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The efficacy of DNA vaccination is enhanced in mice by targeting the encoded protein to dendritic cells

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DNA vaccines promote an immune response by providing antigen-encoding DNA to the recipient, but the efficacy of such vaccines needs improving. Many approaches have considerable potential but currently induce relatively weak immune responses despite multiple high doses of DNA vaccine. Here, we asked whether targeting vaccine antigens to DCs would increase the immunity and protection that result from DNA vaccines. To determine this, we generated a DNA vaccine encoding a fusion protein comprised of the vaccine antigen and a single-chain Fv antibody (scFv) specific for the DC-restricted antigen-uptake receptor DEC205. Following vaccination of mice, the vaccine antigen was expressed selectively by DCs, which were required for the increased efficacy of MHC class I and MHC class II antigen presentation relative to a control scFv DNA vaccine. In addition, a DNA vaccine encoding an HIV gag p41–scFv DEC205 fusion protein induced 10-fold higher antibody levels and increased numbers of IFN-γ–producing CD4+ and CD8+ T cells. After a single i.m. injection of the DNA vaccine encoding an HIV gag p41–scFv DEC205 fusion protein, mice were protected from an airway challenge with a recombinant vaccinia virus expressing the HIV gag p41, even with 1% of the dose of nontargeted DNA vaccine. The efficacy of DNA vaccines therefore may be enhanced by inclusion of sequences such as single-chain antibodies to target the antigen to DCs.

Introduction
DNA vaccination is at the forefront of efforts aimed at developing vaccines against challenging infectious diseases, including HIV AIDS as well as emerging strains of influenza and SARS (reviewed in refs. 1–4). Innovations in vaccine design have improved the immunogenicity of DNA vaccines, leading to licensure of 2 candidate DNA vaccines for nonhuman applications (reviewed in refs. 5, 6). However, in humans and experimental animals, repeated and high doses of DNA must be injected.

A fundamental initial step in vaccination is that the vaccine proteins need to be taken up, processed, and presented by DCs, which are antigen-presenting cells that initiate and control many aspects of the immune response. Following intradermal or i.m. injection of a plasmid DNA vaccine in mice, the encoded gene is expressed in transfected keratinocytes and myocytes at the site of inoculation (7) as well as a small number of DCs (8–12). Keratinocytes and myocytes lack MHC class II and costimulatory molecules, which renders them poorly effective at presenting antigen and priming naive immune cells (reviewed in ref. 13), and they do not have ready access to T cells in lymphoid tissues, as is the case for DCs (14, 15). Therefore, it is thought that immune priming begins through the function of a few transduced DCs and that antigen secreted from other long-lived transfected cells then boosts immunity (16–23). We hypothesized that the weak immunogenicity of DNA vaccines could be overcome by directly targeting the encoded vaccine protein to DCs, particularly during the later stages of immunization.

Recent studies have demonstrated that antigens can be targeted selectively to DCs in vivo when antigen is incorporated into an antibody against the DC endocytic receptor, DEC205/CD205 (24). This targeting is accompanied by efficient antigen processing and presentation on MHC class I and II products as well as strong and protective T cell immunity (25, 26). Demangel et al. reported on the capacity of a single-chain antibody to murine anti-DEC205 (scDEC) to target DNA vaccine antigens in mice (27). They observed approximately a 2-fold increase in B and T cell responses to a mycobacterial antigen but did not establish whether targeted delivery of vaccine to DCs was taking place or whether the dose of DNA vaccine could be reduced and maintain efficacy. As a result, it is not clear whether the targeted delivery to DCs overcomes some of the key obstacles to current DNA vaccines.

We have evaluated the incorporation of a scDEC antibody into DNA vaccines. We will show that this approach greatly enhances antigen presentation by DCs in lymphoid tissues relative to nontargeted DNA vaccine in mice, and concomitantly, that single and much lower doses of DNA are effective in inducing immunity and protection.

Results
Construction and expression of scDEC DNA vaccines. To better target DNA vaccines to DCs, a eukaryotic expression plasmid was constructed to encode scDEC fused to a vaccine protein. DEC205

Nonstandard abbreviations used: CHO, Chinese hamster ovary; DT, diphtheria toxin; scDEC, single-chain antibody to murine DC205; scFv, single-chain Fv antibody; TAP, transporter for antigenic peptides.

