The catalytic A1 domains of cholera toxin and heat-labile enterotoxin are potent DNA adjuvants that evoke mixed Th1/Th17 cellular immune responses

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DNA encoded adjuvants are well known for increasing the magnitude of cellular and/or humoral immune responses directed against vaccine antigens. DNA adjuvants can also tune immune responses directed against vaccine antigens to better protect against infection of the target organism. Two potent DNA adjuvants that have unique abilities to tune immune responses are the catalytic A1 domains of Cholera Toxin (CTA1) and Heat-Labile Enterotoxin (LTA1). Here, we have characterized the adjuvant activities of CTA1 and LTA1 using HIV and SIV genes as model antigens. Both of these adjuvants enhanced the magnitude of antigen-specific cellular immune responses on par with those induced by the well-characterized cytokine adjuvants IL-12 and GM-CSF. CTA1 and LTA1 preferentially enhanced cellular responses to the intracellular antigen SIVmac239-gag over those for the secreted HIVBal-gp120 antigen. IL-12, GM-CSF and electroporation did the opposite suggesting differences in the mechanisms of actions of these diverse adjuvants. Combinations of CTA1 or LTA1 with IL-12 or GM-CSF generated additive and better balanced cellular responses to both of these antigens. Consistent with observations made with the holotoxin and the CTA1-DD adjuvant, CTA1 and LTA1 evoked mixed Th1/Th17 cellular immune responses. Together, these results show that CTA1 and LTA1 are potent DNA vaccine adjuvants that favor the intracellular antigen gag over the secreted antigen gp120 and evoke mixed Th1/Th17 responses against both of these antigens. The results also indicate that achieving a balanced immune response to multiple intracellular and extracellular antigens delivered via DNA vaccination may require combining adjuvants that have different and complementary mechanisms of action.

Introduction

For various known and unknown reasons, the promise of DNA vaccination has translated poorly from mice to primates. While all of the reasons for the loss of potency between mice and primates have not been fully elucidated, 2 reasons are believed to be most important - poor DNA uptake/expression and reduced immune-stimulatory signals at the inoculation site.1-5 Ballistic particle bombardment with the gene gun6 and in vivo electroporation7 have dramatically enhanced DNA uptake generating substantially improved immunogenicity in primates.2,5,5-21 However, delivery of DNA with either of these techniques is not ideal for large scale vaccination programs, particularly in underdeveloped countries. DNA expressed adjuvants, such as cytokines22-26 or chemokines,27-29 have also been used to boost immunogenicity. Unfortunately, like DNA vaccination itself, many of these adjuvants did not translate well from mice to primates.30-32 A notable exception is the Th1 polarizing cytokine, IL-12, that is perhaps the best DNA adjuvant tested to date in non-human primates and humans.33-39 Unfortunately, IL-12 is not sufficiently potent to alleviate the need for delivery with electroporation.

Cholera Toxin (CT), heat-labile enterotoxin (LT) and their enzymatically active A1 domains have also shown great promise as DNA vaccine adjuvants.6,19,21,40,41 CT and LT are closely related AB5 enterotoxins produced by Vibrio cholera and E. coli, respectively and are well known mucosal immunogens and adjuvants (reviewed in ref.42). These toxins consist of catalytic A domains (separated into A1 and A2 subunits) anchored in rings of 5 identical B pentamers.43 Their enzymatic active sites reside within their A1 subunits, while the A2 subunits anchor the A1 subunits into the B pentamers.43 The B pentamers engage
gangliosides on cell membranes and facilitate the entry of the A subunits into lysosomes. The A1 subunits then exploit host protein retention and degradation pathways to gain access to the cell cytoplasm (reviewed in ref.45).

In the cytoplasm, the A1 subunits of CT and LT (CTA1 and LTA1, respectively) catalyze the transfer of ADP-ribose moieties from nicotinamide adenine dinucleotide (NAD) to stimulatory α-subunits of G proteins (Gαs). The ADP-ribosylation of Gαs locks it into the GTP bound active state and subsequently causes the constitutive activation of adenylate cyclase leading to sustained high levels of cAMP within cells. The ADP-ribosyltransferase activities of CTA1 and LTA1 are responsible for their toxicity and for a large portion of their adjuvant effects.41,47-53 Although Gαs is a well-established cellular target of both CTA1 and LTA1, other signaling proteins are known to be ADP-ribosylated by these toxins.54,55 Whether Gαs is the principle target responsible for adjuvant activity is unknown.

Unfortunately, the CT and LT holotoxins cannot be used as mucosal adjuvants in humans since they are extremely toxic when administered at mucosal sites even at very small doses.56 Several methods have been used to minimize toxicity including toxin-inactivating mutations,51,57-59 immunizing at sites distant from the mucosa, such as the skin,60-64 or by re-targeting the A1 subunits using plasmid DNA40,41 or other proteins47,48 absent the promiscuous native B subunit. CTA1, expressed as the native bacterial DNA sequence as a DNA adjuvant, enhanced both cellular and antibody responses to HIVgalgp120 when delivered intramuscularly (i.m.).41 Human codon-optimization enhanced the in vitro biological activity of CTA1 by 250-fold and enhanced antibody responses to SIV gag when administered by gene gun.40 Cellular responses, however, were absent when delivered by the gene gun.40

Much of the promise of DNA based adjuvants lies in their abilities to “tune” immune responses in directions better suited to protect vaccinees from target infectious agents. Enabling this outcome requires more in depth profiling of the immune responses induced by the selected adjuvants. Since CTA1 and LTA1 have proven to evoke potent antibody responses when administered via gene gun, we wanted to profile the antibody and T cell responses provided by these adjuvants when administered i.m. We also compared the immune profiles of CTA1 and LTA1 against IL-12 and GM-CSF and show that their adjuvant effects are on par with those of IL-12 and GM-CSF. More importantly, these enterotoxin subunit adjuvants had complementary effects when combined with either of these cytokine adjuvants.

