Vaccination Using Recombinants Influenza and Adenoviruses Encoding Amastigote Surface Protein-2 Are Highly Effective on Protection against Trypanosoma cruzi Infection

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Abstract

In the present study we evaluated the protection raised by immunization with recombinant influenza viruses carrying sequences coding for polypeptides corresponding to medial and carboxi-terminal moieties of Trypanosoma cruzi amastigote surface protein 2 (ASP2). Those viruses were used in sequential immunization with recombinant adenovirus (heterologous prime-boost immunization protocol) encoding the complete sequence of ASP2 (Ad-ASP2) in two mouse strains (C57BL/6 and C3H/He). The CD8 effector response elicited by this protocol was comparable to that observed in mice immunized twice with Ad-ASP2 and more robust than that observed in mice that were immunized once with Ad-ASP2. Whereas a single immunization with Ad-ASP2 sufficed to completely protect C57BL/6 mice, a higher survival rate was observed in C3H/He mice that were primed with recombinant influenza virus and boosted with Ad-ASP2 after being challenged with T. cruzi. Analyzing the phenotype of CD8+ T cells obtained from spleen of vaccinated C3H/He mice we observed that heterologous prime-boost immunization protocol elicited more CD8+ T cells specific for the immunodominant epitope as well as a higher number of CD8+ T cells producing TNF-α and IFN-γ and a higher mobilization of surface marker CD107a. Taken together, our results suggest that immunodominant subpopulations of CD8+ T elicited after immunization could be directly related to degree of protection achieved by different immunization protocols using different viral vectors. Overall, these results demonstrated the usefulness of recombinant influenza viruses in immunization protocols against Chagas Disease.

Introduction

Over a hundred years after its first description, Chagas Disease remains as an important public health problem, mostly in Latin America. Nonetheless, the infection rate is increasing in other continents, mostly by blood transfusion [1,2]. According to WHO, there are currently over 10 million people infected in Latin America and more than 100 million people live at risk areas in endemic countries. Moreover, this disease kills approximately 15 thousand people every year, due to the clinical complications and to the poor efficacy of the pharmacological treatment which is highly toxic and effective mostly during the acute phase of disease [3,4]. In addition, the resistance of parasites to chemotherapy is another major drawback to the pharmacological treatment [5,6,7]. Thus, the development of vaccines is an important approach to be used in therapy and prophylaxis of Chagas disease [3,8].

Many vaccination studies against Chagas’ disease already provided evidence that CD8+ T cells play pivotal role on the development of protective immunity [9,10,11,12]. Mechanisms used by these cells to eliminate the parasite include directly killing of infected cell or secretion of cytokines such as IFN-γ [13,14]. Among the antigens that have been studied as potential candidates for vaccine development, the surface amastigote protein 2 (ASP2) has been found as one of the most promising [15,16]. In addition, different strategies have already been tested to deliver this antigen in mice, including the use of recombinant protein, plasmid DNA...
and recombinant viruses [17,18,19,20]. For instance, our group demonstrated that two sequential immunizations with recombinant HA5 adenovirus encoding ASP2 were able to significantly reduce the parasitemia and improve the survival of vaccinated mice, when they were challenged with Y strain of *T. cruzi* [18]. However, in spite of these very promising results, a drawback in the same viral vector in sequential immunizations rely on the risk that anti-vector antibodies generated after the priming could neutralize the vector when it is used in further immunizations and, consequently, hinder the boost of heterospecific immune response [21,22]. The limitation of anti-vector response elicited by homologous prime-boost immunization could be surpassed by different strategies, such as the use of two different recombinant viruses on prime and boost immunizations [23,24].

Live recombinant influenza viruses have some features that make them attractive to be used in vaccination protocols against protozoan infections, as we can mention: They are well known inducers of Cytotoxic T Lymphocytes (CTLs) by direct infection of immature dendritic cells (DCs) and monocytes, facilitating antigen (Ag) presentation both local and systemically [25,26,27]; It is feasible to generate recombinant influenza viruses by reverse genetics techniques [28]; There are different influenza A strains and subtypes, which could be used in sequential immunizations to overcome previous immune responses directed to the vector [29].

