responsible for the rapid production of IL-12 and IFN-γ following CpG-ODN administration (Supplemental Figure 2), cell-depletion studies were also performed (Figure 2B). Depletion of pDC had no effect on survival, suggesting additional sources of IL-12 (i.e., cDC) as shown in supplemental data (Supplemental Figure 2). Similarly, depletion of NK or CD8 T cells had no effect on survival, while depletion of CD4 T cells completely abrogated protection. Finally, given the requirement for IFN-γ and IL-12 to drive nitric oxide (NO) production for parasite elimination (28), NO depletion was also performed. In keeping with the requirement for CD4 T cells, IFN-γ and IL-12, depletion of NO abrogated protection.

Despite the moderate level of parasite-specific antibodies induced by the low-dose vaccine, we set out to study their role in protection. First, we determined the kinetics of the antibody response during immunization and challenge. Total parasite-specific IgG antibody levels increased slowly through immunization to reach a titer of 1:2,000–5,000 prior to challenge (day 49). After challenge and coincident with increasing parasitemia, IgG titers decreased to less than 1:500, the lowest at the time of peak parasitemia (days 57–60). During parasite clearance (days 61–80), however, antibody levels progressively increased, reaching a final titer of 1:20,000–30,000. The initial reduction and subsequent increase suggested that vaccine-induced antibodies were being adsorbed during acute infection and produced during recovery. To better define a role for antibodies, adoptive transfer studies were also performed. For this purpose, sera from mice immunized with parasite in CpG-ODN were collected after completion of the immunization regime (day 49 after immunization) and adoptively
transferred into naive recipients (Figure 2D). Here, we found that recipients of sera from CpG-immunized mice reached antibody levels comparable to donors (3 to $5 \times 10^3$) after 3 injections (Figure 2D). Upon challenge, however, titers decreased, suggesting adsorption of parasite-specific antibodies as observed following direct immunization (Figure 2C). Despite this, all recipients of sera from CpG-immunized mice succumbed to infection (Figure 2E) as did recipients of sera from ODN-immunized mice. Therefore, the
Figure 2
Mechanisms of protection. (A) To determine the profile of cytokine secretion, levels of IFN-γ (first panel), IL-12 (second panel), TNF-α (third panel), and IL-10 (fourth panel) were assessed in sera collected from A/J mice during immunization with 10^3 (AS) prbc in control ODN (gray bars) or CpG-ODN (black bars) as well as following challenge (dotted boxes). Immunizations (1 priming and 2 boost injections) are indicated as black arrows and challenge as a gray arrow on the x axis (days after immunization). Results represent mean ± SEM. † Animals that succumbed to infection. (B) To identify cytokines and cell types essential for protection, A/J mice were immunized with 10^3 prbc in CpG-ODN and subsequently depleted of IFN-γ, IL-12, TNF-α, IL-10, pDC, NK, CD8, CD4, or NO as described in Methods. Mice were then challenged i.v. with 10^5 homologous (AS) prbc and the outcome of infection monitored. Data for individual mice are shown. (C) To determine the kinetics of antibody responses during immunization and challenge, A/J mice were immunized with 10^3 (AS) prbc in CpG-ODN (black circles) or control ODN (white circles) and titers of parasite-specific IgG against homologous antigen assessed. Reciprocal median total IgG titers and interquartile ranges are shown. (D, E) To ascertain a role for antibodies in protection, naive A/J mice were adoptively transferred on days −1, 0, and 1 with 200 ml i.p. of serum collected from immune (10^3 [AS] prbc in CpG-ODN) or control (10^3 [AS] prbc in control ODN) donors. Recipients were challenged with homologous 10^5 (AS) prbc on day 0. Parasitemia and parasite-specific IgG titers against homologous (AS) were monitored. Reciprocal median total IgG titers and interquartile ranges are shown. All data sets are representative of 5 mice per group of 2 independent experiments performed.
indicated by reduced parasitemia and disease severity. As shown in Figure 2, cytokine secretion indicated the predominance of a Th1 profile. As such, high levels of TNF-α, IFN-γ, and IL-2 were produced not only in response to AS but also to AJ and YM parasites (Figure 4A).