In contrast to the cytokine adjuvants IL-12 and GM-CSF that preferentially enhanced cellular responses to the secreted antigen gp120 over the intracellular antigen gag, CTA1 and LTA1 preferentially enhanced cellular responses to gag over gp120. LTA1 and CTA1 also evoked a mixed Th1/Th17 cellular response consistent with what has been observed with the holotoxin adjuvants and the CTA1-DD adjuvant.65-69 Together, the results of this study suggest that an optimal combination of DNA expressed adjuvants that enhance the magnitude of immune responses to vaccine antigens and tune the immune responses toward those best suited to combating targeted infectious agents may eliminate the need for suboptimal delivery techniques such as gene gun and electroporation for some vaccines, particularly when they are used as a prime for a recombinant subunit boost.

**Results**

One inoculation evokes near maximal magnitude cellular responses when CTA1 is used as an adjuvant

We first determined how many inoculations are necessary to evoke high-level cellular and/or antibody responses when a codon-optimized CTA1 is used as a DNA vaccine adjuvant. For this experiment, mice were immunized 1, 2 or 3 times with or without the CTA1 adjuvant on days 0, 14 and/or 28. Mice were inoculated with pHIV-gp120 and pSIV-gag plus either empty plasmid (as a no adjuvant control) or with pCTA1. Splenocytes from all mice were harvested on day 48 and stimulated with peptide pools encompassing HIVgalgp120 or SIVmac239gag for IFN-γ ELISpot analysis. Day 48 correlates to 20 d post the final immunization for mice vaccinated 2 or 3 times and 20 d post the single immunization for mice vaccinated a single time.

As expected, the number of HIV gp120 and SIV gag IFN-γ ELISpots increased after each vaccination in those animals that did not receive an adjuvant (Fig. 1A). By contrast, CTA1 adjuvanted mice reached high-magnitude cellular responses after a single inoculation and these cellular responses did not increase further with additional inoculations. SIV gag and HIV gp120 ELISpots were 7.3-fold and 5.1-fold higher, respectively than those of control mice that did not receive an adjuvant after a single inoculation (Fig. 1A). Importantly, the number of gag-specific ELISpots from mice inoculated a single time with CTA1 was significantly higher (3.8-fold, p = 0.001) than the number of gag-specific ELISpots from mice inoculated 3 times in the absence of adjuvant (Fig. 1A). Similarly, the number of gp120-specific ELISpots from mice inoculated a single time with CTA1 was higher than the number of gp120-specific ELISpots from mice inoculated 3 times in the absence of adjuvant (Fig. 1A); however, the difference between these responses did not reach statistical significance.

In contrast to the cellular responses, the codon optimized CTA1 did not enhance antibody responses to HIV gp120 (Fig. 1B). The antibody titers to gp120 increased with subsequent immunizations whether CTA1 was included or not, however, mice that received CTA1 had gp120 titers that were significantly lower (8-fold) than those of unadjuvanted controls after 3 inoculations. The gp120 titers for most mice in this study did not rise above background levels (data not shown). Subsequent experiments focus comparison on the cellular responses as little to no differences were found between the antibody responses when we used CTA1 or LTA1 delivered i.m.

Dose response of CTA1

To determine the optimal dose range of the codon optimized CTA1, mice were immunized i.m. once on day 0 with
pHIV-gp120 and pSIV-gag plus either an empty plasmid, pIRES-IL-12 or different doses of pCTA1. Enough empty plasmid was included to keep the total DNA delivered in each vaccination to 70 μg. An additional group that received unadjuvanted DNA delivered i.m. with electroporation was included as a comparator since delivery by electroporation has been shown to dramatically increase antigen expression and ensuing immune responses.7,13-16,70-75 Splenocytes were harvested on day 14 and stimulated with HIV-BaLgp120 or SIVmac239gag peptide pools for IFN-γ ELISpot assays.

Figure 2 shows that CTA1 significantly enhanced the anti-gag ELISpot responses over the unadjuvanted controls at all doses tested. Increases in the anti-gp120 ELISpot responses were also observed but did not reach statistical significance. These observations are similar to those from the previous study and demonstrate that CTA1 preferentially enhances cellular responses to the intracellular expressed antigen gag over those to the secreted antigen gp120. IL-12 did the opposite by significantly increasing the ELISpot responses to gp120 but not significantly enhancing the responses to gag (Fig. 2). Strikingly, CTA1 induced anti-gag ELISpot responses that were equal or greater in magnitude than those evoked by electroporation at the 25 and 75 μg doses (Fig. 2). We can conclude that CTA1 has a very wide effective dose range (over 3 orders of magnitude) and that it has the unusual ability to preferentially enhance responses to the intracellular antigen gag over the secreted antigen gp120.