Conflict of interest: R.M. Steinman and M.C. Nussenzweig have a financial interest in Cellex Therapeutics, which is developing antibody-based vaccines that target dendritic cells.

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is an antigen uptake receptor on DCs, and the open reading frames for the variable regions of the light and heavy chains of the parental anti-DEC205 antibody were fused to the protein sequence. A control plasmid from a nonreactive antibody was also constructed with different variable light- and heavy-chain V regions (Figure 1A). cDNAs for OVA, HIV gag p41 (anti-p41) and p24 (anti-p24), and GFP were cloned in frame with the scDEC or the control antibody. To verify that the scDEC protein retained the antigen-binding activity of the parental antibody, we measured specific binding to Chinese hamster ovary (CHO) cells stably expressing murine DEC205 receptor. Only supernatants from scDEC-OVA–transfected 293T cells (Figure 1, B and C) showed binding to DEC205-expressing CHO cells. When we compared the DEC205-binding efficacy of the scDEC-OVA relative to the parental bivalent antibody (DEC-OVA) compared with scDEC-OVA, we found that the majority of TCR transgenic T cells went into cell cycle, i.e., both CD8+ MHC-I–restricted OT-I cells and CD4+ MHC-II–restricted OT-II cells. The OT-I cells responded to the DNA vaccine at all doses, 3–100 μg, whereas the OT-II cells only showed proliferation at the higher doses of DNA. This reflects the fact that OT-I cells are unusually efficient in being able to recognize low pM doses of specific OVA peptide. In contrast, a 100-μg DNA dose of scControl OVA DNA was required to induce OT-I responses that were comparable to those obtained with 3 μg of scDEC-OVA, and no proliferation was observed for CD4+ OT-II T cells at any dose of scControl OVA DNA (Figure 2A).
To prove that MHC class I presentation by scDEC-OVA was DEC205 dependent, we performed similar experiments with DEC205-deficient (DEC–/–) mice and observed a lack of T cell proliferation (DEC–/–; Figure 2B). To prove that MHC class I presentation by the scDEC-OVA was transporter for antigenic peptides–dependent (TAP-dependent), we verified that TAP-deficient (TAP–/–) mice could not induce OT-I T cell proliferation (Figure 2B).

To assess the duration of antigen presentation after scDEC-targeted DNA vaccination, B6 mice were immunized with 30 μg DNA, and CFSE-labeled OT-I TCR transgenic T cells were injected i.v. at day 7. Draining lymph node cells were harvested and cell division was evaluated by flow cytometry in CD4+ OT-I cells with Vα2 T cell receptor (20.1) antibody and in CD8+ OT-I cells with Vβ5.1/5.2 T cell receptor (MR9-4) antibody. (A) Control mice received similar amounts of the scControl–OVA DNA vaccine. (B) As in A, but the experiments were performed with DEC–/–, TAP–/–, or WT B6 mice using 30 μg of each DNA vaccine. Representative results from 3 experiments are shown. (C) Longevity of in situ antigen presentation after vaccination with 30 μg DNA, then evaluation by antigen presentation to 2 × 10^6 OT-I T cells injected at the indicated time points. (D) scDEC-OVA targets antigens to DCs after DNA vaccination. Mice were immunized with 30 μg of the indicated scFv DNA vaccines or backbone vector. 4 days later, CD11c+ and CD11c– cells were isolated and cocultured with CFSE-labeled OT-I T cells at the indicated APC to T cell ratios to determine CFSE dilution in vitro at day 7.5 of the experiment. (E) CD11c-DTR mice were vaccinated with 30 μg scDEC-OVA or backbone vector and were depleted of CD11c+ DCs 4 days later by injection of 4 ng/g of DT (DT+); then CFSE-labeled OT-I T cells were injected to monitor antigen presentation as in A. C–E are representative of 2 experiments.

