A Comprehensive Preclinical Model Evaluating the Recombinant PRAME Antigen Combined With the AS15 Immunostimulant to Fight Against PRAME-expressing Tumors

Catherine Gérard, Nathalie Baudson, Thierry Ory, Lawrence Segal, and Jamila Louahed

Summary: The PRAME tumor antigen is a potential target for immunotherapy. We assessed the immunogenicity, the antitumor activity, and the safety and the tolerability of a recombinant PRAME protein (recPRAME) combined with the AS15 immunostimulant (recPRAME + AS15) in preclinical studies in mice and Cynomolgus monkeys. Four groups of 12 C57BL6/F1 mice received 4 injections of phosphate-buffered saline (PBS), recPRAME, AS15, or recPRAME + AS15. Immunized mice were injected with tumor cells expressing PRAME (CT26-PRAME) 2 weeks or 2 months after the last injection. The mean tumor surface was measured twice a week. Two groups of 10 monkeys received 7 injections of saline or recPRAME + AS15. T-cell responses were measured by flow cytometry using intracellular cytokine staining (ICS). In C57BL6/F1 mice, repeated injections of recPRAME + AS15 induced high PRAME-specific antibody titers and mostly CD4+ T cells producing cytokines. This immune response was long-lasting in these animals and was associated with protection against a challenge with PRAME-expressing tumor cells (CT26-PRAME) applied either 2 weeks or 2 months after the last injection; these data indicate the induction of an immune memory. In HLA-A02:01/HLA-DR1 transgenic mice, recPRAME + AS15 induced both CD4+ and CD8+ T-cell responses, indicating that this antigen can be processed by the human leukocyte antigen and is potentially immunogenic in humans. In addition, a repeated-dose toxicity study in monkeys showed that 7 biweekly injections of recPRAME + AS15 were well tolerated, and induced PRAME-specific antibodies and T cells. In conclusion, these preclinical data indicate that repeated injections of the PRAME cancer immunotherapeutic are immunogenic and have an acceptable safety profile.

Key Words: PRAME tumor antigen, AS15 immunostimulant, animal models, cancer immunotherapy, tumor protection

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From the GSK Vaccines, Rixensart, Belgium. Reprints: Catherine Gérard, GSK Vaccines, Rue de l’Institut 89, Rixensart 1330, Belgium (e-mail: catherine.gerard@gsk.com).

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lung carcinoma, and breast and cervical cancer cell lines, suggesting that PRAME is immunogenic. The induction of PRAME-specific T-cell immune responses has been reported ex vivo in studies of melanoma and leukemia in humans. In addition, PRAME-specific T cells were highly reactive against different PRAME tumor cell lines or freshly isolated metastatic melanoma and primary leukemic cells, but not against nonmalignant cells. Thus, PRAME-targeted immunotherapy, able to induce strong T-cell responses against tumors, could potentially provide significant benefit to a large number of cancer patients.

Our immunotherapeutic approach is based on the use of a PRAME recombinant protein (recPRAME) combined with the GSK (Rixensart, Belgium) proprietary immunostimulant AS15, to form the PRAME cancer immunotherapeutic (recPRAME + AS15). The PRAME cancer immunotherapeutic is currently in phase I or phase I/II clinical development for the treatment of NSCLC (NCT01159964) and melanoma (NCT01149343), respectively. Given that PRAME expression is not purely restricted to tumor cells, the safety of the potential PRAME-targeted immunotherapy needs to be assessed carefully. Therefore, to support the clinical development of the PRAME cancer immunotherapeutic, we performed nonclinical studies evaluating the immunogenicity and the antitumor activity of recPRAME + AS15 against a PRAME-expressing mouse tumor model. In addition, the safety and the tolerability of recPRAME + AS15 were assessed in Cynomolgus monkeys.

**MATERIALS AND METHODS**

**Experimental Animal Models, Housing, and Husbandry**

Mice studies were ethically reviewed and approved by the GSK Belgian ethical Committee for Animal Experimentation. They were carried out in accordance with the European Directive 2010/63/EU and the GSK Policy on the Care, Welfare, and Treatment of Animals. GSK facilities are AAALAC (Association for Assessment and Accreditation of Laboratory Animal Care) accredited. All efforts were made to minimize suffering: tumors exceeding a maximum allowable size of 17 mm x 17 mm, ulceration, tumor necrosis, convulsion, morbidity, and circling behavior were conditions requiring euthanasia by intraperitoneal injection with a barbituric acid derivative (overdose).