Therefore, in the present study we exploited the use of recombinant influenza viruses carrying truncated sequences of ASP2 in sequential immunization with adenovirus encoding ASP2. This immunization protocol elicited potent anti-ASP2 cellular immune response, reduced the parasite burden and improved the survival of vaccinated mice when they were challenged with *T. cruzi*.

**Materials and Methods**

**Mice and Ethics**

Male of eight- to ten-weeks-old C57BL/6 and C3H/He mice were obtained from René Rachou Research Institute’s (CPqRR) animal facility center (Fiocruz, Belo Horizonte, Brazil) and housed according to institutional standard guidelines. All animal studies were approved by the Ethical Commission on Animals’ Use (CEUA) at Oswaldo Cruz Foundation (Fiocruz), license LW-949, and performed following institutional Guide for the Care and Use of Laboratory Animals.

**Cells and Parasites**

MDCK and 293T cells (obtained from Pasteur Institut, FR) were grown at 37°C under 5% CO₂ in complete Dulbeccos modified Eagle Medium (DMEM; SIGMA) with 1 mM sodium pyruvate, 4.5 mg/ml L-glucose, 100 U/ml penicillin and 100 µg/ml streptomycin (herein called complete DMEM medium) and respectively supplemented with 5% or 10% heat inactivated fetal calf serum (FCS; CULTILAB) [30]. Trypomastigotes from respectively supplemented with 5% or 10% heat inactivated fetal blood in frame to the sequence coding for k chain of mice immunoglobulin that allows the secretion of the foreign sequence [17]. Those constructs were used as PCR templates to generate IgSP-M or C-ASP2 segments which were site directed cloned into XhoI and XbaI digested pRNA38 vector (Figure 1B). All primers sequences are available under request and the respective presenting haplotype were referenced within the correspondent portion (Figure 1A) [34,35]. The generated plasmids (pPRNA38-M-ASP2 and pPRNA38-C-ASP2) were analyzed using Dynamic ET Dye Terminator Cycle Sequencing KIT® (AMESHAM) and a Megabace 1000 automatic sequencer (AMESHAM).

Details of Laboratory Animals.

**Viral RNA Extraction, RT-PCR Analysis**

Viral RNA (vRNA) extraction from cell-free supernatants of infected MDCK cultures and RT-PCR analysis were performed as previously described [33]. Amplicons were analyzed on 1% agarose gel and visualized by ethidium bromide staining. RT-PCR products were purified and presence of mutations was determined by sequencing using Dynamic ET Dye Terminator Cycle Sequencing KIT® (AMESHAM) and a Megabace 1000 automatic sequencer (AMESHAM).

**Peptides**

Peptides VNHRFTLV and TEWETGQI were purchased from Genescript (Piscataway, NJ). Peptide purity was in higher than 90%. Their identities were confirmed by Q-TOF Micro equipped with an electrospray ionization source (Micromass, United Kingdom).

**ELISPOT and Intracellular Cytokine Staining**

ELISPOT assay was performed essentially as previously described [37]. Spleens cells of immunized mice were obtained
three weeks after boost immunization. They were treated with ACK buffer for erythrocytes lysis and washed twice in RPMI containing 5% FBS before to be resuspended in cell culture medium consisting of RPMI 1640 medium (pH 7.4) supplemented with 10 mM HEPES, 0.2% sodium bicarbonate, 59 mg of penicillin/liter, 133 mg of streptomycin/liter, and 10% fetal bovine serum (CULTILAB) containing recombinant IL-2 (100 U/ml). The viability of the cells was evaluated by using 0.2% trypan blue exclusion dye to discriminate between live and dead cells. The number of spleen cells was adjusted to 1\(\times10^6\) cells per well in cell culture medium and stimulated with specific peptides at final concentration of 10\(\mu\)g/ml of VNHRFTLV (aa 553–560; for C57BL/6 splenocytes) or TEWETGQI (aa 320–327; for C3H/He splenocytes). The spots were counted on a S5 Core ELISPOT Analyser (CTL).

