Immune Responses following Neonatal DNA Vaccination Are Long-Lived, Abundant, and Qualitatively Similar to Those Induced by Conventional Immunization†

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Virus infections are devastating to neonates, and the induction of active antiviral immunity in this age group is an important goal. Here, we show that a single neonatal DNA vaccination induces cellular and humoral immune responses which are maintained for a significant part of the animal's life span. We employ a sensitive technique which permits the first demonstration and quantitation, directly ex vivo, of virus-specific CD8+ T cells induced by DNA immunization. One year postvaccination, antigen-specific CD8+ T cells were readily detectable and constituted 0.5 to 1% of all CD8+ T cells. By several criteria—including cytokine production, perforin content, development of lytic ability, and protective capacity—DNA vaccine-induced CD8+ memory T cells were indistinguishable from memory cells induced by immunization with a conventional (live-virus) vaccine. Analyses of long-term humoral immune responses revealed that, in contrast to the strong immunoglobulin G2a (IgG2a) skewing of the humoral response seen after conventional vaccination, IgG1 and IgG2a levels were similar in DNA-vaccinated neonatal and adult animals, indicating a balanced T helper response. Collectively, these results show that a single DNA vaccination within hours or days of birth can induce long-lasting CD8+ T- and B-cell responses; there is no need for secondary immunization (boosting). Furthermore, the observed immune responses induced in neonates and in adults are indistinguishable by several criteria, including protection against virus challenge.

Conventional vaccines provide the safest and most effective prophylactic measures against viral diseases. A focused vaccination program has led to the eradication of smallpox and, by the early years of the next millennium, global vaccination should have eliminated poliomyelitis. Despite these successes, infectious diseases remain major contributors to human morbidity and mortality. The World Health Organization mortality figures for 1998 place infections (acute lower-respiratory-tract infections and human immunodeficiency virus) as the third and fourth most frequent causes of adult death worldwide, and infectious diseases kill 2 to 5 million children (<5 years of age) annually. For several reasons, neonates and infants are at heightened risk of viral infection and disease, but vaccination is often delayed in this age group, in part because maternal antibodies can inactivate conventional vaccines, preventing the induction of active immunity in the young. The lack of a widely applicable vaccination strategy which could safely and effectively confer lifelong protective immunity after a single administration early in life has led us to examine the efficacy of DNA vaccination in neonates. We (13) and others (6, 24, 26, 27, 39) have previously shown that DNA vaccination in the first few hours of life can prime both cytotoxic T lymphocyte (CTL) and antibody responses. Furthermore, protective immunity can be induced against some pathogens even when the vaccine recipient carries maternal antibodies (13, 21), validating one proposed advantage of DNA vaccination. However, the longevity, quantity, and quality of DNA-induced neonatal responses have not been characterized.

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MATERIALS AND METHODS

Mice. All mice were BALB/c (H-2d haplotype). Adult animals were purchased from the Scripps Research Institute breeding facility and were bred (by D.E.H.) to generate mice inoculated with DNA immunizations.

DNA immunizations. The plasmid pCMVNP encodes the full length LCMV NP (Armstrong strain); pCMV, the vector control, contains no LCMV sequences. The construction of both of these plasmids has been previously described (47). Plasmids were propagated in Escherichia coli by standard techniques and purified with an Endofree plasmid purification kit (Qiagen, Chatsworth, Calif.) according to the manufacturer’s instructions. Adult mice (6 to 10 weeks of age) received a single 50-μg dose of plasmid DNA dissolved in 50 μl of saline in the tibialis anterior muscle. Neonatal mice (3 days old) were injected in the upper left thigh with 50 μg of plasmid dissolved in 25 μl of saline.