The study in monkeys was conducted in an AAALAC-accredited European Contract Research Organization in compliance with the principles of Good Laboratory Practice (GLP), in particular the OECD Principles of Good Laboratory Practice, and “Arrêté du 14 Mars 2000.” Compliance with animal health regulations, in particular the Council directive 86/609/EEC and the European Directive 2010/63/EU, was also ensured. Serology experiments and the evaluation of T-cell responses in the monkey study were carried out as a post hoc analysis in the GSK facilities.

**Antigen Description, Production, and Purification**

The recPRAME antigen is a 626-aa recombinant fusion protein produced in *Escherichia coli* comprising an N-terminal tripeptide (aa 1–3), containing the truncated methionine and 2 unrelated aa (aspartic acid and proline), aa residues 20–127 of *Haemophilus influenzae* protein D (PD) (aa 4–111), the full-length 509-aa-long PRAME sequence (aa 112–620), and a hexahistidine tag (His) (aa 621–626) enabling protein purification.

The manufacturing process of recPRAME purified bulk consisted of the following key steps: (i) antibiotic-free fermentation of the recombinant E. coli cell culture; (ii) disruption of E. coli cells and extraction of inclusion bodies (IB) containing the recPRAME protein; and (iii) purification of recPRAME from the IB pellet. The purification process involved the following steps: (i) extraction of the PD1/3-PRA-ME-His protein from the IB pellet using the centrifuge system; (ii) carbamidomethylation treatment with iodoacetamide to avoid the formation of disulfide bridges between the cysteine residues; (iii) an immobilized metal ion (Ni²⁺) affinity chromatography (Ni²⁺-IMAC)
procedure (the interaction between the antigen’s His-tag and immobilized Ni$^{2+}$ from the resin is responsible for the reversible capture of the antigen on the resin); (iv) hydroxypatite chromatography of the IMAC eluate to remove E. coli-derived impurities including DNA and endotoxins; (v) ultrafiltration for buffer exchange; and, lastly, (vi) filtration through a 0.45-mm or a 0.22-mm cellulose acetate membrane.

AS15 is an immunostimulant containing 3-O-desacyl-4′-monophosphoryl lipid A produced by GSK, QS-21 Stimulon (Quillaja saponaria Molina, fraction 21; licensed by GSK from Antigenics Inc., a wholly owned subsidiary of Agensys Inc., a Delaware, USA corporation) and a synthetic oligodeoxynucleotide containing unmethylated CG dinucleotides (CpG 7909) in a liposomal formulation. CpG 7909 is a clinical-grade material of CpG 2006 that has been shown to work in both humans and mice and is efficacious in tumor models.

Each dose of recPRAME + AS15 contained either 0.4 or 50 μg of recPRAME for the injection in CB6F1 mice or HLA-A02.01/HLA-DR1 transgenic mice, respectively, and a fixed dose of AS15 (50 μL, 1/10 of a human dose).

Each dose of recPRAME + AS15 for injection in monkeys contained 500 μg of recPRAME and a fixed dose of the AS15 immunostimulant. Each dose corresponded to 1 full human dose and represented an approximately 15–20-fold overexposure of the animals based on a 60–70-kg human and a 3–4-kg monkey.

Control items used were PBS for mice and saline (0.9% NaCl) for monkeys.

**Study Objectives**

The objectives of the studies in mice included the characterization of immune responses and antitumor effects induced by repeated injections of recPRAME alone or recPRAME + AS15 in CB6F1 mice, and the evaluation of immune responses induced by recPRAME + AS15 in HLA-A02.01/HLA-DR1 transgenic mice.

The objectives of the study in monkeys included the evaluation of the immune responses in the context of a repeated-dose toxicity study (CIT35519), using a human dose of recPRAME + AS15.