CTA1 has complementary adjuvant effects with IL-12 and GM-CSF

IL-12 is a Th1 polarizing cytokine with direct effects on T cells76-81 and is one of the most potent DNA vaccine adjuvants ever tested in mice and primates.11,33-39 GM-CSF is another well-studied and potent DNA vaccine adjuvant.82-85 Since CTA1 likely has a different mechanism of DNA adjuvant activity than IL-12 or GM-CSF, we speculated that their adjuvant effects would be complementary (i.e., additive or synergistic). To test these hypotheses, mice were immunized i.m. 1 time on day 0 with pHIV-gp120 and pSIV-gag plus either empty plasmid, pCTA1, pIRES-IL-12, pGM-CSF or combinations of pCTA1 and pIL-12 or pGM-CSF.
**Figure 3.** CTA1 has complementary adjuvant effects with IL-12 and GM-CSF: Groups of 6 BALB/c mice were immunized i.m. 1 time on day 0 with 30 μg of pSV-gag and 15 μg of pHIV-gp120 plus either 40 μg of empty plasmid, 40 μg of pCTA1, 40 μg of pIL-12, 40 μg of pGM-CSF or 20 μg of pCTA1 + 20 μg of pIL-12 or 20 μg of pCTA1 + 20 μg of pGM-CSF. Day 14 splenocytes were stimulated with SIVmac239gag or HIV-BaLgp120 peptide pools for IFN-γ ELISPOT assays. Consistent with the previous experiments using CTA1, LTA1 preferentially enhanced responses to gag (8.8-fold) over gp120 (3.6-fold) (Fig. 4A). Again, IL-12 did the opposite by preferentially improving responses to gp120 (2.7-fold) over gag (1-fold) (Fig. 4A). Similar to when CTA1 was combined with IL-12, the LTA1 + IL-12 group had total gp120 + gag ELISPot responses (2432) approximately additive when compared to the LTA1 alone (1377 total spots) and the IL-12 alone (502 total spots) groups. The differences in the total gp120 + gag ELISPot responses between the LTA1 + IL-12 group and the LTA1 or IL-12 alone group were also statistically significant (p = 0.032 for the LTA1 alone group compared to the LTA1 + IL-12 group and p = 0.001 for the IL-12 alone group compared to the LTA1 + IL-12 group).

Similar to the ELISPot results, LTA1 preferentially enhanced IFN-γ secretion in response to gag peptides (23.7-fold) over gp120 peptides (20-fold) whereas IL-12 did the opposite by preferentially enhancing IFN-γ secretion in response to gp120 (7.2-fold) over gp120 (1.2-fold) (Fig. 4B). Strikingly, the animals that received LTA1 secreted significantly more IFN-γ when calculated on a per cell basis (3.26 pg/spot) when compared to both the unadjuvanted group (0.82 pg/spot, p = 0.016), and the IL-12 adjuvanted group (1.53 pg/spot, p = 0.008). The LTA1 + IL-12 group also had strong IFN-γ secretion in response to both gp120 and gag (Fig. 4B) with an average secretion of 2.56 pg/spot.

Granzyme B is a Cytotoxic T cell (CTL) effector molecule that directly participates in the killing of target cells. This release of Granzyme B by peptide stimulated splenocytes is often used as a measure of the killing capacity of the evoked cellular immune responses. Figure 4C shows that LTA1 significantly enhanced Granzyme B secretion from gag peptide stimulated splenocytes 3.8-fold over the unadjuvanted group, but did not enhance Granzyme B secretion from gp120 peptide stimulated splenocytes affirming the preferential boosting of gag specific cellular immune responses by the A1 subunit adjuvants. IL-12, on the other hand, did not boost Granzyme B secretion for either antigen. The LTA1 + IL-12 combination yielded enhanced combined total Granzyme B secretion (gp120 + gag) from

better balanced cellular responses to both antigens as compared to the adjuvants used alone (Fig. 3).

**LTA1 and CTA1 have similar adjuvant effects**

CT and LT are highly homologous enterotoxins with the majority of their differences lying in the targeting domains of their respective B subunits. Subtle differences have been intermittently observed between immune responses when the holotoxins are used either as antigens or adjuvants. Since there are only 26 amino acid differences between the 196 amino acid A1 domains of our CTA1 and LTA1 DNA adjuvants (87% homology), we wanted to confirm that they would have similar DNA vaccine adjuvant effects and that LTA1 would also have additive effects with IL-12.

In this experiment, mice were immunized i.m. on days 0 and 21 with pHIV-gp120 and pSIV-gag plus either an empty plasmid, pIL-12, pLTA1, or a combination pLTA1 and pIL-12. Day 35 splenocytes were stimulated with HIV-BaLgp120 or SIVmac239 gag peptide pools for IFN-γ ELISPot assays. The error bars represent the standard errors of the means. The P values (compared to the no adjuvant control) were calculated with a Students T Test using SigmaPlot v12 software. NS stands for not significantly different from the unadjuvanted control. The results shown are from a single experiment of 3 performed.
spleenocytes stimulated by peptides that was balanced between both antigens and was significantly higher than the unadjuvanted group (p = 0.008), the IL-12 group (p = 0.008) and the LTA1 group (p = 0.032) (Fig. 4C). Together, these data imply that the main benefit of the CTA1/LTA1 + IL-12 or CTA1 + GM-CSF combinations is that these combinations significantly enhance the combined gp120 + gag responses over the responses evoked by the individual adjuvants. This represents mostly an additive effect rather than a synergistic effect. In other words, when CTA1 or LTA1 (that preferentially enhance responses to gag) are combined with IL-12 or GM-CSF (that preferentially enhance responses to gp120) the result is an enhancement to both antigens and the total response (the additive total for gag and gp120) is significantly enhanced compared to the individual adjuvants alone.