For Intracellular Cytokine Staining, the cell concentration was adjusted to 1\(\times10^6\) cells per well in cell culture medium containing GolgiStop™ and GolgiStop™ (according to manufacturing instructions; BD Pharmingen) and -phycoerythrin (PE) anti-CD107a (BD Pharmigen). In half of the cultures, a final concentration of 10\(\mu\)g/ml of VNHRFTLV (for C57BL/6 splenocytes) or TEWETGQI (for C3H/He splenocytes) peptide was added. The cells were cultivated in U-bottom 96-well plates (Corning) in a final volume of 200 \(\mu\)l at 37°C in a 5% CO\(_2\) humid atmosphere. After 12 hour-incubation, cells were stained for surface markers fluorescein isothiocyanate (FITC)-labeled dextramer TEWETGQI (Immudex), after 10 minutes incubation, cells were also stained with peridinin chlorophyll protein complex (PerCP) anti-CD8, avidin-phycoerythrin (PeCy7) anti-CD8, or FITC-labeled anti-CD3 (in samples without dextramer) antibodies (BD Pharmigen). The cells were fixed and permeabilized using Cytofix/Cytoperm kit (BD, Biosciences) according to manufacturer’s recommendations. Cells were then stained for intracellular markers allophycocyanin (APC) anti-IFN-\(\gamma\), APC-Cy7 anti-TNF-\(\alpha\), or PE Cy7 anti-TNF-\(\alpha\) (BD Pharmigen). Finally, the cells were fixed in 2% PBS-paraformaldehyde and at least 100,000 cells were acquired on a FacsCanto, LSRFortessa or FacsAria II (BD, Biosciences) flow cytometers and then analyzed with FlowJo software (ThreeStar). The ancestry gates are represented in Figure S1.

**ELISA and Western Blot**

Recombinant ASP2 (rASP2) protein was produced in Escherichia coli as previously described [17]. The presence of sera specific anti-ASP2 antibodies were assessed by enzyme-linked immunosorbent assay (ELISA) on immunized mice sera obtained fourteen days after the boost immunization. Briefly, plates (Maxisorb, NUNC) were coated with 4 \(\mu\)g/ml (His\(_{6}\)KDa, rASP2) and incubated at 4°C overnight. Mice sera were diluted 1:100 in blocking buffer and incubated for 2 hours at 37°C. Plates were incubated with peroxidase-conjugated goat anti-mouse IgG (SIGMA) one hour at room temperature, and reactions were developed with 3,3',5,5'-tetramethylbenzidine (TMB) reagent (SIGMA) and read at 450 nm. Alternatively, 0.5 \(\mu\)g of His\(_{6}\)KDa, rASP2 were loaded on 12% polyacrylamide gels and transferred to nitrocellulose membranes. Membranes were then blocked and incubated with individual sera of mice immunized with recombinant viruses. After extensive washes, membranes were incubated with peroxidase-conjugated...
goat anti-mouse IgG (SIGMA) and detection was performed by membrane exposure to X-ray films after a standard chemoluminiscence reaction (ECL Detection System, Amersham Biosciences).

To measure IFN-γ production, spleen cells were obtained as described above and incubated for 72 hours at 37°C, 5% CO₂. The IFN-γ concentration was determined in cell culture supernatant with DuoSet ELISA Development System mouse IFN-γ kit (R&D Systems) according to manufacturer’s recommendations.