Figure 2
Antigen presentation to OVA-specific T cells following in vivo targeting of DCs with OVA DNA fused to a single-chain antibody to DEC205. (A) Mice were immunized i.m. with graded doses of DNA, and 4 days later, 2 × 10^6 CFSE-labeled CD8+ OT-I or 3 × 10^6 CD4+ OT-II TCR transgenic T cells were injected i.v. At day 7, draining lymph node cells were harvested and cell division was evaluated by flow cytometry in CD4+ OT-I cells with Vα2 T cell receptor (20.1) antibody and in CD8+ OT-I cells with Vβ5.1/5.2 T cell receptor (MR9-4) antibody. (A) Control mice received similar amounts of the scControl–OVA DNA vaccine. (B) As in A, but the experiments were performed with DEC–/–, TAP–/–, or WT B6 mice using 30 μg of each DNA vaccine. Representative results from 3 experiments are shown. (C) Longevity of in situ antigen presentation after vaccination with 30 μg DNA, then evaluation by antigen presentation to 2 × 10^6 OT-I T cells injected at the indicated time points. (D) scDEC-OVA targets antigens to DCs after DNA vaccination. Mice were immunized with 30 μg of the indicated scFv DNA vaccines or backbone vector. 4 days later, CD11c+ and CD11c– cells were isolated and cocultured with CFSE-labeled OT-I T cells at the indicated APC to T cell ratios to determine CFSE dilution in vitro at day 7.5 of the experiment. (E) CD11c-DTR mice were vaccinated with 30 μg scDEC-OVA or backbone vector and were depleted of CD11c+ DCs 4 days later by injection of 4 ng/g of DT (DT+); then CFSE-labeled OT-I T cells were injected to monitor antigen presentation as in A. C–E are representative of 2 experiments.
cessful antigen presentation. We could only detect presentation of the OVA peptide and consequent proliferation of the OT-I TCR transgenic T cells from cocultures with CD11c+ DCs and not with CD11c-DCs (Figure 2D). In addition, no proliferation was observed when we cocultured OT-I TCR transgenic T cells with CD11c- DCs from DEC205-deficient mice, indicating that DEC205 was necessary for the uptake of the targeted vaccine by CD11c+ DCs (Figure 2D). Finally, CD11c+ DCs isolated from mice treated with the control DNA vaccine only induced background levels of proliferation (Figure 2D). We conclude that the targeted DNA vaccine delivered antigen to CD11c+ DCs in vivo, in a DEC205-dependent manner.

To investigate the role of DCs in targeted DNA vaccination in vivo, we used a transgenic mouse (CD11c-DTR/GFP) that allows for conditional ablation of CD11c+ DCs in vivo after administration of diphtheria toxin (DT) (30). These mice carry a transgene encoding a fusion protein of the DT receptor and GFP under control of the mouse CD11c promoter, allowing for depletion of DCs in vivo upon injection of DT. The CD11c-DTR/GFP mice as well as transgene-negative littermates and WT C57BL/6 mice were vaccinated with 30 μg DNA i.m. and injected with DT on day 4 (4 ng/g body weight of DT i.p.). 24 hours later, the mice each received 3 × 10^6 CFSE-labeled TCR transgenic OT-1 T cells. Three days later, draining lymph node lymphocytes were evaluated for proliferation of the T cells. As shown in Figure 2E, temporal ablation of CD11c+ DCs resulted in marked reduction in CFSE dilution. In contrast, WT littermates lacking the CD11c-DTR/GFP transgene and treated similarly with DT behaved like WT B6 mice, with virtually all the OT-I T cells going into cell cycle in mice following targeted DNA vaccination. Thus, the ablation of DCs hampered the ability of the targeted vaccine to induce OT-I T cell responses in vivo.

Targeted DNA vaccination improves antibody responses. To evaluate whether scDEC-OVA vaccination would improve B cell responses, we injected B6 mice with graded doses of the vaccine or scControl OVA and measured anti-OVA antibodies. Another control group received a dose of the backbone (empty) vector pcDNA3.1 equivalent to the highest dose of DNA vaccine (data not shown). At all doses of DNA, mice primed with scDEC-OVA DNA produced OVA-specific IgG1 antibody titers at least 10 times higher than those of groups primed with scControl OVA (Figure 3A). IgG2b and IgG2c titers were lower than IgG1, and the differences between scDEC-OVA and scControl-OVA were less than observed for IgG1 responses (Figure 3A).