LTA1 evokes mixed Th1/Th17 antigen-specific T cell responses

The dramatically enhanced secretion of IFN-γ induced by the LTA1 adjuvant prompted us to profile the cytokines and chemokines induced by the adjuvant more thoroughly. Mice were immunized i.m. on days 0 and 21 with 25 μg of pSIV-gag and 15 μg of pHIV-gp120 plus either 30 μg of empty plasmid, 30 μg of pLTA1, 30 μg of pLTA1-15 or 15 μg of pIL-12. Day 35 splenocytes were stimulated with SIVmac239 gp120 or HIV-Bal gp120 peptide pools for IFN-γ ELISpot assay and for supernatant cytokine analysis. (A) Splenocytes assayed by IFN-γ ELISpot assay. (B) Supernatants from peptide-stimulated splenocytes plated at 8 × 10^5 cells/well analyzed by IFN-γ ELISA. (C) Supernatants from peptide-stimulated splenocytes plated at 8 × 10^5 cells/well analyzed by Granzyme B ELISA. The error bars represent the standard errors of the means. The P values in panel (A) (compared to the no adjuvant group) were calculated with a Students T Test using SigmaPlot v12 software. The P values in panels (B and C) (compared to the no adjuvant group) were calculated with a Mann-Whitney Rank Sum-test using SigmaPlot v12 software. NS stands for not significantly different from the unadjuvanted control. The results shown are from a single experiment of 2 performed.

As expected, the LTA1 and IL-12 adjuvants evoked IFN-γ ELISpot responses that were consistent with the previous study (Fig. 5A). Immunization with IL-12 enhanced the gp120 IFN-γ ELISpot response 3.8-fold but did not enhance the gag response as compared to the unadjuvanted controls (Fig. 5A). LTA1 enhanced gag IFN-γ ELISpot 5.1-fold and anti-gp120 ELISpot 3.5-fold compared to the no adjuvant controls (Fig. 5A). Peptide-stimulated splenocytes from LTA1 adjuvanted mice again secreted more IFN-γ per ELISpot than those from unadjuvanted controls (Fig. 5B) (0.19 pg/spot for LTA1 mice verses 0.084 pg/spot for unadjuvanted mice). Examining the production of other cytokines revealed that LTA1 induced the secretion of more IL-2, IL-3, IL-6, RANTES, GM-CSF, TNF-α and MIP-1β compared to
unadjuvanted or IL-12 adjuvanted mice (Fig. 5B). The concentrations of these cytokines and chemokines from LTA1 adjuvanted mice were also higher for gag than for gp120 further highlighting the preferential enhancement of gag responses over gp120 responses by this adjuvant. The concentrations of the remaining 24 cytokines and chemokines were either undetectable or were not different between groups (data not shown). LTA1 also dramatically enhanced the secretion of IL-17 (both the IL-17A and IL-17F isoforms) from gp120 and gag peptide stimulated splenocytes indicating that LTA1 evokes a mixed Th1/Th17 response (Fig. 5C). Although LTA1 evoked IL-17 secretion from both gag and gp120 peptide stimulated splenocytes, again the IL-17 concentrations were no statistically significant differences in the fold-enhancements in the gp120 ($p = 0.96$) or gag ($p = 0.72$) responses evoked by CTA1 and LTA1. The average fold-enhancements for gag were also 2-times higher for CTA1 and 1.7-times higher for LTA1 than the corresponding fold-enhancements for gp120 (Fig. 6). These data further demonstrate the preferential enhancing effects of CTA1 and LTA1 for gag over gp120. Figure 6 also shows that IL-12 did the opposite by preferentially enhancing IFN-γ ELISpot responses to gp120 over those to gag. In this regard, the average fold-enhancements for gp120 were 2.1-times higher for IL-12 than the corresponding fold-enhancements for gag.

CTA1 and LTA1 preferentially enhance cellular responses to gag over gp120

The studies above indicate that CTA1 and LTA1 behave similarly as DNA adjuvants and that both adjuvants preferentially enhance cellular responses to gag over gp120. We next determined if the preferential enhancement of cellular responses to gag over gp120 by these adjuvants was retained between different studies performed at different times using different reagents. A total of 9 individual mouse studies were performed using CTA1, 6 were performed using LTA1 and 9 were performed using IL-12. These studies were performed over a period of 4 years and often used different batches of DNA and/or peptides.

Figure 6 shows that CTA1 and LTA1 evoked similar average fold enhancements in the gp120 and gag specific IFN-γ ELISpot responses between these studies. In fact, there were no statistically significant differences in the fold-enhancements in the gp120 ($p = 0.96$) or gag ($p = 0.72$) responses evoked by CTA1 and LTA1. The average fold-enhancements for gag were also 2-times higher for CTA1 and 1.7-times higher for LTA1 than the corresponding fold-enhancements for gp120 (Fig. 6). These data further demonstrate the preferential enhancing effects of CTA1 and LTA1 for gag over gp120. Figure 6 also shows that IL-12 did the opposite by preferentially enhancing IFN-γ ELISpot responses to gp120 over those to gag. In this regard, the average fold-enhancements for gp120 were 2.1-times higher for IL-12 than the corresponding fold-enhancements for gag.
This optimized CTA1 dramatically increased cellular responses to SIVmac239gag, and to a lesser extent to HIV_Bal gp120 (Figs. 1–3), but did not improve the magnitude of the anti-gp120 antibody response (Fig. 1B). Similar results were observed with an optimized LTA1 (Figs. 4 and 5). This apparent discrepancy from our previous report where we used a non-optimized CTA1 is most likely an artifact of the experimental design. Here, the mice were sacrificed 20 d post the final immunization which may have been too short for antibody responses to fully develop since it took 26 weeks for the antibody response to reach its maximum after immunization with the native CTA1 sequence. 41 Also, when delivered by gene gun, CTA1 dramatically enhanced antibody responses to SIV-gag (over 2 orders of magnitude) but had little to no enhancing effects on antibody responses to HIV-gp120.40