**Immunizations**

Heterologous prime-boost immunizations were performed as previously described [30]. Briefly, the animals were lightly anesthetized with a mixture of ketamine and xylazine and inoculated by intranasal route (IN) with 10⁶ plaque-forming unit (pfu) of recombinant influenza viruses (Flu-CT or Flu-nASP2) diluted in 25 µl of PBS. Four weeks later, the animals were boosted with 5x10⁷ pfu of recombinant Ad-ASP2 or Ad-CT in 100 µl of PBS by subcutaneous route (SC). Alternatively, some animals received two immunizations with 5x10⁶ pfu of recombinant Ad-ASP2 or Ad-CT in 100 µl of PBS by subcutaneous route (SC). In contrast, some animals received two immunizations with 5x10⁶ pfu of recombinant Ad-ASP2 or Ad-CT by SC route four weeks apart (homologous prime-boost immunization protocol). Finally, some mice received only one immunization with 5x10⁷ pfu of recombinant Ad-ASP2 by SC route.

**Statistical Analysis**

Data are expressed as ± SEM and analyzed using GraphPad Prism ver.5 Software. Statistic significance for ELISA, ELISPOT and cytokine staining assays were evaluated using One-Way ANOVA and non-parametric test followed by Bonferroni post-test. Statistical significance for parasitemia was evaluated by 2-way ANOVA with Bonferroni post-test. The Gehan-Breslow-Wilcoxon test was performed to compare mouse survival curves.

**Results**

Generation and Characterization of Recombinant Influenza Viruses

Recombinant influenza viruses harboring the medial or the carboxi-terminal sequence of ASP-2 protein were recovered using the 12 plasmid driven reverse genetics as previously described [30]. These recombinant viruses, which were respectively named Flu-M-ASP2 and Flu-C-ASP2, displayed lysis plaques in MDCK cells similar in size to those found in cells infected with the recombinant Flu-CT. In contrast, these viruses displayed lysis plaques that were slightly smaller than those of the wild type WSN virus (Figure 1C). In addition, their infectious titers (1.4x10⁶ pfu/ml Flu-M-ASP2 and 2.8x10⁶ pfu/ml Flu-C-ASP2) were significantly lower than those of WSN virus (1x10⁷ pfu/ml).

As shown in figure 1D, amplifications products of expected size (~1000 bp) were found for each recombinant influenza virus assayed. Moreover, when these amplicons were analyzed by sequencing, we found no mutations, demonstrating that those recombinant influenza viruses were genetically stable in cell culture (data not shown).

Evaluation of Humoral Immune Response

Immunization protocols were carried out according to the schedule depicted at figure 2A. Two weeks after the boost immunization, specific anti-ASP2 IgG serum antibodies were measured by ELISA and western blot, using the recombinant ASP2 (His65KDa) protein as capture antigen. Western blot results showed that specific anti-ASP2 IgG antibodies could be found in sera of all C57BL/6 mice primed with Flu-C-ASP2 and boosted with Ad-ASP2 (figure 2B), whereas only one animal that received a single immunization with Ad-ASP2 displayed detectable levels of specific anti-ASP2 antibodies. In addition, we detected higher levels of specific anti-ASP2 antibodies in the sera of mice primed with recombinant influenza than those found in animals that received only one immunization with Ad-ASP2 (Figure 2C). Interestingly, neither by Western blot (data not shown) nor ELISA (figure 2D), we were able to detect specific anti-ASP2 antibodies in sera of C3H/He mice immunized with recombinant viruses, irrespective the immunization protocol used in vaccination. It is noteworthy that previous studies demonstrated that B epitopes are located in C-terminal moiety of ASP-2 protein and humoral immune response against intra-cellular amastigote proteins is not essential for protection [34,30].

Specific Cellular Immune Response Against Protective Epitopes

The activation of specific anti-ASP2 CD8⁺ T cell response was measured in spleen of immunized mice by stimulating their splenocytes with VNHRFTLV (H-2Kk restricted, C57BL/6) or TEWETGQI (H-2Kk restricted, C3H) peptides, three weeks after the boost immunization. As depicted in figure 3, specific IFN-γ producing CD8⁺ T cells could be found in spleen cells of mice primed with Flu-C-ASP2 or Flu-M-ASP2 and boosted with Ad-ASP2 (figure 3A and C). In addition, high amounts of IFN-γ could be measured in spleen cell culture supernatants stimulated ex-vivo with their respective peptides (Figure 3B and D). Interestingly, in both cases, there was a clear improvement in the prime-boost immunization, as we could find a significant increase in IFN-γ production on prime-boosted groups compared to single Ad-ASP2 immunized mice (Figure 3).