We next tested to determine whether a single dose of scDEC-HIV gag p41 could induce better gag-specific antibody responses in BALB/c mice (Figure 3B). Six 8-week-old mice were injected once with 3 and 9 μg doses of DNA i.m. in combination with electroporation. Five weeks later, serum was tested for HIV gag p41-specific antibodies. Again, at least 10-fold better antibody titers were detected after a single injection of just 3 μg of the targeted vaccine, and the increase in titers was greatest for the IgG1 subclass of antibodies. Antibody responses due to the backbone vector (pcDNA3.1) alone were insignificant (data not shown). Thus, DEC205 targeting enhances the humoral responses to a DNA vaccine.

scDEC targeting improves HIV gag-specific T cell responses. To examine T cell immunity to HIV gag, we immunized mice with graded doses of scDEC gag p41 and control DNA vaccine and measured T cell responses with a library of 15-mer “mimotope” peptide staggered every 4 amino acids along the HIV gag p41 sequence (26). Multiparametric intracellular cytokine assays showed that Cxβ6 F1 mice generated T cell responses after vaccination with low doses of the targeted vaccine (0.3 and 3 μg DNA) whereas controls did not (Figure 4A). In contrast, at the higher doses that are typically used during DNA vaccine experiments in mice (30 μg), both scControl and scDEC vaccines performed similarly in inducing IFN-γ–producing, gag-specific T cells. Similar findings were made with ELISPOT assays, including the fact that no IL-4– or IL-10–producing cells were detectable (data not shown).

T cells producing multiple cytokines appear to be especially effective in protective immunity (31–33). We therefore examined the quality of the T cell responses induced by DNA vaccination in terms of the capacity of gag-specific T cells to secrete multiple cytokines such as IFN-γ, TNF-α, and IL-2. Fourteen days after immunization, scDEC-targeted vaccination preferentially induced gag-specific polyfunctional CD4+ T cells, including triple cytokine producers, at all doses of the vaccine tested (Figure 4B). In contrast, triple producing cells were not detected after 1 dose of the scControl DNA vaccine. More double cytokine–producing CD4+ T cells
were also induced at all doses of the scDEC vaccine (Figure 4B). The CD4+ T cell responses to scDEC vaccine were broad since reactivity was noted to peptides in p24 pools 1 and 4 and the p17 pool (data not shown), as previously observed with anti-DEC antibody immunization (26). Responses to the backbone DNA vector were insignificant (data not shown). Likewise, at 28 days after vaccination (Figure 4C), scDEC–gag p41 induced cytokine-producing CD4+ and CD8+ T cells, including T cells with a predominant production of IL-2 as well as some double cytokine–producing T cells. These T cells expressed high levels of CD44, as is characteristic for memory T cells (not shown). These data indicate that scDEC targeting of low doses of DNA improves the induction of broad T cell–mediated immunity.

Targeted DNA vaccination elicits protection at a mucosal surface. To determine whether targeted vaccination produces long-lasting protective immunity at a mucosal surface, we challenged BALB/c and Cx6B6 F1 mice intranasally with recombinant vaccinia-gag virus 3 months after a single dose of targeted and control vaccine, using a wide range of DNA doses, i.e., 0.3, 3, and 30 μg per mouse. Control mice vaccinated with the empty vector were not protected relative to PBS-injected mice, and they lost weight and developed high lung virus titers over 7 days (10^7–10^8 PFU/ml per lung; data not shown). In contrast, all mice receiving graded doses of the scDEC-targeted vaccine were protected, even with a single dose of 0.3 μg DNA, whereas only mice vaccinated with the highest 30-μg dose of the scControl vaccine showed protection (Figure 5A).
Data from a total of 4 different experiments, 2 each in BALB/c and BALB/c × C57Bl/6 F1 mice, are summarized in Figure 5B. It is evident that scDEC vaccine (Figure 5B) provides better and long-lived protective immunity at lower (0.3 and 3 μg) but not higher (30 μg) doses of DNA than the scControl (Figure 5B). Thus, the targeting of protein within a DNA vaccine to DCs in situ enhances protective efficacy by almost 100-fold.