Intracellular cytokine staining (ICCS) and perforin staining. One year after DNA vaccination, splenocytes were harvested and single cell suspensions of splenocytes were prepared. Cells (2 × 10^6) were incubated for 5 h in 200 μl of RPMI 1640 medium containing 50 μg 2-mercaptoethanol, 150 U of recombinant human interleukin-2 per ml, and 2 μg of brehelin A per ml in the presence or absence of 10^{-7} M peptide corresponding to the immunodominant H-2d restricted LCMV NP CD8+ T-cell epitope (NP118–126; sequence RPQASGVYM). After stimulation, the cells were washed with phosphate-buffered saline (PBS) and 5% fetal bovine serum (FBS), stained for 30 min with an anti-mouse CD8+ cytochrome-conjugated antibody (Pharmingen, San Diego, Calif.), washed with PBS and 5% FBS, fixed and permeabilized with Cytofix-Cytoperm (Pharmingen). Intracellular cytokines were stained with fluorescein isothiocyanate-labeled antibody specific for mouse IFN-γ (PharMingen). To detect intracellular perforin, cells were fixed with 2% formaldehyde in PBS and 5% FBS, stained with propidium iodide (PI; Sigma, St. Louis, Mo.) in PBS containing 1% FBS. Perforin antibody (clone P1-8, Kamiya Biomedical Co.) was diluted 1:400 in 0.3% saponin-PBS and added to the cells. Following a 30-min incubation on ice, the cells were washed with 0.1% saponin and incubated with polyclonal goat anti-rat Ig-phycoerythrin (PharMingen) for 30 min. After washing, cells were stained for 30 min with anti-CD8-PE-conjugated (PharMingen) and anti-IFN-γ-fluorescein isothiocyanate. The cells were washed with 0.1% saponin and then 1% FBS. FBS and were stored at 4°C in PBS containing 2% formaldehyde until analysis. In all cases, stained cells were acquired on a FACScan flow cytometer (100,000 to 800,000 events per sample) and analyzed with CELLQUEST software (Becton Dickinson, San Jose, Calif.).

Identification of CTL priming by DNA vaccination. Directly ex vivo, CD8+ memory T cells have little or no detectable lytic activity and therefore cannot be identified in a standard in vitro cytotoxicity assay. However, the CD8+ memory T cells present in a successfully vaccinated mouse can rapidly proliferate in response to virus infection, yielding a lytic response that is detectable by 4 days postinfection (p.i.). In contrast, naive mice must generate a primary cytotoxic T-cell response from a limited number of naive T cells of the appropriate specificity, a process that takes about 5 to 6 days to become detectable in a standard in vitro assay. Therefore, virus-induced and DNA-induced CTL activity detected 4 days p.i. indicates that the mouse was previously successfully vaccinated. Neonatal or adult mice were inoculated with DNA as described above, and 6 months or 1 year later each animal was infected with LCMV (2 × 10^7 PFU; Armstrong strain) intraperitoneally. Spleen cells were harvested 4 days p.i. and half was reserved for later virus titration (see below), and the remainder was assayed for in vitro cytolytic activity by a standard chromium release assay, described elsewhere (41).

Virus titration. Spleen samples were snap frozen by immersion in liquid nitrogen. Each sample was weighed and then homogenized in 1 ml of complete 199 medium (199 medium [Gibco BRL] plus 10% FBS plus penicillin, streptomycin, and β-glutamine). Aliquots of serial dilutions were plated on subconfluent monolayers of Vero cells. After a 1-h incubation period, the virus was removed and the monolayers were overlaid with complete 199 medium containing 0.5% agarose and incubated at 37°C. Three to four days later, the cells were formalin fixed, the agarose plugs were counted, and the monolayers were stained with crystal violet. Plaques were counted, and the data were used to calculate the number of PFU per gram of spleen. The lower limit of detection was approximately 200 PFU per g.

Evaluation of plasmid DNA-induced serum IgG responses. Serum was prepared from coagulated whole blood from mice at 6 months postimmunization. Ninety-six-well plates (Falcon 3912 Microtest III flexible assay plates; Becton Dickinson) were coated with 100 μl of target antigen, comprising purified LCMV in PBS (200 ng of total protein/well). Following overnight incubation at room temperature, the unbound antigen was washed and the wells were blocked with blocking buffer [3% bovine serum albumin (BSA fraction V; Sigma), 0.2% Tween 20 in 1× PBS] and washed with wash buffer (0.1% Tween 20, 1× PBS). The serum samples were serially diluted in blocking buffer and added to the target plate (100 μl/well). Following a 1-h incubation at room temperature, the liquid was aspirated and the wells were washed three times with wash buffer. Total bound IgG, IgG1, or IgG2a was detected with appropriate secondary antibodies conjugated to horseradish peroxidase (1:4,000 dilution, 100 μl per well; Jackson Immunoresearch, West Grove, Pa.). After a 1-h incubation at room temperature, the secondary antibody was removed and the wells were washed three times with wash buffer. Horseradish peroxidase activity was measured by incubation for 30 min with 100 μl of the substrate o-phenylenediamine dihydrochloride in urea hydrogen peroxide (Sigmafast OPD tablets; Sigma), followed by the addition of 1 N HCl (100 μl per well). Absorbance at 492 nm was measured with a Titertek Multiscan Plus (Flow Laboratories, McLean, Va.). For each mouse, the endpoint titer was defined as the first serum dilution at which the mean optical density of triplicate samples did not lie at least 3 standard deviations above the background optical density.