Protection Against Experimental Infection

The protection afforded by the vaccination protocols was evaluated by challenging the vaccinated mice with 500 (C3H/He) or 1000 (C57BL/6) bloodstream Y strain trypomastigotes. Regarding the resistant mice strain, C57BL/6, a single immunization with Ad-ASP2 sufficed to reduce the parasitemia and to completely protect the animals comparing to control immunized groups (Figure 4A, p<0.05; and 4B, p<0.001).

Regarding the C3H/He mice, which display remarkable susceptibility to T. cruzi, infection groups that received at least a single immunization with recombinant adenovirus-ASP2 were able to reduce the peak of parasitemia (Figure 4C, p<0.001), control tissue pathology (Figure S2) and prolong survival compared to the groups immunized with control recombinant viruses (Figure 4D p<0.0005). Remarkably, a higher survival rate was found in mice that were primed with Flu-M-ASP2 and boosted with Ad-ASP2 as close to 80% of vaccinated mice survived, comparing to mice that were primed with Flu-C-ASP2 and boosted with Ad-ASP2 (p=0.0019). They also showed significant improvement of survival when compared to single or prime-boosted Ad-ASP2 immunized mice (p=0.05, single and 0.08 homologous groups; Figure 4D).

In order to verify if the improvement of survival rate induced by the Flu-Ad protocol could be due to recombinant influenza properties, we tested the usefulness of a homologous intranasal prime subcutaneous-boost immunization using Flu-M-ASP2 virus in C3H/He mice strain. As demonstrated in Figure S3, we could not observe the production of specific immune response under stimulation neither by ELISPOT (Figure S3) nor intracellular staining for IFN-γ and TNF-α (data not shown) in splenocytes derived from homologous immunized mice. This could be expected since a single immunization with recombinant influenza
is known to elicit neutralizing antibodies that can prevent a proper boost against the heterologous M-ASP2 polypeptide [30,32].

**Cellular Immune Response Profile Elicited by Different Immunization Protocols**

The survival results found in C3H/He mice prompted us to study more deeply the cellular immune profile elicited by the immunization protocols. To this aim, C3H/He mice were immunized as previously described and three weeks after the boost immunization, spleen CD8\(^+\) T cells were evaluated for intracellular staining of IFN-\(\gamma\) and TNF-\(\alpha\) cytokines and for the surface mobilization of CD107a upon ex vivo stimulation with peptide TEWETGQI, as described in Material and Methods section. As depicted in figure 5A, the percentage of CD8\(^+\) T cells positive for at least one of the parameters evaluated were similar in mice that received two immunizations with recombinant viruses, irrespective the immunization strategy employed.

Regarding the phenotype of subpopulations of CD8\(^+\) T cells found in vaccinated mice, triple (IFN-\(\gamma\), TNF-\(\alpha\), CD107a) and double (IFN-\(\gamma\)+CD107a+) and IFN-\(\gamma\)+TNF-\(\alpha\)+ positive cells are the major populations that were found after immunization with recombinant viruses encoding ASP2 (Figure 5B). Interestingly, mice immunized with recombinant viruses encoding ASP2 displayed similar percentage of CD8\(^+\) T cell subpopulations, irrespective if they were immunized according to heterologous or homologous immunization protocols, and similar to IFN-\(\gamma\) production seen by ELISPOT and ELISA, there was a clear impact of boost immunization in the frequency of specific effector CD8\(^+\) T cells comparing prime-boosted groups with Ad-ASP2 single immunized group (Figure 5A and 5B).