**The Study Design, Treatment, and Administration**

Studies in mice were conducted at GSK (Rixensart, Belgium) between 2008 and 2011. CB6F1 mice were allocated randomly into 4 groups of 12 mice to receive 4 intramuscular (IM) injections of PBS, recPRAME alone, AS15 alone, or recPRAME + AS15 on days 0, 14, 28, and 42. Immune responses were assessed 2 months after the last injection. Eight immunized mice in each group were injected with 100 μg of CT26-PRAME cells 2 weeks (day 56) or 2 months (day 98) after the last injection. Three extra groups of 12 mice were included and were immunized 4 times 2 weeks apart with recPRAME alone, AS15 alone, and recPRAME + AS15, followed by immune response assessment 2 weeks after the last injection.

Another group of 10 CB6F1 mice were enrolled for tumor specificity experiments. Two groups of mice were immunized with PBS, and 2 with recPRAME + AS15, on days 0 and 14, and then were challenged with either 10$^{6}$ of CT26-PRAME cells or 10$^{5}$ of CT26-MAGE-A3 cells 2 weeks after the last immunization.

HLA-A02.01/HLA-DR1 transgenic mice were allocated to 2 groups of 5 mice to receive 4 injections of PBS or recPRAME + AS15 on days 0, 14, 28, and 42, and their immune responses were assessed 2 weeks after the last injection.

The repeated-dose toxicity study in monkeys was conducted at the CToxLAB (Evreux, France) between 2008 and 2009. The study design and the injection schedule were described previously. In brief, 20 monkeys (10 male and 10 female) were allocated into 2 groups (by sex) using a manual randomization procedure to receive 7 IM injections (500 μL/injection) of saline (control group) or the full human dose of recPRAME + AS15 (treatment group) on days 1, 15, 29, 43, 57, 71, and 85. At the end of the treatment period (3 d after the last injection), the first 3 animals from each group were killed, whereas the last 2 surviving animals from each group were held for a 28-day treatment-free period before being killed.

In all studies, the IM route was selected to mimic the intended route of administration in human therapeutic use.

**The Assessment of Immune Responses**

Blood samples for the assessment of the immune responses were taken from CB6F1 mice on days 56 and 98 and from HLA-A02.01/HLA-DR1 mice on day 56. Blood samples from monkeys were taken at a pretreatment time point, and on days 57, 88, and 113.

**Antibody Responses**

In CB6F1 mice, PRAME-specific antibody levels were measured by an enzyme-linked immunosorbent assay 2 weeks (day 56) or 2 months (day 98) after the fourth injection. Before the addition of sera, the immunoplate was coated with the PRAME antigen overnight at 4°C. After reaction with the sera for 90 minutes at 37°C, a biotinylated sheep whole antibody against mouse immunoglobulins was added for 90 minutes at 37°C. The antigen-antibody complex was then revealed by incubation with a streptavidin-biotinylated peroxidase complex for 30 minutes at 37°C. This complex was then revealed by the addition of tetramethyl benzidine for 10 minutes at room temperature; the reaction was stopped with 0.2 M H$_{2}$SO$_{4}$. Optical densities were recorded at 450 nm.

In monkeys, full details for the determination of the antibody response are presented elsewhere.

Geometric mean titers were calculated with 95% confidence intervals.

**T-cell Responses**

In mice, the frequency of PRAME-specific T cells producing cytokines (at least IFN-γ; IFN-γ single-positive cells or IFN-γ/TNF-α double-positive cells) were detected using ICS and flow cytometry [LSR-II flow cytometer, Becton Dickinson (BD) Biosciences] on individual spleen cells (CB6F1 mice; n = 4) or on peripheral blood lymphocytes (1 pool/group of HLA-A02.01/HLA-DR1 transgenic mice). Cells (10$^{6}$ in each well) were restimulated in vitro for 2 hours at 37°C with either medium (background) or a pool of 123 peptides [15-mer peptides with 11 aa overlap (1 μg/mL)] covering the entire sequence of the PRAME protein in the presence of anti-CD49d and anti-CD28 antibodies (BD Biosciences). Cells were then incubated overnight with Brefeldin A (BD Biosciences).