The CTA1 and LTA1 adjuvants are closely related enzymes with likely identical activities and targets. 74 Our data indicate that these enzymes also behave indistinguishably from each other as DNA adjuvants. For this reason, we believe that CTA1 and LTA1 can be thought of interchangeably as DNA adjuvants. We have chosen LTA1 as a lead only because there is a general perception that CTA1 is more toxic than LTA1. The preferential enhancing effects that CTA1 and LTA1 have for the intracellular antigen gag over the secreted antigen gp120 may be due to their ability to facilitate the release of intracellular antigens into the extracellular milieu. In this regard, we found that CTA1 and LTA1 increased the release of an intracellular retained reporter protein into the extracellular milieu by approximately 2 orders of magnitude (manuscript in preparation). This may facilitate the transfer of intracellular expressed antigens from transfected muscle cells to APCs. This ability of CTA1 and LTA1 may have also caused or contributed to the reduced antibody responses to gp120 observed in Figure 1. In this regard, it is plausible that increased antibody responses to gag may have decreased the antibody responses to gp120 (i.e., antigen competition). We are currently studying the mechanism(s) of the adjuvant effects of CTA1 and LTA1 in greater detail. The results of these studies should shed more light on the mechanism(s) of action of these adjuvants and may better answer the questions of why CTA1 and LTA1 preferentially enhance responses to gag over gp120 and why CTA1 reduced antibody responses to gp120.

Interestingly, the results of this study, where CTA1 boosted cellular responses but not antibody responses are the converse of those of our previous study where CTA1 was delivered by gene gun. 40 In that study, CTA1 boosted antibody responses but not cellular responses. However, in that study CTA1 dramatically enhanced antibody responses to gag, but did not significantly enhance antibody responses to gp120.40 This is consistent with the preferential enhancing effects that CTA1 and LTA1 had for gag over gp120 in the current studies. The results of that study may also be explainable by the ability of CTA1 and LTA1 to cause the release of normally intracellular retained proteins into the extracellular milieu.

We do not believe that our failure to detect cellular responses when the adjuvants were delivered by gene gun represents a failure of the adjuvants, but rather a failure of the DNA delivery. In

![Figure 6. CTA1 and LTA1 preferentially enhance cellular responses to gag over gp120. The results from 9 independent mouse studies for CTA1, 6 independent studies for LTA1 and 9 independent studies for IL-12 were compared. Each study had groups of 5 BALB/c mice immunized i.m. on days 0 and 14 or days 0 and 21 with 25 μg of pHIV-gp120 and 15 μg of pHIV-gp120 plus either 25 μg of empty plasmid, pSIV-gag or HIV_Balgp120 peptide pools for IFN-γ ELISpot assays. For each study, the average gp120 and gag-specific IFN-γ ELISpot responses were determined for the no adjuvant groups and the CTA1, LTA1 or IL-12 adjuvanted groups. Within each study, the average fold increases for the gp120 (light gray bars) and gag (dark gray bars)-specific ELISpot responses were calculated by dividing the mean gp120 and gag-specific ELISpot responses from the adjuvanted groups by the corresponding mean gp120 and gag-specific ELISpot responses from the unadjuvanted groups. The error bars represent the standard errors of the means. The P values (provided in the text) were calculated with a Students T test using SigmaPlot v12 software.](image-url)
this vein, others have consistently measured robust cellular responses after gene gun immunization, even in the absence of adjuvants.6,19,94-101 We, however, failed to measure cellular responses with gene gun delivery even when IL-12 was used as the adjuvant (data not shown). Our inability to evoke cellular responses with the gene gun likely stems from undetermined differences in formulation and/or delivery parameters. Differences in the local tissue environments between the epidermis (where DNA is delivered by gene gun) and muscle (where DNA is delivered by i.m. inoculation) also likely impacts how well the vaccines evoke humoral immune responses. For instance the epidermis is rich in Langerhans cells which may be efficient at priming humoral responses, whereas muscle tissue has a much smaller population of antigen presenting cells (APCs).

The results of the current study also indicate that a single immunization with CTA1 provides maximal, or near maximal, cellular immune responses that are not improved by further boosting (Fig. 1A). The antibody response, on the other hand, did benefit from additional boosts (Fig. 1B). The adjuvant effects of combinations of CTA1 with IL-12 or GM-CSF were approximately additive (Fig. 3) principally because the adjuvants showed an effective preference for either the gag or gp120 when used alone that combined to give more balanced responses when the adjuvants were used together. The propensity for CTA1 and LTA1 to preferentially enhance cellular responses to the intracellular SIVmac239 gag antigen (Figs. 1–6) may be a unique feature of these adjuvants since using no adjuvant, adjuvanting with IL-12 or GM-CSF, or delivery of DNA with electroporation all favored cellular responses to secreted HIV gag (Figs. 3–6). We are currently evaluating whether this observation can be generalized to other intracellular antigens. If so, then CTA1 and LTA1 may be ideal adjuvants for therapeutic vaccines for chronic viral infections and cancer where the majority of targeted antigens are intracellular or surface bound.