To determine the longevity of this response, proliferation (Figure 4B) and IFN-γ secretion (Figure 4C) were assayed at 2, 8, and 12 weeks after immunization. We found that a significant proportion of CD4 T cells from mice immunized with parasite in CpG-ODN proliferated (Figure 4B) and secreted IFN-γ (Figure 4C) against all parasites at 2, 8, and even 12 weeks after immunization. We also tested any antibody cross-reactivity against all parasite strains (Figure 4D). Here, modest IgG titers against homologous parasite (AS) were detected at 2 weeks after immunization (3 to $1 \times 10^3$) but declined to less than $1 \times 10^3$ at 12 weeks. In contrast to cellular responses, antibody titers against heterologous parasites were negligible (less than $1 \times 10^3$) at all time points, suggesting that humoral responses were neither sustained nor cross-reactive.

Heterologous immunity. Next, we set out to ascertain whether the broad recognition of multiple parasites observed in vitro would afford some cross-protection in vivo. For this purpose, strains of mice with differing genetic background (A/J [H-2a], B6 [H-2b], and BALB/c [H-2d]) were immunized with $P. c. chabaudi$ AS and challenged with homologous (AS) or heterologous (AJ or YM) parasites at 2, 8, or 12 weeks (Figure 5). Consistent with the results described in Figure 4, B–C, A/J mice were protected against challenge with all parasites at 2, 8, and 12 weeks after immunization. Interestingly, B6 and BALB/c mice demonstrated long-term protection against AS and AJ but limited protection against YM parasites. Better protection against YM challenge, however, was achieved when animals were immunized with lower doses of homologous parasite ($10^3$ YM prbc) in CpG-ODN (Figure 6). These results showed that cross-protection against highly virulent YM parasites was only achieved in A/J mice despite cross-protection against AS and AJ parasites being achieved in all mouse strains.

Induction of CD4 T cell memory. Because a population of cross-reactive CD4 T cells was known to persist up to 12 weeks after immunization (Figure 4B), we set out to determine whether these cells were required for long-term protection. Two approaches were undertaken. First, to ascertain an effector role for memory CD4 T cells, 12 weeks after immunization, CD4 T cells were depleted and mice given a lethal challenge. Similar to the results described when depletion was undertaken at 2 weeks (Figure 2B), removal

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**Figure 3**

Protection in B cell–deficient mice. To define the role of B cells in protection following immunization, B cell–deficient (μMT) or B cell–sufficient (B6) mice were immunized with $10^5$ (AS) prbc in CpG-ODN or control ODN. 2 weeks later, mice were challenged i.v. with $10^6$ homologous (AS) prbc and (A) parasitemia or (B) disease severity monitored. Disease severity was assessed by determining clinical scores (C). Data for individual mice are shown and are representative of 3 independent experiments. Note that mice of B6 background (B6 or μMT) are naturally resistant to $P. c. chabaudi$ AS and thus, protection was determined by reduced parasitemia and disease severity. ms1, mouse 1.

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modest antibody response induced by vaccination appeared not to play a role in protection.

To further examine the requirement for B cells in the establishment of protective responses after immunization, B cell knockout mice (μMT) were used. These mice are derived from a C57BL/6 (B6) background [H2b] and are known to display functional T cell activation and memory formation (29). However, unlike A/J mice, B6 mice do not succumb to $P. c. chabaudi$. Therefore, protection is indicated by reduced parasitemia and disease severity. As shown in Figure 3, A–C, protection was equivalent in CpG-immunized B cell–deficient (μMT; Figure 3A) or B cell–sufficient (B6; Figure 3B) mice. Altogether, the serum transfer and B cell KO experiments confirmed that neither antibodies nor B cells were necessary for protection following low-dose immunization.

Cross-reactivity and longevity of cellular responses. The known promiscuity of CD4 T cells in human and rodent malaria (16, 18, 19) prompted us to also investigate whether broadly reactive CD4 T cell responses were elicited by the low-dose vaccine. Different lethal strains and species of parasite were selected to stringently assess heterologous immunity in vitro and in vivo. First, we assessed the type of CD4 T cell responses against different strains of parasite in vitro. Mice were immunized with $P. c. chabaudi$ AS (AS), and the profile of CD4 T cell cytokine secretion (Th1 vs. Th2) against homologous (AS) and heterologous (P. c. chabaudi AJ [AJ] and P. Yoelii YM [YM]) parasites was assessed after in vitro restimulation (Figure 4A). In accordance with the results described in Figure 2, cytokine secretion indicated the predominance of a Th1 profile. As such, high levels of TNF-α, IFN-γ, and IL-2 were produced not only in response to AS but also to AJ and YM parasites (Figure 4A).