For cell staining, cell suspensions were washed, and resuspended in 50 μL PBS/1% FCS containing 2% Fc
the frequency of CD4⁺/CD69⁺ T cells producing cytokines and CD8⁺ T cells producing at least IFN-γ in the PBS/1% FCS solution, and then with PBS, and resuspended in FACS buffer before FACS analysis. Live cells were gated (FSC/side-scattered light) and acquisition was performed on ~15,000 CD4⁺ T cells and ~10,000 CD8⁺ T cells for CB6F1 mice. For HLA-A02.01/HLA-DR1 transgenic mice, ~35,000 CD4⁺ T cells and ~5,000 CD8⁺ T cells were acquired. The analyses were performed using the Cellquest or the Diva software (BD Biosciences). The percentage of cells producing at least IFN-γ (single IFN-γ or double IFN-γ/TNF-α-positive cells) in response to antigen stimulation were calculated for the CD4⁺/CD8⁺ gated T-cell population by subtracting the response obtained with medium stimulation from the response upon in vitro stimulation with the PRAME peptide pool. The geometric means of CD4⁺ or CD8⁺ T cells producing at least IFN-γ are shown.

In monkeys, to focus on the activated CD4⁺ T cells, the frequency of CD4⁺/CD69⁺ T cells producing cytokines (at least 1 among IFN-γ, TNF-α, and IL-2) was measured by flow cytometry (LSRII, BD Biosciences) using ICS in peripheral blood lymphocytes before immunization, and after 7 injections (day 113).

### The Assessment of Antitumor Responses

After the immunization of CB6F1 mice and challenge with CT26-PRAME tumor cells, individual tumor growth was recorded twice a week by measuring the product of the 2 main diameters of the tumor during the monitoring phase, starting 7 days after the day of challenge. If a mouse was killed during the study because the tumor size reached the maximum acceptable limit of 289 mm², the value of the last measurement obtained before sacrifice was carried forward to the next time points. The specificity of tumor protection induced by recPRAME + AS15 was assessed twice a week by measuring the tumor surface in CB6F1 mice immunized on days 0 and 14, and challenged with either CT26-PRAME or CT26 cells genetically engineered to express another tumor antigen, MAGE-A3 (CT26-MAGE-A3) on day 28. Although the tumor protection effects are generally more pronounced after 4 compared with 2 immunizations, as certain protection could already be observed after 2 immunizations, for the specificity of tumor response experiments, mice received only 2 immunizations to gain some time and observe the results faster.

### Assessment of the Safety and the Tolerability (Study in Monkeys)

During the study period, each monkey was checked for mortality or signs of morbidity, clinical signs, skin or ophthalmologic reactions, body weight, and food consumption. Rectal temperatures of each animal were recorded and electrocardiography examinations were performed on all animals. Blood samples for the assessment of hematology/biochemistry parameters were taken before the beginning of the treatment period, and on days 2, 4, 15 (before dosing), 84, 86, 88, and 113. A complete microscopic and macroscopic postmortem examination was performed on all monkeys. All assessments were performed as described previously.

### Statistical Analysis

#### Studies on Mice

A comparison of the percentages of tumor-free mice was performed using the Fisher exact test. Pairwise comparisons were adjusted for multiplicity using the Bonferroni method. The tumor size measured at the last time point was the main variable and was analyzed using an analysis of variance model with groups as factors. Each group was compared with the recPRAME + AS15 group, and Dunnett’s adjustment for multiple comparisons was performed. Serum titers were compared after log-transformation using the same approach, and geometric mean ratios between each group and recPRAME + AS15 were derived.

Nonparametric comparisons between the percentages of CD4⁺ T cells were performed after a rank transformation. Dunnett’s adjustment was also applied.

For the study on HLA-A02.01/HLA-DR1 transgenic mice, the analysis of the tumor size measured at the last time point was performed using an analysis of variance model; comparisons of the recPRAME + AS15 with the other groups were adjusted using the Dunnett method.

#### Study on Monkeys

The percentages of CD4⁺/CD69⁺ T cells producing cytokines were compared between the monkey groups in an analysis of covariance model with group as the factor; the percentage of CD4⁺/CD69⁺ T cells producing cytokines without stimulation was used as a covariate in the model.