To date, no single adjuvant has been identified that can alleviate the need for delivery by electroporation. IL-12 has been promising in primates,33,38,39,98,102 including humans,39 but is still not sufficiently potent to negate the need for electroporation. Combining adjuvants may provide additive or synergistic effects that may eclipse those provided by the best individual adjuvants or delivery regimens. Our results with combinations of CTA1 or LTA1 with IL-12 or GM-CSF support this possibility (Figs. 3 and 4). With the right combination of adjuvants, it may be possible to eliminate the need for electroporation and potentially reduce the cost, and increase the deployability of some DNA vaccines, particularly those dependent solely on cellular responses. One caveat is that none of the adjuvants or combinations that we have tested have substantially boosted humoral responses when delivered i.m. without electroporation. For this reason, vaccines that require a humoral component for protection may require an enhanced delivery regimen such as electroporation or a heterologous boost with recombinant protein. We are currently exploring the effects of CTA1 and LTA1 delivered by electroporation.

Perhaps the most interesting finding of this study is that CTA1 and LTA1 evoked mixed Th1/Th17 cellular responses (Fig. 5 and data not shown). This may also be a unique feature of these bacterial toxin A1 subunit adjuvants since we have not detected the secretion of IL-17 from mice immunized with DNA alone or co-formulated with IL-12 or GM-CSF (Fig. 5 and data not shown). Th17 cells are a T helper CD4 subset103-105 that can secrete IL-17A104,105 and IL-17F,103 as well as IL-21106,107 and IL-22.108,109 Th17 cells generally orchestrate the eradication of extracellular bacteria and fungi by activating neutrophils through production of IL-17A and IL-17F (reviewed in ref.110). Like Th1 cells, Th17 cells also promote B-cell class switching to opsonizing IgG antibodies.111 Th17 cells have also been found to play roles in combating certain viruses and cancers and have been found to play important roles in several autoimmune disorders.112,113 Th17 cells can also acquire the ability to co-express IFN-γ.114 A number of recent studies have demonstrated that the CT and LT holotoxins evoke Th17 responses when delivered to mucosal sites.65-68 Additionally, the novel CTA1-DD adjuvant that retargets the A1 subunit of CT to B cells also stimulates mixed Th1/Th17 responses69. The results of that study69 along with our results here indicate that this Th17 skewing activity resides in the A1 subunits of these toxins, and occurs whether delivered mucosally or systemically. This suggests that CTA1 and LTA1 might be ideal adjuvants for anti-bacterial and anti-fungal vaccines (reviewed in ref.115). Interestingly, the CTA1-DD adjuvant targets B cells47,116,117 whereas i.m. delivered CTA1 or LTA1 almost exclusively targets muscle cells, yet both adjuvants evoke similar mixed Th1/Th17 responses. This indicates that the cell types targeted by CTA1-based adjuvants may be less important than the biological effects that these enzymes have on the targeted cells.

Numerous DNA-based adjuvants, including CTA1 and LTA1, have been discovered and studied, almost always individually. Given the cacophony of cytokines and chemokines that are induced upon infection, we believe that no single DNA adjuvant will solve the shortcomings of DNA vaccines. In this regard, during natural infections, the innate immune system senses multiple pathogen associated molecular patterns (PAMPs) and danger associated molecular patterns (DAMPs) released by host cells in response to infection. It is the integration of signals from these multiple PAMPs and DAMPs that fully triggers innate and adaptive immune responses. Single DNA adjuvants can at best only mimic one of the many signals triggered by natural infections. We therefore propose that combining adjuvants with different and potentially complimentary mechanisms of action will yield improved adaptive immune responses. The data generated under this study support this hypothesis and provide incentive to continue exploring new adjuvant combinations for unique and useful immunological outcomes. Additionally, our data show that different adjuvants can enhance immune responses to different antigens, suggesting that a single adjuvant may not be appropriate for every antigen.

Although the studies presented in this manuscript provide new insights into improving DNA vaccine performance, there are limitations to these studies. For instance, we focused on delivering our vaccines and adjuvants as naked DNA by i.m. injection, while most vaccines that are in clinical trials or are nearing clinical trials are delivered using electroporation. We have also
evaluated CTA1 and LTA1 and combinations of CTA1 and LTA1 with IL-12 and other adjuvants delivered by electroporation. We found similar enhancements in the cellular immune responses when these adjuvants were delivered using electroporation yielding dramatically higher cellular responses than those achieved by delivery with electroporation without adjuvants or by i.m. delivery with the same adjuvant combinations (manuscript in preparation).

Another limitation of these studies is that they do not elucidate the mechanism(s) of action of the CTA1 and LTA1 adjuvants. However, we have determined that their adjuvant effects are a result of their enzymatic activities and that their adjuvant effects cannot be mimicked by the elevation of cAMP levels within transfected muscle cells (manuscript in preparation). Therefore, Gs- and Gt-specific effects cannot be achieved by delivery with electroporation without adjuvants or by i.m. delivery with the same adjuvant combinations (manuscript in preparation). Achieving adjuvants that are worthy of further study, particularly in combination with other adjuvants.