Discussion
Our results directly show that DNA vaccines can be configured to improve antigen presentation by DCs in vivo and that this in turn improves immunity and lowers the dose of DNA vaccine that is required. Multiple strategies have been developed to augment immune stimulation following DNA vaccination (reviewed in refs. 34–36). One approach is to include plasmids that encode different types of immunostimulatory molecules, such as cytokines (IL-2, IL-12, GM-CSF, flt-3L), chemokines (MIP-1α, SLC), and membrane costimulators (CD80, CD86) (37–43). Another approach is for the DNA vaccine to encode a fusion protein that allows the vaccine to target to specific immune molecules that could enhance immunization, e.g., MHC class II, CTLA-4, Fc receptors, and chemokine receptors (refs. 44–50; reviewed in ref. 6). A third, newer approach has been for the DNA vaccine to target pattern recognition and “alarmin” receptors to increase triggering of the innate immune system, e.g., DEC205/CD205, TLR5, and HSP70 (27, 51, 52). This extensive literature has repeatedly shown that the immune response to DNA vaccines can be enhanced, but the findings have yet to
prove that enhanced presentation of vaccine antigens by DCs is being achieved or that the dose of DNA required for vaccination can be greatly reduced.

To overcome these obstacles to improved DNA vaccines, we made plasmids in which the vaccine protein was fused to a single-chain antibody that could target the DEC205 receptor on DCs following secretion from transfected muscle. The rationale originated in prior research showing that hybrid antibodies carrying vaccine protein to target the DEC205 antigen uptake receptor on DCs could greatly increase antigen presentation and immune responses (24–26, 53–55). Here, we found that “hybrid” DNA vaccines, composed of scDEC205 or a single-chain Fv antibody (scFv) control Ig fused to another protein, were both secreted well from transfected cells. However, the introduction of an scDEC205 antibody sequence into the DNA vaccine led to improved delivery of vaccine protein to CD11c+ DCs in situ relative to scFv control Ig and resulted in a large increase in the efficacy with which the vaccine induced T cell immunity and protection against a challenge infection. Demangel et al. have previously reported on the use of scDEC205 to improve DNA vaccination (27). However, their study did not demonstrate targeting to DCs in lymphoid organs in a receptor-dependent manner, and they did not report a marked reduction in the dose of vaccine DNA that could elicit immunity, including protective immunity. Our data indicate that the scDEC vaccine greatly enhances, by about 100-fold, the delivery of vaccine protein to DCs in vivo and allows for a corresponding reduction in the dose of DNA required to induce protective immunity.

Transfected muscle cells at sites of DNA injection may contribute to vaccine immunogenicity (21, 56). However, we feel that our results are best explained by the secretion of scFv anti-DEC vaccine from transfected muscle or other cells and then presentation by DCs targeted by the anti-DEC. This event is termed cross presentation because muscle cells synthesize the vaccine antigen, which then crosses into DCs for processing and presentation to T cells. Cross priming acts as a major mechanism for the induction of T cell responses following DNA vaccination (16, 17, 19, 20, 22, 23).

Hybrid αDEC antibody protein vaccines (24–26, 53–55) represent one strategy for preparing DC-targeted vaccines, but the extension of this concept to DNA vaccines is limited by the need to coexpress the heavy and light chains in vivo to form a functional antibody within specialized antibody-producing B cells. The use of scFv αDEC-antigen fusion plasmids allows the anti-DEC antibody to be secreted in tandem with the vaccine fusion protein from transfected non-B cells, and it allows for prolonged and greatly improved presentation of vaccine by DCs in lymphoid organs (Figures 1 and 2). In addition, we used an electroporation instrument to facilitate the delivery of DNA following i.m. injection, as has been reported by many others (e.g., refs. 29, 57–60). A newer finding is that electroporation may also make more DNA available for detection by intracellular DNA sensors, thereby activating the production of cytokines as an innate reaction (61). Thus, DNA electroporation in combination with DC targeting could provide a much more efficient platform for developing clinically effective DNA vaccines.

DCs are known to express several receptors with the potential to enhance antigen uptake (62). To date, the DEC205/CD205 receptor has been studied in the greatest detail in vivo and shown to deliver many different proteins for presentation on both MHC class I and II products (24–26, 53, 55). DEC205 is expressed at high levels on a large fraction of DCs in the T cell areas of lymphoid organs, including human lymphoid tissues (15). However, other uptake receptors such as DCIR2 are preferentially expressed on different DC subsets with distinct antigen-presenting properties, and these too need to be assessed in single-chain antibody–based targeting vaccines (54). DEC205 targeting may be advantageous because it targets a subset of DCs in mice that is specialized to process antigens on MHC class I but also mediates improved immunization of MHC class II–restricted CD4+ helper T cells (25, 26, 53).