### Results

#### Studies on Mice

**Immune Responses**

Two weeks after the fourth injection, recPRAME-specific antibody levels were significantly higher in CB6F1 mice immunized with recPRAME + AS15, compared with mice injected with PBS, recPRAME alone, or the AS15 immunostimulant alone; 2 months after the fourth injection, these levels remained significantly higher in mice from the recPRAME + AS15 group compared with the levels in the recPRAME and the AS15 groups (Fig. 1). Similarly, 2 weeks after the fourth injection, the percentage of T cells (mostly CD4⁺ T cells) producing at least IFN-γ (IFN-γ single-positive cells and IFN-γ/TNF-α double-positive cells) was also significantly higher in CB6F1 mice immunized with recPRAME + AS15 compared with the other groups, and remained significantly higher 2 months after injection (Fig. 2). Of note, this response was slightly lower than the one measured 2 weeks after the last immunization, suggesting that the cellular response decreased with time.

Because the PRAME-specific CD8⁺ T-cell response could not be measured in the mouse strain used, the immunogenicity of recPRAME + AS15 was also evaluated in HLA-A02.01/HLA-DR1 transgenic mice. These mice have the advantage to present antigens in the context of the
most common human HLA classes I and II. In these mice, in contrast to CB6F1 mice, both CD4\(^+\) and CD8\(^+\) T-cell responses could be measured 2 weeks after the fourth injection. The percentages of CD4\(^+\) and CD8\(^+\) T cells producing cytokines were substantially higher in the recPRAME + AS15 group, as compared with the PBS group (Fig. 3).

In addition, CD8\(^+\) T-cell responses were also evaluated in an experiment with outbred mice (CD1). Two weeks after the 2 injections of recPRAME + AS15, CD8\(^+\) T cells producing at least IFN-\(\gamma\) were induced in 50% of the mice (Supplementary Fig. S1, Supplemental Digital Content 1, http://links.lww.com/JIT/A395).

Antitumor Responses and Long-term Protection

To assess the capacity of recPRAME + AS15 to induce an immune memory, tumor-free mice immunized with recPRAME + AS15 were challenged with PRAME-expressing tumor cells (CT26-PRAME cells), either 2 weeks (Fig. 4A) or 2 months (Fig. 4B) after the last injection. At both time points of assessment, most mice were protected and remained tumor free (Figs. 4A, B). In contrast, mice immunized with PBS, recPRAME alone, or AS15 alone, and subjected to a similar tumor challenge, had significantly higher mean tumor sizes compared with mice injected with recPRAME + AS15 at 2 weeks after the challenge, with no tumor-free mice detected in these groups (Fig. 4A). At 2 months after the tumor challenge with CT26-PRAME cells, all mice that received recPRAME + AS15 were tumor free, whereas only 3/8 mice were tumor free after the repeated injections of recPRAME alone, and none were tumor free after injections of PBS or AS15 (Fig. 4B). Mean tumor sizes in mice injected with PBS or AS15 were higher compared with mice injected with recPRAME alone (Fig. 4B). These data suggest that long-term antitumor immunity induced by recPRAME + AS15 injections was still able to protect the mice against the tumor challenge 2 months after the last immunization.

In addition, although not a part of the experiments presented in this manuscript, the role of CD4\(^+\) and CD8\(^+\) T cells in tumor protection was assessed in a T-cell deple- tion experiment on CB6F1 mice. After the tumor challenge with PRAME-expressing CT26 tumor cells, and CD4\(^+\) T-cell depletion with the monoclonal anti-CD4 antibody, the mean tumor surface at 1 month after the challenge was markedly higher than that of mice subjected to CD8\(^+\) T-cell depletion, suggesting the crucial role of CD4\(^+\) T cells in tumor protection (Supplementary Fig. S2, Supplemental Digital Content 2, http://links.lww.com/JIT/A396).

The Specificity of Tumor Protection in CB6F1 Mice Induced by recPRAME + AS15

To assess whether the observed tumor protection was PRAME specific, mice immunized with recPRAME + AS15 were challenged either with CT26-PRAME cells or with CT26 cells expressing an irrelevant antigen, MAGE-A3 (CT26-MAGE-A3). After 2 injections of recPRAME + AS15, CB6F1 mice were specifically protected against a challenge with PRAME-expressing tumor cells, but not against a challenge with tumor cells expressing MAGE-A3 (Fig. 5). The mean tumor size of mice immunized with recPRAME + AS15 and challenged with CT26-PRAME significantly lower compared with mice challenged with tumor cells expressing MAGE-A3 or mice receiving PBS and challenged with CT26-PRAME or CT26-MAGE-A3.