In order to perform a more accurate analysis on CD8\(^+\) T cells elicited by immunization, we used a specific H-2K\(d\)/TEWETGQI dextramer. Mice were immunized as previously described and the phenotype of specific CD8\(^+\) T cells was assessed in TEWETGQI stimulated pooled spleen cells of vaccinated mice three weeks after the last immunization. As depicted in figure 5C, mice vaccinated with Flu-ASP2/Ad-ASP2 displayed the highest number of total dextramer positive CD8\(^+\) T cells. The main subpopulation of dextramer positive CD8\(^+\) T cells that were found in mice immunized irrespective the tested protocols were triple positives (IFN-\(\gamma\), TNF-\(\alpha\), CD107), followed by single (CD107+) positives CD8\(^+\) T cells (Figure 5D). On the other hand, we could observe a higher frequency of single IFN-\(\gamma\)+TEWETGQI+ CD8\(^+\) T cells (CD107a- TNF-\(\alpha\)-) in heterologous (19%) and Ad-ASP2 single (12.5%) immunized groups compared to Ad-ASP2/Ad-ASP2 group (2.9%).

Accordingly, Table 1 shows that besides heterologous Flu-Ad immunization elicited higher numbers of CD8\(^+\)TEWETGQI+ T cells, also the frequency of CD8\(^+\)TEWETGQI+ CD107a and/or IFN-\(\gamma\) and/or TNF-\(\alpha\) positive cells increase above two fold compared to homologous or single immunized groups. We could also find a significant increase of perforin production under stimulation only in the heterologous vaccinated group (Figure S4). This results suggest the importance of those effector factors on protection and could also indicate that the improvement of survival by heterologous could be due to a higher number of effector specific CD8\(^+\) T cells.

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**Figure 2. Immunization Schedule and Induction of specific anti-ASP2 humoral immune response in mice vaccinated with recombinant viruses.** Timeline representation of immunization schedule and experimental procedures (A). C57BL/6 mice were immunized as described in Material and Methods. Two weeks after the boost immunization, the animals were bled and the presence of specific anti-ASP2 total IgG antibodies in mice sera was evaluated by western by incubating individual (lanes 1–9) or pooled (lanes 10 and 11) sera of C57BL/6 mice with nitrocellulose membranes loaded with recombinant ASP2 protein (His65KDa) as capture antigen blot (B). Alternatively, the antibodies levels were measured by ELISA using individual sera of C5BL/6 (C) or C3H/He (D) mice sera diluted 1:100 and recombinant ASP2 protein as capture antigen. Optical Density (OD) was measured at 450 nm.

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Discussion

Recombinant viruses carrying foreign sequences have been proven to be useful tools as vaccines against many pathogens, including those which require the induction of potent type I T cell immune responses, such as *Leishmania s.p.*, *Toxoplasma gondii* and *Trypanosoma cruzi* [39]. Studies carried out by our group demonstrated that two immunizations with recombinant adenovirus carrying *T. gondii* or *T. cruzi* antigens were able to elicit specific humoral and cellular immune response and to protect different mouse lineages after challenge with those protozoan parasites [18,37]. In spite of these very promising results, two immunizations with recombinant adenovirus (homologous immunization protocol) raises some concerns, mostly due to the elicited anti-vector immune response, which could hurdle the immune response directed against the foreign sequence in subsequent vaccinations. This problem could be surpassed by using two different vectors for each immunization [24].

Therefore, we evaluated the use of recombinant influenza viruses encoding ASP2 derived polypeptides as a tool for priming the specific anti-ASP2 immune response, followed by sequential immunization with a recombinant adenovirus encoding ASP2. Regarding the naturally resistant C57BL/6 mice, the prime with recombinant influenza virus encoding the carboxy-terminal portion of ASP2 was as useful as recombinant adenovirus in priming specific anti-ASP2 immune response. Indeed, antibodies levels and the number IFN-γ producers CD8+ T cells specific for ASP2 were similar in mice primed with recombinant influenza or adenovirus. Interestingly, regarding C57BL/6 mice strain, a single immunization with recombinant adenovirus suffice to control parasitemia and to completely protect the animals after challenge. Similar findings were obtained by Duan and collaborators using recombinant Sendai virus encoding ASP2, which was able to significantly reduce the parasitemia and to completely protect C57BL/6 mice after the challenge with Tulahuen strain [19].