RESULTS

Antigen-specific CD8+ T-cell responses are detectable directly ex vivo 1 year after neonatal or adult DNA vaccination. The exquisite specificity of the T-cell receptor for its cognate peptide/MHC complex, coupled with the ability to detect intracellular cytokines in responding cells, has recently made it possible to detect extremely small numbers of primary or memory antigen-specific T cells within a large and heterogeneous population of nonresponsive cells. Primed antigen-specific T cells initiate cytokine production within minutes of T-cell receptor engagement (37), and in the presence of inhibitors of secretion, these cytokines accumulate within the cell where they can be detected by staining with fluorescently labeled anti-cytokine antibodies, followed by flow cytometry (11). By this ICCS technique, we have analyzed neonatal and adult DNA vaccinees for the presence of LCMV NP-specific CD8+ T cells 1 year after vaccination. Neonatal (3-day-old) or adult (at least 6-week-old) mice were vaccinated either with pCMV (as a negative control) or with pCMVNP. One year later, spleen cells were harvested and virus-specific cells were measured directly ex vivo by ICCS (see Materials and Methods). As positive controls, similar analyses were carried out using cells from an acutely infected mouse 7 days p.i. and from a mouse immunized by conventional means (live virus) 246 days previously (Fig. 1). As shown in Fig. 1, virus-specific cells were abundant in day 7 mice, representing approximately 43% of all CD8+ T cells, a number similar to that seen previously in our laboratory (37) and by others using MHC class I tetramers (25); LCMV-specific memory cells constituted >10% of all splenic CD8+ T cells at 246 days p.i. Most importantly, antigen-specific CD8+ T cells were readily detectable in DNA-immunized mice at 1 year postvaccination, forming discrete populations of CD8+ IFN-γ+ (Fig. 1) which were not present in negative control (pCMV) vaccinees. This is the first identification, directly ex vivo, of CD8+ T-cell responses induced by DNA immunization. Between 0.24 and 1% of CD8+ T cells in neonatal and adult vaccinees were epitope-specific memory cells. There was no significant difference in the percentage of memory cells present in adult compared to neonatal vaccinees. Although the numbers of memory cells are 10- to 40-fold lower than those induced by live-virus immunization, it is remarkable that a year after a single inoculation of DNA within days of birth up to 1 in 100 CD8+ cells remains specific for the encoded antigen. Mice immunized with pCMV showed no discrete CD8+ IFN-γ+ signal after peptide stimulation (Fig. 1), and similar background levels were observed in pCMVNP vaccinees in the absence of peptide stimulation (data not shown).

DNA-induced and virus-induced CD8+ memory T cells produce similar levels of IFN-γ upon antigen contact. Virus-induced and DNA-induced epitope-specific CD8+ T cells produced IFN-γ in response to a 5-h peptide stimulation in vitro (Fig. 1). We estimated the extent of IFN-γ production in both cell populations by determining the mean fluorescence intensity of cells stained for IFN-γ. IFN-γ staining was similar in all antigen-specific cells analyzed (data not shown), suggesting that the mode of immunization (live virus or DNA) has little effect on expression of this effector function.
stimulation (3, 10), low-level lytic activity has been reported (31). We therefore quantitated perforin levels directly ex vivo in CD8+ memory T cells present 1 year after DNA vaccination of adults and neonates and compared them to the levels in virus-induced memory cells and virus-specific CD8+ T cells at the peak of the antiviral immune response. In individual mice, perforin levels were evaluated in IFN-γ (antigen-specific) CD8+ cells and in nonresponding (IFN-γ−) CD8+ cells. As shown in Fig. 2, perforin was detectable in antigen-specific CD8+ T cells following both DNA immunization and virus infection. The highest median level of perforin (205.4) was found during acute virus infection. In all three long-term immune groups (DNA-immunized neonates and adults and virus-immune subjects), the median level of perforin was consistently higher in virus-specific CD8+ T cells than in nonresponding CD8+ T cells from the same mouse. We have not yet determined whether this reflects persistently low levels of perforin in the virus-specific cells or the initiation of perforin synthesis during the 5-h peptide stimulation. Collectively, these data indicate that the age at which animals were vaccinated had no measurable impact on CD8+ T-cell effector functions. Furthermore, the mode of immunization has minimal impact on the effector status of individual CD8+ memory T cells, since virus-induced memory cells contained levels of IFN-γ and perforin similar to those found in their DNA vaccine-induced counterparts.