Materials and Methods

DNA vaccine construction

The pMAX-PRO DNA vaccine vector was created by replacing the yellow fluorescent protein (YFP) gene from pMAX-FP-Yellow (Amaza Corp., Cologne, Germany) with a multiple cloning site using the Nhe-I and Bss-HII sites. The DNA sequences of CTA1 and LTA1 (minus their bacterial secretion signals), mouse GM-CSF, SIVmac259-gag and HIVBal-gp120 were codon-optimized for expression in human and mouse cells by GeneArt (Resensburg, Germany). The optimized sequences that were provided in the GeneArt shuttle vector were sub-cloned into pMAX-PRO using the 5' Kpn-1 and 3' Xho-1 sites. The plasmids pMAX-PRO-CTA1, pMAX-PRO-LTA1, pMAX-PRO-SIVmac259-gag, pMAX-PRO-HIVBal-gp120 and pMAX-PRO-mGM-CSF are referred to as pCTA1, pLTA1, pGM-CSF, pSIV-gag and pHIV-gp120 respectively in the text of this manuscript.

In early studies, we used an IL-12 expressing vector that expressed the mouse IL-12 p35 subunit from a CMV promoter and the p40 subunit from an Internal Ribosome Entry Site (IRES). This plasmid (pMAX-PRO-mIL-12) was created by excising the mIL-12 expression cassette from pUMVC3-mIL-12 (Aldeveron, Fargo, ND) and sub-cloning it into the pMAX-PRO backbone. The expression of mIL-12 from pMAX-PRO-mIL-12 was equivalent to that of pUMVC3-mIL-12 (data not shown). The pMAX-PRO-mIL-12 plasmid is referred to as pIL-12 in the text of this manuscript.

Vaccination procedures

DNA vaccinations given i.m. without electroporation were given in the quadriceps muscle(s). The mice were anesthetized using a Ketamine-based anesthetic before vaccination. Hair was removed from the areas over the quadriceps muscle(s) using clippers. The animals were inoculated in both quadriceps muscles using a 27 gauge needle and an insulin syringe. A total injection volume of 100 µl (50 µl/muscle) was injected. For mice receiving electroporation, the vaccines were also delivered into the quadriceps muscles. Within one minute after the inoculation, the muscles were subjected to electroporation with the following parameters: 6 pulses of 100 V lasting 50 ms each with 200 ms intervals between pulses. Electroporation was carried out using an 830 electroporation generator (BTX Harvard Apparatus, Holliston, MA) and a 2 needle gene probe with a 5 mm gap.

Animal housing and handling

For the studies in Figures 1 and 2, mice were housed in the Advanced BioScience Laboratories (ABL) Animal Facility in Rockville, MD. The facility is AAALAC-International Accredited, USDA Registered, and has a Category 1 Assurance from the Office of Laboratory Animal Welfare (Assurance # A3467-01). The Animal Research Facility complies with USDA regulations pertaining to animal care (USDA registration #51-R-0059) and with the “PHS Policy on Humane Care and Use of Laboratory Animals” The facility also complies with all policies of the “Guide for the Care and Use of Laboratory Animals” (National Academy of Sciences, 1996). ABL also performed the IFN-γ ELISPOT assays shown in Figures 1 and 2.

For the remaining studies, performed by Profectus, the mice were housed at the vivarium of the Department of Comparative Medicine of New York Medical College. This facility is AAALAC International accredited, USDA Registered and has an Assurance from the Office of Laboratory Animal Welfare (formerly OPRR) OLAW Assurance #A3362-1. The Comparative Medicine Department also complies with all the policies promulgated in the “Guide for the Care and Use of Laboratory Animals” (National Academy of Sciences, 1996).

Humoral responses to SIVmac259 gag

Solid-phase ELISAs were used to determine SIV-gag-specific antibody (Ab) titers in the sera. Briefly, 96 well high-binding microtiter plates (Nunc, Rochester, NY) were coated with 100 µl of 10 µg/ml of SIVmac259-gag (Immunodiagnostics, Woburn, Massachusetts) in PBS overnight at 4°C. Plates were washed 3 times with Tris Buffered Saline (TBS) and then blocked with 300 µl of Blotto (5% w/v non-fat dried milk in TBS) at room temperature for 30 minutes. Serially diluted sera were added to the wells and incubated at room temperature for one hour, then washed 3 times with TBS. Horseradish peroxidase-conjugated goat anti-mouse IgG (Kirkegaard & Perry, Gaithersburg, MD) diluted 1:1000 in Blotto was added (100 µl/well) and incubated at room temperature for one hour. The plates were washed 3 times with TBS before adding SureBlue TMB peroxidase substrate (Kirkegaard & Perry) (100 µl/well) and incubating for 3–5 minutes. The reactions were stopped by adding 50 µl/well of 1N H2SO4. Absorbance was read at 450 nm using a Beckman Coulter AD 200 Plate Reader (Brea, CA). Half-maximal serum binding titers were calculated using SigmaPlot v11 software.