Regarding the susceptible strain C3H/He, all immunizations protocols employing Ad-ASP2 in our study were able to significantly reduce the parasitemia, control at certain extent tissue pathology and prolong survival of challenged animals. Considering Y strain of *Trypanosoma cruzi*, there is a variable correlation between blood parasitemia and survival rate, as demonstrated in different mice strains [40,41]. However, we could observe a correlation of parasitemia control with prolonged survival in our model. Notwithstanding, an improvement on the survival rate was observed in mice primed with
recombinant influenza-M-ASP2 and boosted with recombinant adenovirus even when compared to the survival of the animals that were immunized once or twice with recombinant adenovirus, or primed using Flu-C-ASP2 which does not contain an immunodominant epitope to C3H/He MHC-I haplotype. These results seemed quite surprising because the specific anti-ASP2 cellular immune response, measured by the number of specific CD8$^+$ T as well as by production of IFN-$\gamma$ was similar in mice that were submitted either to the homologous or heterologous prime and boost immunization protocols.

Phenotype analyses performed on total CD8$^+$ T cells obtained from vaccinated C3H/He showed that most effector CD8$^+$ T cells were polyfunctional and mostly triple (IFN-$\gamma$, TNF-$\alpha$ and CD107a) and double (IFN-$\gamma$, CD107a) positives. These results were similar to those obtained previously in C57BL/6 mice that were immunized with naked DNA and adenovirus encoding ASP2 [42]. Our results also showed that mice that received one immunization with Ad-ASP2 displayed similar CD8$^+$ T cells phenotype than those observed in mice that received two immunizations with recombinant viruses encoding ASP2, suggesting that just one immunization with Ad-ASP2 suffice for shaping the CD8$^+$ T cells phenotype. Thus, our observations indicate that a single immunization using Ad-ASP2 suffice to stimulate a significant production of effector cytokines IFN-$\gamma$, TNF-$\alpha$ and mobilize CD107a, elicit an immunodominant effector population which can control parasitemia, reduce tissue pathology and prolong survival when compared to control immunized mice even in a susceptible model. This is particularly important because often, studies using different recombinant viruses or other vaccine vectors, mice models and Trypanosoma cruzi strains without the single immunized group could be overestimating their protection using prime-boost protocols.

Remarkably, C3H/He mice that were primed with Flu-M-ASP2 displayed higher number of dextramer positives CD8$^+$ T cells than mice that were immunized with Ad-ASP2. Moreover, our results showed that a boost immunization with Ad-ASP2 did not augment the number of TEWETGQI dextramer positive CD8$^+$ T cells in mice primed with Ad-ASP2. To discuss the reason by which the heterologous prime-boost protocol could improve protection and enhance the frequency of TEWETGQI CD8$^+$ T cells we hypothesized that immunization with Flu-M-ASP2, which encodes only the medial moiety of ASP2, primed the immunodominant CD8$^+$ response towards TEWETGQI epitope resulting in the expansion of this population after boosting with Ad-ASP2. In contrast, priming with Ad-ASP2, which carries the entire sequence of ASP2, could possibly elicit immune response also against subdominant epitopes of ASP2, resulting in a lower secondary response against TEWETGQI immunodominant epitope after boost [34,38,42,43,44]. Accordingly, previous results of our group suggest that immunization with plasmids or adenovirus encoding parasitemia, reduce tissue pathology and prolong survival when compared to control immunized mice even in a susceptible model. This is particularly important because often, studies using different recombinant viruses or other vaccine vectors, mice models and Trypanosoma cruzi strains without the single immunized group could be overestimating their protection using prime-boost protocols.