Accelerated antiviral CTL responses after virus challenge. We have demonstrated that, for at least 1 year postvaccination and even in the absence of boosting, both neonatal and adult vaccinees maintained a population of antigen-specific CD8+ T cells. Following antigen contact, these memory cells produced IFN-γ (Fig. 1) and contained low levels of perforin (Fig. 2). We considered it important to demonstrate that DNA-induced memory cells or their progeny could lyse target cells, since lysis is required for the clearance of LCMV infection (16, 38). Neonates or adults received a single injection of pCMVNP or the vector control plasmid pCMV, and 6 months or 1 year later were challenged with LCMV (Fig. 3). Four days postchallenge, splenocytes from each individual animal were tested in vitro for lytic activity against MHC-matched target cells coated with the immunodominant LCMV NP epitope peptide. As explained in Materials and Methods, the presence of detectable cytolytic T-cell activity 4 days p.i. is indicative of preexisting virus-specific immunity. Six months after DNA immunization, 3 of 4 adult vaccinees and 5 of 5 neonatal vaccinees showed accelerated lytic activity (Fig. 3A and B). CTL activity also was detectable 1 year postvaccination in 4 of 4 adult vaccinees and in 4 of 5 neonatal vaccinees (Fig. 3C and D). In each experiment, CTL activity at 4 days p.i. was also evaluated using splenocytes from several nonimmune mice; the maximum background level of lysis seen in any of these mice is shown (Fig. 3). Thus, DNA vaccination of neonatal or adult mice with no subsequent boosting induces CD8+ memory T cells that are detectable up to 1 year postimmunization. These cells respond to antigen contact by secreting IFN-γ (Fig. 1), contain low levels of perforin (Fig. 2), and mount rapid cytolytic responses upon virus challenge (Fig. 3). This is the first study to demonstrate the lifelong maintenance of CTL following a single neonatal administration of plasmid DNA.

Similar long-lived IgG responses in DNA-immunized neonates and adults. Many vaccines exert their protective effect at least in part by eliciting humoral immune responses. Although long-term antiviral antibody responses have been described after adult and neonatal DNA vaccination (14, 26, 28, 39, 45), there have been few comparative studies of adult and neonatal vaccinees. To address this issue, LCMV-specific IgG serum
antibodies were measured by enzyme-linked immunosorbent assay in individual animals 6 months after DNA vaccination (Fig. 4). For clarity, only the means and standard deviations for each group are shown. All animals that received pCMVNP as either neonates or adults were LCMV seropositive 6 months later (Fig. 4), with endpoint titers of $1:3,200$. The difference in antibody titers between neonatal and adult vaccine groups was not statistically significant ($P < 0.17$). As expected, none of the adult or neonatal pCMV vaccinees showed evidence of LCMV-specific antibodies (Fig. 4). We conclude from these data that a single vaccination with pCMVNP within 3 days of birth leads to the generation of long-lived humoral responses that are similar to those seen in adult vaccinees.

**Similar long-term antibody isotypes in adult and neonatal vaccinees.** Immunization regimens that lead to high levels of IgG2a in adults often result in IgG1-dominated responses in neonates (4, 34). The characteristic IgG1 or IgG2a skewing of the humoral response is established shortly after initial priming and remains unaltered by boosting regimens which lead to alternate skewing among naïve animals (22). To determine the isotype profiles several months after neonatal and adult pCMVNP vaccination, sera were tested for the presence of LCMV-specific antibodies (Fig. 4). We conclude from these data that a single vaccination with pCMVNP within 3 days of birth leads to the generation of long-lived humoral responses that are similar to those seen in adult vaccinees.