Unlike protein-based DEC205 vaccines, which require the coinjection of additional DC maturation stimuli to induce T cell responses (24–26, 53, 55), scDEC targeting in a DNA format induces immunity with no additional external adjuvant (27), possibly because the DNA itself provides some signals for DC maturation (61). In this study, we constructed several single-chain antibody-antigen fusion proteins that maintained the DEC205-binding specificity of the parental antibody. Injection of the scDEC DNA vaccine i.m. in combination with electroporation allowed antigen to be targeted to DCs in a DEC205-dependent fashion, resulting in a marked increase in presentation of antigen on MHC class I and II products. It needs to be determined whether the efficacy of scDEC-targeted vaccines can be further improved by the inclusion of additional stimuli that increase DC numbers and maturation state and/or improve the function of the responding T cells.

In summary, in vivo targeting of DNA-encoded vaccine proteins to DCs with scFv to DEC205 has the following features: (a) a precisely defined ratio of antigen to targeting antibody; (b) absence of binding via Fc receptors; (c) relative ease of large-scale production of DNA vaccine; and (d) increased immunogenicity (antibody as well as CD4+ and CD8+ T cells) because of the combination of DC targeting and stimulatory sequences within the vaccine. We propose that DC targeting via DEC205 or perhaps other DC receptors in the context of DNA vaccination will significantly enhance efficacy and may be particularly valuable for protection against diseases such as AIDS, in which broad-based immune responses will be helpful.

Methods

Vector construction. To generate scDEC-OVA plasmids, the cDNA of the heavy- and light-chain variable regions of the monoclonal antibody NLDC145 to DEC205 were amplified by PCR from the plasmids DEC-heavy HEL and DEC–κ HEL (24). Using overlap extension PCR with primers encoding an interchain linker (GGGGS), TCCCTCGAGGCGGTCGGTGAGGAGGTGGTGGGAGCGGTGGCGGCGGATCC, the heavy- and light-chain cDNAs were fused and cloned into the pcDNA3.1 expression plasmid (Invitrogen). A fragment containing the myc and histidine tags of plasmid VP22/myc-his (Invitrogen) was subsequently cloned downstream of the light chain. The OVA cDNA was amplified by PCR from plasmid pcDNA–OVA-L (63) and cloned in frame between the light-chain and the tag, resulting in scDEC-OVA (Figure 1A). To generate scControl-OVA, heavy- and light-chain variable region cDNAs of the III/10 control antibody (which is the same isotype as the NLDC145 antibody but does not bind to DCs) (24) were also fused by overlap extension PCR and inserted in scDEC-OVA, replacing the respective DEC variable regions. Constructs were confirmed by sequence analysis. Similarly, HIV gag p24 and p41 were amplified from the plasmid synag (64) and inserted in place of the OVA gene in the scFv vectors. For in vivo experiments, plasmid DNA was prepared using the QIAGEN Endo-Free plasmid kit.

Cell lines, media, and antibodies. 293 cells, 293T cells (293ts/A1609), CHO cells, and CHOMDEC205 cells were maintained in DMEM supplemented with 10% FCS, penicillin, streptomycin, and glutamine. CV-1 cells were

The Journal of Clinical Investigation
http://www.jci.org Volume 118 Number 4 April 2008
maintained in the same medium but with Genetin in place of streptomycin. Antibodies specific for the TCRs of the transgenic OT-I and OT-II cells (Vβ1/3,2/MR9-4, Vβ2/20.1) and to other cell surface markers (CD11c, CD8α, and CD4) were purchased from BD Biosciences — Pharmingen. Magnetic microbeads were from Miltenyi Biotec. Other antibodies included rabbit anti-OVA (Chemicon), FITC-labeled anti-p24 (KC75-FITC; Coulter) and HRP-conjugated monoclonal antibody to HIV gag p24 (ImmunoDiagnostics), and FITC-labeled goat anti-rabbit and goat anti-mouse polyclonal antibodies (BioSource).