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Humoral responses to HIV_BaLgp120

Capture ELISAs were used to detect HIV_BaL-gp120-specific Ab titers in the sera of mice. Briefly, 96 well high-binding microtiter plates (Nunc, Rochester, NY) were coated with 200 ng/ml of the sheep capture antibody D7324 (Aalto Scientific, Carlsbad, California) (18,19) (100 µl/well) overnight at 4°C. After three washes with TBS, 1 µg/ml of nLgp120 (purified recombinant) in PBS (100 µl/well) was added for one hour at 37°C. Plates were washed 3 times with TBS, then blocked with (300 µl/well) Blotto at room temperature for 30 minutes. Serially diluted sera were added to the wells and incubated at room temperature for one hour, then washed 3 times with TBS. Horseradish peroxidase-conjugated goat anti-mouse IgG (Kirkegaard & Perry, Gaithersburg, MD) diluted 1:1000 in Blotto was added after each wash and incubated at room temperature for one hour. The plates were washed 3 times with TBS before adding SureBlue TMB peroxidase substrate (100 µl/well) and incubating for 3–5 minutes. The reactions were stopped by adding (50 µl/well) of 1 N H2SO4. Absorbance was read at 450 nm using a Beckman Coulter AD 200 plate reader. Half-maximal serum binding titers were calculated using SigmaPlot v11 software.

IFN-γ ELISPot Assays

Ninety six-well flat-bottom ELISPot plates (Millipore, Bedford, MA) were coated overnight at 4°C with an anti-mouse IFN-γ monoclonal antibody (BD-PharMingen, San Jose, CA) at a concentration of 10 µg/ml. After which the plates were washed 3 times, then blocked for 2 hours with PBS containing 5% heat-inactivated FBS. Mouse splenocytes were resuspended in complete RPMI 1640 medium containing either medium alone, 50 µg/mL PHA-M (Sigma), or peptide pools (15 mers overlapping by 11 amino acids; 1 µM each final peptide concentration) spanning HIV_BaLgp120 or SIVmac239_gag, and plated in 96-well flat-bottom tissue culture plates (Becton Dickenson). Input cell numbers were 8 x 10^5 cells/well for the study shown in Figure 4 or 1 x 10^5 cells/well for the study shown in Figure 5. Cells were incubated for 24 hours at 37°C and then supernatants were removed for cytokine ELISAs or CBA. IFN-γ and Granzyme B Ready-SET-Go! ELISA kits were purchased from Affymetrix/eBioscience (San Diego, CA) and were performed according to the manufacturer’s instructions. CBA was performed by Affymetrix/ Panomics (Santa Clara, CA) using their Luminex-based platform (ProcartaPlex™ Multiplex). This service included sample handling, performance of the assays and data analysis. The concentrations of the following cytokines or chemokines were determined in the supernatants: IL-1α, IL1-β, IL-2, IL-3, IL-4, IL-5, IL-6, IL-9, IL-10, IL-12, IL13, IL-15, IL-17A, IL-17F, IL-21, IL-23, EOTAXIN, G-CSF, GM-CSF, IFN-γ, IP-10, KC, LIX, MCP-1, MCP-3, MIP-1α, MIP-1β, MIP-2, M-CSF, sRANKL, RANTES, TGF-β, TNF-α and VEGF. The concentrations of the cytokines and chemokines not shown in the figures were either below the limits of detection or were not significantly different between the groups.

Cytokine ELISAs and Cytokine Bead Array (CBA)

Splenocytes were resuspended in complete RPMI 1640 medium containing either medium alone, 50 µg/mL PHA-M (Sigma), or peptide pools (15 mers overlapping by 11 amino acids; 1 µM each final peptide concentration) spanning HIV_BaLgp120 or SIVmac239_gag, and plated in 96-well flat-bottom tissue culture plates (Becton Dickenson). Input cell numbers were 8 x 10^5 cells/well for the study shown in Figure 4 or 1 x 10^5 cells/well for the study shown in Figure 5. Cells were incubated for 24 hours at 37°C and then supernatants were removed for cytokine ELISAs or CBA. IFN-γ and Granzyme B Ready-SET-Go! ELISA kits were purchased from Affymetrix/eBioscience (San Diego, CA) and were performed according to the manufacturer’s instructions. CBA was performed by Affymetrix/ Panomics (Santa Clara, CA) using their Luminex-based platform (ProcartaPlex™ Multiplex). This service included sample handling, performance of the assays and data analysis. The concentrations of the following cytokines or chemokines were determined in the supernatants: IL-1α, IL1-β, IL-2, IL-3, IL-4, IL-5, IL-6, IL-9, IL-10, IL-12, IL13, IL-15, IL-17A, IL-17F, IL-21, IL-23, EOTAXIN, G-CSF, GM-CSF, IFN-γ, IP-10, KC, LIX, MCP-1, MCP-3, MIP-1α, MIP-1β, MIP-2, M-CSF, sRANKL, RANTES, TGF-β, TNF-α and VEGF. The concentrations of the cytokines and chemokines not shown in the figures were either below the limits of detection or were not significantly different between the groups.

Statistical methods

The error bars in the figures are the standard errors of the means (SEMs). The SEMs were calculated by dividing the standard deviations of the means (STDEVs) (determined using MicroSoft Excel software) by the square root of the number of animals in the group. To make comparisons between groups in a study, the medians of the immune measures for each group were calculated and then evaluated for normality to determine if the distribution required parametric or non-parametric statistical tests for analysis. When a parametric test was required, a student t-test comparison was used. When a non-parametric statistical test was required, a Wilcoxon rank-sum test (2-group comparison) was used. Tests were performed using SigmaPlot v11 or v12 software. A p value ≤0.05 indicates that there is a significant difference in the immune measure between groups.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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