**Long-term DNA-induced immunity protects against virus challenge.** Viral clearance and protective immunity against LCMV are conferred by CD8$^+$ CTLs. We have demonstrated the long-term maintenance of LCMV-specific CD8$^+$ T cells (Fig. 1) and CTLs (Fig. 3) in mice immunized with pCMVNP as neonates and adults. To establish whether these long-lived responses were biologically relevant, we next evaluated their ability to limit LCMV multiplication at 6 months and 1 year postvaccination (Fig. 6). The mice were infected with LCMV and 4 days p.i., their splenic titers were determined; these are shown for individual mice (log$_{10}$ scale). To demonstrate...
the strong correlation between protective immunity and the presence of CTLs, the bars in Fig. 6 are color coded; white bars represent animals in which lytic CTL activity was detectable (Fig. 3), while black bars denote animals devoid of detectable CTLs. Virus which was below the level of detection in five mice is indicated by an asterisk in Fig. 6; all of these mice were positive for CTLs (Fig. 3) and therefore had been infected and had cleared the virus.

Six months after DNA immunization (Fig. 6A), 7 of 8 pCMVNP vaccinees (3 of 4 adult and 4 of 4 neonatal subjects) showed >1,000-fold reductions in LCMV titer 4 days after LCMV challenge. A similar reduction was seen in 8 of 9 pCMVNP vaccinees (4 of 4 adult and 4 of 5 neonatal subjects) 1 year postimmunization (Fig. 6B). None of the mice immunized with pCMV showed a reduction in titer approaching these levels. There is an excellent correlation (26 of 27 DNA-immunized mice) between CTL activity and protective immunity. Only one mouse with detectable CTLs did not meet the stringent criterion of a 1,000-fold reduction in titer (Fig. 6B); however, infectious virus was markedly (~95%) reduced, even in this animal.

Therefore, DNA vaccination within the first 3 days of life primes antigen-specific effector T cells (Fig. 1) and CTLs (Fig. 3) and is as effective as immunization in adulthood for priming long-lived protective antiviral immunity (Fig. 6).

**FIG. 3.** Accelerated antigen-specific lytic CD8⁺ T cell responses at 6 months and 1 year after DNA vaccination of neonatal or adult mice. Adult (▼, panels A and C) or neonatal (●, panels B and D) mice received a single 50-μg injection of pCMVNP. Six months (panels A and B) or 1 year (panels B and D) later, mice were infected with LCMV, and 4 days thereafter were evaluated for the presence of lytic CTL activity. Each line represents the percent specific ³¹Cr release for an individual mouse, determined at the indicated effector-to-target cell (E:T) ratio. For each experiment, the nonimmune mouse (either pCMV inoculated or given no DNA) showing the highest level of background lysis at day 4 p.i. is shown (open squares).

**DISCUSSION**

Neonates are profoundly sensitive to a variety of viral infections, and early antiviral immunization is vital to reduce the incidence and severity of viral diseases in this age group. However, early postnatal immunization is not entirely without risk. Live viral vaccines at doses administered to adults might overwhelm the neonatal immune system and cause inadvertent pathological consequences. Killed vaccines, although safer, often do not stimulate the strong responses necessary to provide long-lived protective immunity, and they are rather ineffective in stimulating the CD8⁺ T-cell responses which are critical for protection against a wide variety of intracellular pathogens. In addition, numerous examples indicate that neonates and adults may respond differently to immunological challenges (33). For these reasons, few vaccines are given in the neonatal period; most are withheld until infancy, and repeated booster immunizations are administered throughout infancy and early childhood to ensure adequate levels of protection. A preferable situation would be the single administration of a safe vaccine immediately after birth, capable of providing a high level of protective immunity that could be maintained throughout life in the absence of boosting. DNA vaccines are an attractive alternative to the immunization of neonates, and in our previous study we documented the acquisition of short-term humoral and cell-mediated immune responses against LCMV.
after a single administration of pCMVNP within hours of birth. These responses were capable of successfully controlling a challenge infection until at least 6 weeks after birth (13). Protective immunity after neonatal DNA vaccination has also been observed with rodent models of influenza virus (5, 6), herpesvirus (21) and rabies virus (39, 40) infections and in a primate model of hepatitis B virus (27). In contrast, plasmid vaccination against the circumsporozoite antigen of malaria induces tolerance when administered to neonates and protective immunity when given to adults (15, 24).