SDS-PAGE and Western blot analysis. To assess production of scDEC and control antibody fusion protein, both the cell lysate and crude supernatants of the 293 transfectants were resolved on 10% SDS-PAGE under reducing conditions and blotted onto nitrocellulose (Amersham) by standard techniques. The blots were blocked with 10% fat-free milk and then incubated with rabbit anti-OVA polyclonal antibody (1:10,000; Chemicon) followed by an anti-rabbit peroxidase conjugate (1:5,000). Bands were visualized with ECL substrate (Amersham). For the detection of HIV gag p24 and p41, the cell lysates and crude supernatants were similarly resolved on a 10% SDS-PAGE and blotted with 1:7500 HRP-conjugated anti-p24 mAb.

DNA vaccination. BALB/c, Cx66 F1 mice were obtained either from Charles River, Taconic, or Harlan and maintained under specific-pathogen-free conditions and used at 6 to 10 weeks under the guidelines of the Rockefeller University Institutional Animal Care and Use Committee. All experiments were performed in compliance with the institutional guidelines and approved by the Rockefeller University Institutional Animal Care and Use Committee. Mice were injected with DNA i.m. in combination with electroporation in the anterior tibialis muscles of both hind legs. In brief, the mice were anesthetized with 1.25 mg/mouse Nembutal. The skin overlying the anterior tibialis muscles of the hind legs was shaved and sterilized with ethanol swabs. DNA vaccine dissolved in 0.95% NaCl was injected i.m. and electroporation was achieved using a 2-nee gene pulse. Alternatively, DNA delivery and electroporation were achieved using an Ichor device (Ichor Medical Systems).

Cell transfer. CD8+ or CD4+ T cells were prepared from spleen and lymph nodes of OT-I or OT-II mice, respectively, by negative selection using hybridoma supernatants directed against MHC-II, F4/80, B220, NK1.1, and CD4 or CD8 followed by goat anti-rat Dynabeads (Dynal). Purified OT-I and OT-II that had been vaccinated 4–5 days previously either with scDEC–OVA or sterilized with ethanol swabs. DNA vaccine dissolved in 0.95% NaCl was injected i.m. followed by goat anti-rat Dynabeads (Dynal). The presence of the T cells was confirmed by immunofluorescence staining with antibodies to CD8 or CD4 followed by goat anti-rat Dynabeads (Dynal). Purified OT-I and OT-II cells, 107 cells/ml, were incubated with 5 μM CFSE (Molecular Probes) for 10 minutes at 37°C. An equal volume of FCS was added, and the cells were washed 2 times with PBS/0.1% BSA and twice with PBS. 2 × 106 labeled OT-I and 3 × 106 labeled OT-II cells were injected i.v. into B6 recipients that had been vaccinated 4–5 days previously either with scDEC–OVA or scControl–OVA DNA vaccines. After 3 days, cells suspensions of draining lymph nodes and spleen were stained with Vβ1/3/2 and CD8 (OT-I) or Vβ2 and CD4 (OT-II) using mAbs from BD Biosciences. The CFSE fluorescence intensity of OT-I and OT-II cells was then evaluated by multicolor flow cytometry using a FACScalibur (BD Biosciences) with subsequent analysis of data in CellQuest (BD Biosciences) or FlowJo (Tree Star Inc.).

Analysis of DC targeting. C57BL/6 (B6), TAP−/−, DEC−/−, OT-II, and OT-I mice were obtained from Charles River and maintained under standard conditions in the Rockefeller University animal facility. Four to five days after the injection of DNA vaccine, DCs were isolated from lymph nodes (popliteal, inguinal, axillary, brachial, mesenteric) and spleen. In brief, single-cell suspensions were prepared with 400 U/ml collagenase D (Roche) for 25 minutes at 37°C. The cells were incubated with microbeads that had been coated with anti-mouse CD11c antibodies for 30 minutes at 4°C. CD11c+ (DC-enriched) and CD11c− cells were separated by application of a magnetic field. The isolated DCs were added in graded doses to the CFSE-labeled OT-I cells (0.2 μM CFSE/106 splenocytes for 10 minutes at 37°C; 350,000 cells per well) at a DC:T cell ratio of 1:1, 1:3, and 1:9 in a final volume of 200 μl RPMI 1640 culture medium containing 5% FBS, 100 U/ml penicillin streptomycin mixture, 0.25 mg/ml Fungizone, 10 mM HEPES, and 55 μM β-mercaptoethanol. Total cell numbers in each well were normalized by adding an appropriate number of DCs from naive mice. Three and one-half days after, OT-1 T cells were evaluated for proliferation by CFSE dilution. Similarly CD11c+ mice were stained with immunofluorescent antibodies, after which the cell suspensions were were added to the CFSE-labeled T cells. Three days later, draining lymph node lymphocytes were evaluated for proliferation of the T cells.