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Figure 4
Cellular responses and cross-reactivity. (A) To determine CD4 response, A/J mice were immunized with \(10^3\) (AS) prbc in CpG-ODN (black bars) or control ODN (gray bars). Spleen cells collected 2 weeks after immunization were cultured with homologous (AS) or heterologous (P. c. chabaudi AJ [AJ]; P. yoelii YM [YM]) parasites, normal rbc (nrbc), or ConA. After 96 hours, supernatants were assayed for TNF-\(\alpha\), IFN-\(\gamma\), IL-2 (Th1 cytokines), and IL-4 or IL-5 (Th2 cytokines) assessed. Results are mean ± SEM. (B) For proliferation, spleen cells from A/J mice immunized as above were purified at 2, 8, or 12 weeks and incubated for 96 hours with AS, AJ or YM prbc, nrbc, medium, or ConA. Numbers represent percentage of CFSE\(^{\text{dim}}\) CD4 T cells. (C) For cytokine secretion, spleen cells from A/J mice immunized as above were purified at 2, 8, or 12 weeks and incubated for 96 hours with AS, AJ or YM prbc, nrbc, or medium and stained for intracellular IFN-\(\gamma\). Numbers represent percentage of IFN-\(\gamma^+\) CD4 T cells. (D) For cross-reactive antibodies, sera from A/J mice immunized with \(10^3\) (AS) prbc in CpG-ODN (black bars), control ODN (gray bars), or hyperimmune sera (HIS, white bars) were tested for IgG titers against homologous (AS) or heterologous (AJ and YM) parasites. Results represent reciprocal median total IgG titers and interquartile ranges. All data sets are representative of 5 mice per group of 3 independent experiments performed. Significant differences compared with controls are shown. *\(P<0.05\).
of CD4 T cells at 12 weeks resulted in a severe course of infection and death (Figure 7A). Second, to exclude a helper role in recall responses, CD4 T cells were depleted as above, but mice were only challenged 21 days later. This allowed for naive but not memory T cells to replenish. Here, despite the presence of naive/helper CD4 T cells, all animals succumbed to severe infection (Figure 7A). Therefore, vaccine-primed memory CD4 T cells, but not unprimed CD4 T cells, were essential for long-term protection.

Figure 5
Heterologous immunity. To assess for cross-protection, different strains of mice including A/J [H-2^a], B6 [H-2^b] and BALB/c [H-2^d] mice were immunized with 10^3 (AS) prbc and challenged i.v. with homologous (AS) or heterologous (AJ or YM) parasites at 2, 8, or 12 weeks after vaccination. Rechallenge was performed at 12 weeks after parasite clearance. The outcome of infection was monitored by assessing parasitemia. Data for individual mice are shown. nd, not determined. †Animals that succumbed to infection.
Next, we assessed the profile of cytokine secretion of memory CD4 T cells, particularly IL-2 and IFN-γ, as these cytokines exert a profound influence on the maintenance of CD4 T cell memory (30). For this purpose, we estimated the frequency of CD4 T cells able to secrete IL-2, IFN-γ, or both cytokines in the memory phase (Figure 7B). Interestingly, we found that 5.9% ± 0.7% of CD4 T cells (approximately 30%–36% of the parasite-specific CD4 T cell response) were able to secrete both IL-2 and IFN-γ in response to parasite antigen. These data suggested that the low-dose vaccine elicited multifunctional CD4 T cells with potential for memory differentiation.

To formally assess this hypothesis, we analyzed the memory phenotype of CD4 T cells at 12 weeks after immunization. Here, we subdivided CD4 T cell memory cells into central (CM, CD62Lhi) or effector (EM, CD62Llo) subsets based on their expression of the lymph node homing receptor (CD62L) (31). To delineate parasite specificity, we assessed IFN-γ secretion upon in vitro restimulation (Figure 7C). By this method, we found that more than 95% of parasite-specific CD4 T cells were CD62Lhi. The scarcity of the CD62Llo population and the fact that CD62L downregulates upon antigenic stimulation prompted us to also purify individual subsets of CM (CD44hiCD62Llo) and EM (CD44loCD62Lhi) CD4 T cells prior to antigenic stimulation (Figure 7D). By doing so, we confirmed the presence of both types of parasite-specific memory CD4 T cells in CpG-immunized mice. We found that parasite-specific CD62Lhi CD4 T cells (CM) decreased expression of CD62L (data not shown) and exhibited strong proliferative capacity (Figure 7D). In fact, a marked proportion of CD62Lhi CD4 T cells (CM) proliferated and acquired effector function as defined by BrDU incorporation and IFN-γ secretion (97% BrDU⁺ and 39% IFN-γ⁺; Figure 7D). In contrast, CD62Llo CD4 T cells (EM) exhibited less proliferative capacity yet produced considerable more amounts of IFN-γ regardless of their proliferative status (60% BrDU⁺ and 45% IFN-γ⁺; Figure 7D). In conclusion, the low-dose vaccine was able to induce both types of memory subsets where CD62Lhi CD4 T cells (CM) exhibited strong proliferative potential while CD62Llo CD4 T cells (EM) demonstrated rapid effector function on antigen encounter.