The present study had several goals: first, to determine if humoral and cellular immunity primed by neonatal DNA vaccination could be maintained for an extended period of time in the absence of revaccination; second, to quantitate, directly ex vivo, CD8^+ T-cell responses at 1 year postvaccination; third, to compare the effector status of long-term DNA-induced memory cells with the status of memory cells induced by conventional (live-virus) immunization; and fourth, to evaluate the protective efficacy of these memory T cells. In all cases, we carried out parallel analyses in neonatal and adult vaccinees to determine if the age at DNA immunization was an important variable for any of the responses being measured. Our data (Fig. 1) show that antigen-specific CD8^+ T cells can be maintained in vivo without revaccination for at least 1 year (roughly half of the lifespan of a mouse). The relative proportions of antigen-specific CD8^+ effector T cells were similar in neonatal and adult vaccinees. In both cases, ~0.5% to 1% of CD8^+ T cells were NP specific 1 year after DNA vaccination, a number only ~2-fold lower than that observed 1 month after vaccination with a recombinant vaccinia virus expressing LCMV NP (1 to 2%, our unpublished data). NP-specific CD8^+ memory cells constituted ~10% of CD8^+ T cells at 246 days after live virus immunization, a figure consistent with that described with peptide tetramer staining (25). The memory cells induced by DNA immunization appeared qualitatively similar to those induced by conventional immunization, having comparable levels of IFN-γ and perforin. Furthermore, DNA vaccine-induced cells permitted the generation of an accelerated CTL response detected by target cell lysis (Fig. 3), and most importantly, the immunity induced by neonatal DNA vaccination could confer protection against virus challenge 1 year later (Fig. 6). Thus, the mode of immunization affected the quantity, but not the quality, of CD8^+ memory T cells.

Analysis of the anti-NP humoral response 6 months postimmunization revealed that neonatal vaccinates had levels of serum IgG that were indistinguishable from those observed in immunized adults (Fig. 4), confirming that early postnatal exposure to a plasmid-expressed antigen can prime a B-cell response as efficiently as can exposure in adulthood. Since the half-life of antibodies in murine serum is approximately 10 to 14 days (36), high antibody titers at 6 months postvaccination are indicative of ongoing antibody synthesis, presumably by long-lived plasma cells (36). It has been previously reported that neonates show a bias towards an IgG1 antibody response after protein immunization, which is overcome by DNA immunization (22). Our results confirm the presence of a balanced antibody response after neonatal DNA immunization similar to that seen in vaccinated adults and show that this is maintained well into adulthood.

While the above data show conclusively that DNA immunization of neonates and adults can induce long-term immunity,
the underlying mechanisms remain unclear. For example, might the longevity of our DNA-induced memory cells depend on the continued production of NP antigen from persistent plasmid DNA? Plasmid DNA inoculation induces a local antigen-specific inflammatory response (12, 46) which leads to destruction of the transfected muscle fibers (9) and therefore to the eradication of many of the antigen-expressing cells. It is probable that a similar fate would befall transfected cells of all types, including antigen-presenting cells (8). Thus it seems unlikely that significant quantities of NP would persist in the vaccine. Furthermore, several studies—mainly in the LCMV model—suggest that antigen is not required for the maintenance of memory cells (1, 2, 20). However, while we doubt the importance of persistent NP in driving the DNA-induced memory cell response demonstrated here, we cannot altogether exclude the possibility; one report describes the in vivo detection of plasmid DNA and expression of the encoded luciferase protein a remarkable 19 months after DNA inoculation (43).

The data presented here indicate that long-lived protective immunity can be induced by neonatal vaccination with plasmid DNA. By all of the immunological criteria examined, these long-lived responses appear to be efficiently maintained in the absence of boosting and are quantitatively and qualitatively indistinguishable from the responses induced in adults. Furthermore, the DNA vaccine-induced responses are relatively abundant and are qualitatively similar to those induced by conventional (live-virus) immunization. Thus, DNA vaccines may offer a safe and attractive solution to the problems associated with neonatal vaccination.

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REFERENCES