Serum antibody titers. For the detection of HIV gag p41–specific antibodies, high protein–binding ELISA plates were coated overnight at 4°C with 5 μg/ml of gag p41 protein in PBS. Plates were washed 3 times with PBS–Tween 20 (0.02%) and blocked with PBS–BSA 1% for 2 hours at room temperature. Serum antibody titers were then determined as described (53), with the titer being the last dilution showing an OD405 greater than 0.1.

T cell responses to scDNA vaccines. To determine the breadth of HIV gag-specific T cell responses, bulk splenocytes were restimulated in vitro with peptide pools spanning the entire gag p41 protein (26). For the HIV gag-specific IFN-γ/IL-2 ELISPOT assays, ELISPOT plates (MAIPS, Millipore) were coated overnight either with anti-mouse IFN-γ or IL-2 capture antibodies (BD Biosciences — Pharmingen). Bulk splenocytes were then cultured for 18 hours at 37°C with 5% CO2 in the presence of 2 μM of HIV gag p41 peptide pools as previously described (26). Plates were developed with anti–IFN-γ biotinylated antibody (BD Biosciences) and spots visualized with avidin–horseradish peroxidase (Vector Laboratories), followed by DAB as substrate (Invitrogen). Spots were counted in an ELISPOT reader (Autoimmun Diagnostika GmbH). For multiparametric cytokine assays, bulk splenocytes were restimulated either with an H-2Kb–restricted peptide (2 μg/ml; AMQMLKETI, p24 197–205), the entire HIV gag p41 peptide pool mix, a negative control influenza HA peptide mix, or medium alone in the presence of 2 μg/ml of anti-CD28 (clone 37.51) for 6 hours, adding 10 μg/ml of Brefeldin A (Sigma-Aldrich) for the last 4 hours to accumulate intracellular cytokines. Cells were stained as described (26), but dead cells were excluded using live/dead fixable dead cell stain kit (Aqua LIVE/DEAD, Invitrogen). After blocking Fcy receptors, the cells were stained with antibodies CD3-Pacific Blue, CD8-FITC, CD4-percp, CD44–Alexa Fluor 750 (eBioscience) for 15 minutes at room temperature. We used a BD LSRII with data analysis in FlowJo (Tree Star Inc.).

Vaccinia gag protection assays. Nematobul-anesthetized mice were challenged with 2.5 × 106 (B6 mice not shown), 5 × 105 (BALB/C), and 1 × 106 (C57/B6 F1) PFUs of infectious virus by the intranasal route, in 35 μl PBS containing 10 μg/ml of Brefeldin A (Sigma-Aldrich) for the last 4 hours to accumulate intracellular cytokines. Cells were stained as described (26), but dead cells were excluded using live/dead fixable dead cell stain kit (Aqua LIVE/DEAD, Invitrogen). After blocking Fcy receptors, the cells were stained with antibodies CD3-Pacific Blue, CD8-FITC, CD4-percp, CD44–Alexa Fluor 750 (CD27–PE), and Aqua LIVE/DEAD stain for 20 minutes at 37°C. Cells were washed, fixed (Cryofix/Cytoperm Plus, BD Biosciences), permeabized with PermWash and stained with antibodies to IFN-γ (Alexa Fluor 700), TNF-α (Cy7–PE), and IL-2 (APCs) (eBioscience) for 15 minutes at room temperature. We used a BD LSRII with data analysis in FlowJo (Tree Star Inc.).

Statistics. Postchallenge mean vaccinia lung virus titers and mean percentage change in weight were compared between vaccination groups using one-tailed Student’s t test. Differences were considered significant at P < 0.05. Statistical analysis was performed using Prism 3 (GraphPad Software).

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