**Discussion**

The complexity of the malaria parasite coupled with the lack of evidence for robust infection-induced immunity has made malaria vaccine development especially difficult. Most successful vaccines work by mimicking natural immunity to elicit long-lived antibody responses (32). However, humoral immunity to malaria is incomplete and does not eradicate all parasites or provide protection against future challenge (4, 5). For blood-stage malaria, numerous approaches have been pursued, including the use of subunit antigens, prime-boost strategies, or combination of adjuvants to induce robust humoral responses. Here, we report on what we believe to be a novel T cell vaccine against blood-stage malaria that builds on previous observations that (a) IL-12 and ODN are effective adjuvants for immunization against blood-stage infection (33, 34) and (b) low doses of antigenic stimulus facilitate cellular immunity. We combined these strategies with a whole-parasite approach to demonstrate that administration of extremely low doses of killed blood-stage parasites in CpG-ODN is able to engender broad protection against lethal blood-stage challenge through the generation of cross-reactive T cells.

From a vaccine perspective, understanding the elicitation of such a vigorous Th1 response is essential to harness any T cell vaccine candidate into human trials. It is known, for instance, that vaccine-induced Th1 responses are controlled by the type of APC (35), the dose of antigen (36–38), or the cytokine environment induced during vaccination (39). To develop vigorous T cell responses against blood stage, a number of considerations were of paramount importance for us. First was the selection of a potent adjuvant such as CpG-ODN. While it is still under development, there have been extensive trials of ODN in humans in which it has been shown to activate immune cells to secrete cytokines conducive of Th1 differentiation (26). In the case of malaria, for instance, CpG-ODN has been successfully used as an adjuvant to immunize nonhuman primates and mice (40–42). Second was the identification of an optimal antigenic dose. This was based on evidence that low doses of live parasite facilitate cellular immunity (24, 25), while higher parasite burdens tend to result in apoptosis of Th1 effectors (27). Third was our desire to maintain a broad spectrum of target antigens while avoiding the limitations of live parasites. All of the above led us to combine CpG-ODN with low doses of whole killed parasite in a simple prime-boost regime aiming to afford strong cellular immunity against blood-stage infection. Our results confirmed that CpG-ODN was crucial to providing vigorous polarization signals when the dose of antigen was limiting. In fact, the rapid activation of DCs, B cells, and macrophages to secrete IL-12, IL-10, and TNF-α and NK and T cells to secrete IFN-γ confirmed that the strong recognition of pathogen-associated
molecular patterns was central to determining the type of immune response after vaccination. Clearly, the cytokine environment and the broad range of target antigens available facilitated the priming of broadly reactive CD4 T cells, which upon challenge, rapidly engaged to deliver macrophage activation signals (i.e., IFN-γ) to activate NO production for parasite clearance, reminiscent of the protective mechanisms of the bacillus Calmette-Guérin vaccine (43).

Because we aimed to achieve protective cellular responses, it was essential to ensure that the low-dose vaccine was efficacious at inducing memory CD4 T cells. In this case, the induction of