Study on Monkeys

The safety and the tolerability of repeated injections of recPRAME + AS15 were assessed in a GLP toxicology study.
study on Cynomolgus monkeys. Extensive toxicology and antibody response data are presented in a separate publication. In brief, injections of recPRAME + AS15 were well tolerated and did not induce any local or systemic toxicity. High PRAME-specific antibody responses were induced in all monkeys immunized with recPRAME + AS15. Repeated injections of recPRAME + AS15 also induced PRAME-specific CD4+ and CD69+ (producing at least 1 cytokine) T-cell responses; after the seventh dose, the mean percentage of T cells producing cytokines was statistically significantly higher in monkeys from the recPRAME + AS15 group, compared with the saline control (Fig. 6). These data show that although PRAME mRNA expression is observed in a few normal tissues in monkeys, repeated injections of recPRAME + AS15 induced humoral and T-cell immune responses with no local or systemic toxicities.

**DISCUSSION**

To support the clinical development of the PRAME cancer immunotherapeutic, we evaluated the immunogenicity of recPRAME + AS15 injections in CB6F1, HLA-A2 HLA-DR1 transgenic mice, and in nonhuman primates (Cynomolgus monkeys). The capacity of repeated injections...
of recPRAME + AS15 to induce antitumor activity was also evaluated in CB6F1 mice.

The evaluation of the immune responses in CB6F1 mice revealed that injections of recPRAME + AS15 triggered both humoral and cellular PRAME-specific immune responses, with significantly higher antibody titers and percentages of T cells producing cytokines (mostly CD4+ ) compared with mice injected with PBS, recPRAME alone, or the AS15 immunostimulant alone. These results suggest that the combination of recPRAME with a strong immunostimulant is necessary to induce such comprehensive immune responses. This is consistent with findings from several clinical and nonclinical studies showing that a combination of tumor antigens delivered as peptides or proteins in formulations comprising 1 or more immunostimulants among MPL, QS-21, and CpG enhances humoral and cellular immune responses.49–59

Our data also show that this immune response is persistent. The level of PRAME-specific antibodies and T cells induced by recPRAME + AS15 measured 2 months after the last immunization was still relatively high. In contrast to the antibody levels, the T-cell response was slightly lower than 2 weeks after the last immunization, suggesting that the immune response might decrease with time and that booster injections may be needed.

However, CD8+ T-cell responses were weak or time inconsistent, probably due to the genetic background of the mice used in these experiments. This is likely because CD8+ T-cell responses can be detected in some CD1 or OF1 outbred mice injected with recPRAME + AS15 (data not shown). Furthermore, on the basis of the results obtained using adoptive T-cell transfer, it is apparent that CD8+ T cells are important to eradicate an existing tumor; however, other effector cells, such as NK cells or CD4+ T cells, may also play a role. In the current manuscript, we clearly showed that the CD4+ T cells are required for tumor protection.

A similar experiment was conducted in H2-knockout mice transgenic for human HLAs (HLA-A02.01/HLA-DR1 transgenic mice).60 These mice express human HLA Class I (HLA-A02.01) and Class II (HLA-DR1) molecules, the most common HLA alleles in the white population61 and represent a unique in vivo experimental model to study the human immune response without any interference with the mouse MHC response. Although the variability of response is generally higher in transgenic mice compared with regular inbred mice, recPRAME + AS15 induced PRAME-specific CD4+ and CD8+ T-cell responses, suggesting that it can potentially induce cellular immune responses in individuals expressing HLA-A02.01 and HLA-DR1.