Figure 7
Induction of CD4 T cell memory. (A) To assess the role of CD4 T cells in protracted protection, A/J mice immunized with $10^3$ (AS) prbc in control ODN or CpG-ODN were depleted of CD4 T cells 12 weeks after immunization. Mice were challenged i.v. with $10^5$ homologous (AS) prbc immediately after depletion (left panels, effector) or 21 days after depletion to allow for naive/helper CD4 T cells to replenish (right panels, helper). Parasitemias were monitored. Results are representative of 3 separate experiments and data for individual mice are shown. Animals that succumbed to infection. (B) To assess for cytokine secretion in memory CD4 T cells, spleen cells from A/J mice immunized with $10^5$ (AS) prbc in CpG-ODN were purified 12 weeks after immunization, incubated for 96 hours in the presence of homologous (AS) prbc, and cells stained for intracellular IFN-γ and IL-2. Numbers represent percentages of cytokine+ CD4 T cells. (C) To identify the specificity of CD4 T cell memory subsets, IFN-γ secretion and expression of lymph node homing receptor (CD62L) were assessed by harvesting spleen cells from A/J mice immunized with $10^5$ (AS) prbc in CpG-ODN at 12 weeks after immunization. Cells were incubated for 96 hours in the presence of homologous (AS) prbc and stained. Numbers represent percentages of IFN-γ+CD62Llo CD4 T cells. (D) To determine the existence of memory populations, spleen cells from A/J mice immunized with $10^5$ (AS) prbc in CpG-ODN were FACS-sorted into CM (CD44hiCD62Lhi) and EM (CD44hiCD62Llo) populations directly ex vivo. CM and EM cells were then incubated for 96 hours in the presence of homologous (AS) prbc and stained for intracellular cytokine (IFN-γ) and proliferation (BrDU). Numbers represent percentages of CD4 T cells in corresponding quadrants. Results are representative of 3 independent experiments performed.
sufficient but not excessive cytokine signals was desired for the generation of both effector and memory CD4 T cells. It is known, for example, that CD4 effector cells that secrete only IFN-γ have limited capacity to develop into memory cells, while effectors able to secrete IL-2 or IL-2 and IFN-γ are more likely to differentiate into memory cells (44). By combining a strong T cell adjuvant with low doses of killed parasite, we were able to induce a significant proportion of multifunctional parasite-specific CD4 T cells. More importantly, these cells persisted up to 12 weeks after immunization and correlated with protection. It remains to be shown, however, whether similar responses can be induced in humans.

A final consideration for our vaccine was the target population. Here, the development of a formulation able to induce rapid yet long-lived T cell responses is essential if we are to complement current vaccine efforts. This is principally so because in contrast to the induction of protective antibodies that can be inhibited by maternal antibodies, T cell responses can be rapidly elicited (46). While our data indicated that protective responses can be elicited by i.v. injection of 10^8 prbc in normal saline and boosted (i.p.) again 3 weeks later, mice were able to induce a significant proportion of multifunctional parasite-specific CD4 T cells. More importantly, these cells persisted up to 12 weeks after immunization and correlated with protection. It remains to be shown, however, whether similar responses can be induced in humans.

In conclusion, this study demonstrates that a simple vaccine based on low doses of whole killed parasite in CpG-ODN can elicit the robust T cell response usually associated with live (attenuated) vaccines while, at the same time, retaining some of the safety features of a killed (subunit) vaccine. While the hazards involved in the administration of a blood-based product to thousands of otherwise healthy individuals must be addressed, the risk of allograft immunization, blood-borne infections, and parasite purification as well as parasite nonviability are currently being investigated. We envisage that the data provided here will shed some light into the elicitation of cellular responses through whole-organism vaccination and aid in advancing the field of T cell vaccines against other organisms.

**Methods**

**Mice.** Female A/J [H-2b], BALB/c [H-2d], B6 [H-2b], and μMT [H-2b] mice were obtained from the Animal Resources Center (Willetton, Western Australia). μMT mice were bred in-house. All animals were maintained in a specific pathogen-free environment and tested negative for pathogens in routine screening. All animal studies were approved by the Queensland Institute of Medical Research Animal Ethics Committee.

**Parasites.** The rodent malaria parasites used included P. chabaudi AS, P. chabaudi Af, and P. yoelii YM. Parasites were maintained in i.p. passages of 10^8 prbc into naive recipient mice. Challenge infections were performed by i.v. injection of 10^6 prbc (AS or Af) or 10^5 prbc (YM). Parasitemia was monitored by performing Giemsa-stained thin tail blood smears. Disease severity was assessed by performing clinical scores monitoring weight loss (from 0–2), posture (from 0–2), activity (from 0–2), fur texture (from 0–2), and percentage of parasitemia (from 0–2) for a maximal score of 10. Refer to Supplemental Table 1.

**ODN.** Phosphorothioate-modified ODN sequence 1826 containing 2 CpG motifs (underlined: TCCATGACGGTTCCTGAGGTT) and control sequence 1982 (TCCAGGACTCTCTCCAGGTT) were purchased from Sigma-Aldrich.