Because nonhuman primates are considered to be closer to humans than mice in terms of homology to the antigen targeted, it was interesting to characterize the humoral and the cellular immune responses induced by recPRAME + AS15 in these animals. PRAME-specific antibody and T-cell responses were induced in monkeys immunized with recPRAME + AS15. The kinetics of the antibody response in monkeys injected with recPRAME + AS15 revealed that the PRAME-specific antibody levels increased substantially after the fourth injection. The antibody levels did not increase further with any additional immunization, but remained high 3 and 28 days after the seventh injection, indicating that there is no exhaustion of the immune response in these animals after multiple injections.62 Although CD8+ responses were detected in HLA-A02.01/HLA-DR1 transgenic mice, only CD4+ CD69+ T cells were measurable in an ex vivo assay in monkeys after a short in vitro stimulation with a pool of overlapping 15-mer peptides covering the whole PRAME sequence. No CD8+ T-cell responses could be detected under the conditions used. Altogether, immunogenicity results of the studies presented in this article suggest that a vaccine combining a recombinant protein with AS15 is not capable of inducing
to induce a long-term immune memory is a crucial feature of an effective cancer immunotherapy. The immune response induced by 4 injections of recPRAME + AS15 provided a PRAME-specific long-term immune memory able to protect mice against a challenge with PRAME-expressing tumor cells up to 2 months after the last immunization. In mice immunized with recPRAME + AS15, the mean tumor growth remained significantly lower compared with mice injected with PBS, recPRAME alone, or AS15 alone, or was not detectable. These data indicate that both recPRAME and AS15 are required to prevent tumor growth, and that this protection is specific for the PRAME antigen that should be expressed by the tumor. Indeed, after a challenge with a similar tumor expressing another antigen (MAGE-A3), the tumor was not recognized in mice immunized with recPRAME + AS15 and increased in size. In our experiments, we chose a tumor challenge approach, in which tumor-free mice were immunized with recPRAME + AS15 and then challenged with PRAME-expressing tumor cells, as such an approach simulates the adjuvant setting in clinical trials more accurately than would a therapeutic setting approach (ie, first injecting a tumor and then immunizing with recPRAME + AS15). However, unlike mice, patients who underwent surgery and are considered disease free at the time they receive immunotherapy have already been exposed to the tumor. This preexposure is likely to prime the immune system against the targeted antigen or initiate certain immune-suppressive mechanisms. This could possibly impair the efficacy of future immunotherapeutic treatment, although it was not taken into account in the mice experiments. Nevertheless, long-term immunity against the tumor was obtained, which is of particular importance in the context of treating potentially disease-free cancer patients or patients with minimal residual disease undergoing adjuvant treatment and who remain at a high risk of relapse.

In humans, PRAME mRNA expression was detected in a few normal tissues. In nonhuman primates (Cynomolgus monkeys), the PRAME mRNA expression pattern shares some similarities with humans: namely, the antigen can be detected in the adrenal gland, the testis, and the ovaries of both species (unpublished data available to GSK). Furthermore, the monkey PRAME homologous protein presents >90% identity with the human PRAME protein. Thus, before the clinical evaluation of the PRAME cancer immunotherapeutic was launched, the safety of repeated injections of recPRAME + AS15 was evaluated in a GLP toxicity study in nonhuman primates (Cynomolgus monkeys). In this previous study, no signs of inflammation or systemic toxicity were observed after injections of full human doses of recPRAME + AS15 in monkeys. recPRAME + AS15 also induced PRAME-specific antibodies; however, these results should be interpreted with caution due to the higher dose/body mass ratio used compared with the intended administration in humans. To date, only a few reports on the immunogenicity of PRAME in patients with solid tumors have been published. Our results support these very scarce reports on the PRAME tumor antigen as a potential target for immunotherapy of solid tumors.

In conclusion, the results presented in this manuscript indicate that recPRAME + AS15 induced a comprehensive immune response in CB6F1 and HLA-A02.01/HLA-DR1 transgenic mice and provided PRAME-specific long-term protection of CB6F1 mice against a tumor challenge.
recPRAME + AS15 was well tolerated and did not induce any signs of systemic toxicity in nonhuman primates.

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CONFLICTS OF INTEREST/FINANCIAL DISCLOSURES

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All authors are employed by the GSK group of companies. C.G., L.S., and J.L. declare stock ownership in the GSK group of companies. The remaining authors have declared that there are no financial conflicts of interest with regard to this work.

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