**Immunization.** Whole-parasite vaccines were prepared using blood from A/J mice infected with P. chabaudi AS. When the parasitemia of infected mice was 30%–40%, blood collected into heparinized tubes was washed (1,000 g for 10 minutes) and resuspended in sterile PBS. Total rbc counts and parasitemias were determined and parasites adjusted to desired concentrations. Subsequently, parasites were killed by repetitive cycles of freeze-thawing, and vaccine formulation was performed as described (34). Briefly, an amount of malaria antigen equivalent to 10^8 to 10^9 prbc was admixed with 50 μg of CpG-ODN or control ODN and aluminium hydroxide (Pierce) and mixed thoroughly to a final volume of 100 μl. Mice were immunized s.c. on the abdomen. Three weeks later, mice were boosted (i.p.) with the same amount of antigen in normal saline and boosted (i.p.) again 2 weeks later with the same amount of antigen. Animals were challenged 2, 8, or 12 weeks after immunization.

**Flow cytometry and serum cytokines.** FACS was performed on a FACSCalibur flow cytometer (BD Biosciences) and analyzed using Summit V4.3 (Dako-
Cytometry) or CellQuest Pro (BD Biosciences) software. Serum cytokines were measured using the Cytometric Bead Array (CBA) for mouse inflammation or Th1/Th2 cytokine kits (BD Biosciences). Acquisition was performed on a FACScan cytometer equipped with CellQuest Pro and CBA Software (BD Biosciences) according to manufacturer’s instructions.

Serum antibody ELISA. Crude parasite antigen from AS, AJ, or YM parasites was prepared by collecting blood, washing in PBS, and incubating in 0.01% (w/v) in saponin (Sigma-Aldrich) at 37°C for 20 minutes. The pellet was washed in PBS, resuspended in 1.5 ml of PBS, and sonicated. Maxi-Sorp Nunc immunoplates (Nalge Nunc) were coated with parasite antigen (10 μg) and incubated with PBS, washed, and incubated with PBS again for 30 minutes. Plates were washed, blocked with 5% (w/v) non-fat milk in PBS for 30 minutes, and then washed again. Serum was added to each dilution (10 μl) and incubated for 2 hours. The plates were washed again and incubated with HRP conjugated anti-mouse IgG or IgM (1:1000) for 1 hour. After wash, plates were incubated with 3,3’-diaminobenzidine (DAB) substrate (100 μl) for 10 minutes and read in a Spectra Max 190 microplate reader at 492 nm. Values were normalized to the mean of the coating coated wells. Results were considered statistically significant when p < 0.05.

Depletions. To deplete IFN-γ, mice received 1 mg i.p. of anti–IFN-γ antibody (clone XMG-6-6) provided by H. Xu, Queensland Institute of Medical Research) on days –1, 0, and 1 relative to day of challenge. IL-12 depletion was performed by injecting 1 mg i.p. of the mAb (clone C17.8) on days –1 and 0. TNF-α blockade was performed by injecting 100 μg i.p. of Etanercept (Wyeth) on days –1, 0, 2, 4, 6, and 8. IL-10 depletion was performed by injecting 1 mg i.p. (clone 1B1.3a) on days –1, 0, and 1. Depletion of pDC was performed by administering 500 μg of anti-pDCA i.p. (Miltenyi Biotec) on days –1 and 1 relative to day of challenge. To deplete NK cells, mice received a single dose of 675 μg i.v. (anti-asialo GM1; Wako Chemicals) on days –2, 0, and 4. For CD8 or CD4 depletion, mice were injected with 1 mg i.p. (clone 58.5.8 or clone GKL.5, respectively) on days –4, –3, –2, –1, and 0. NO depletion was performed by injecting amino guanidine (AG; Sigma-Aldrich) at 50 mg/kg of body weight in 0.5 ml of saline by gastric lavage twice daily on days –1, 0, 2, 4, 6, and 8. In all studies, control mice received parallel doses of rat Ig control (Sigma-Aldrich) from day –4 to day 10 in the same manner as the test antibodies.

Statistics. All comparisons among experimental groups to establish significance were determined by 2-tailed Student’s t test, assuming unequal variances, or 1-way ANOVA followed by Bonferroni’s comparison test. All analyses were performed using the statistical program GraphPad Prism. Results were considered to be statistically significant at P < 0.05.

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