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Figure 4. Parasitemia and survival curves of immunized mice challenged with T. cruzi. B6 and C3H/He mice were immunized as described in Material and Methods. Four weeks after boost immunization, they were challenged intraperitoneally with 1000 and 500, respectively, T. cruzi $Y$ strain bloodstream trypomastigotes. Parasitemia was monitored on blood and depicted as the number of bloodstream trypomastigotes per milliliter of blood (A, n = 4; C, n = 8). The survival of vaccinated C57BL/6 (B, n = 7) and C3H/He (D, n = 7–9) mice was followed during 50 days and showed as Kaplan-Meier curves. * p<0.05, ** p<0.001.

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C3H/He mice were immunized with recombinant viruses as described in Material and Methods. Two weeks after the last immunization, the spleen cells were harvested and cultivated ex vivo with specific TEWETGQI CD8+ T peptide and incubated with anti-CD3, anti-CD8, permeabilized and fixed with anti-CD107a, anti-IFN-γ and anti-TNF-α antibodies and assessed by flow cytometry. Percentage of effector CD8+ T cells reacting to the presence of TEWETGQI peptide obtained from spleen cells of mice immunized with recombinant viruses (A). Percentage of CD8 T cells which produces IFN-γ or/and TNF-α or/and mobilizes the degranulation marker CD107a after stimulation with TEWETGQI (B), the statistics depicted are compared to groups of mice immunized with control recombinant viruses. The number and frequency of dextramer positive CD8+ T found in 3×10^6 CD8+ T (C). Functional profile of CD8+ T cells subpopulations obtained from mice immunized with recombinant viruses (D). Response were depicted with different color patterns according to the number of assessed functions (IFN-γ, TNF-α and CD107a) displayed by each dextramer negative or dextramer positive CD8+ T cells subpopulations.

Figure 5. Phenotype of anti-ASP2 specific CD8+ T cells elicited by vaccination with recombinant viruses.

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ASP2 subdominants epitopes afforded lower degree of protection when compared to that observed in animals immunized with vectors encoding the immunodominant epitope [42]. A reinforcement to this hypothesis could be found in the low number of proteins encoded by influenza when compared to adenovirus, which potentially reduces the number of viral antigens that could compete with the heterologous antigen for presentation by antigen-presenting cells [45,46]. The correlation between TENVETGQI (present in medial portion of ASP2, M-ASP2) immunodominant frequency and protection is reinforced by the result of C3H/He mice that were immunized using recombinant influenza encoding the C-terminus portion of ASP2 as prime and Ad-ASP2 as boost presented a survival curve similar to single immunized or homologous prime-boost using Ad-ASP2 after infection (p = 0.46).

Another finding of our study was that the number of dextramer stained CD8+ T cells producing IFN-γ, TNF-α or the surface marker CD107a found in animals primed with Flu-M-ASP2 were approximately three times higher than those observed in other vaccinated groups. However, while the role of different T cell subpopulation to control the infection with some viruses, bacteria and Plasmodium was already well documented [47,48,49,50], the biological relevance of CD8+ T cells subpopulations phenotypes to control the infection with T. cruzi remains elusive [51]. The IFN-γ production itself is known to be important for protection against Trypanosoma cruzi infection in many previous work of our and other groups [14,52,53,54,55]. On the other hand, other factors as the effector phenotype of specific CD8+ T cells, the production of perforin, the recirculation of those cells out of spleen [56], their presence in the heart [57,58], apoptosis of specific immunodominant anti-ASP2 CD8+ T cells [59], and the type of memory cells involved are important to be considered [51]. Recently a group has elegantly shown that multiple redundant effector CD8+ T cells factors deriving from transferred Tc1 and Tc17 populations are capable of protecting mice against viral infection [60], and as CD8+ T cells have a major role in protection against Trypanosoma cruzi infection, this statement is an interesting subject of research. Thus, if the improvement of protection observed in mice primed with recombinant influenza-M-ASP2 virus was only due to the higher number of CD8+ T specific for the immunodominant epitope or could also be due to other factors remains to be solved.

In summary, we demonstrated that recombinant influenza viruses encoding an ASP2 derived polypeptide would be useful in heterologous prime-boost studies aiming the development of vaccines against Chagas Disease. The priming with recombinant influenza virus followed by boost with recombinant adenovirus could properly augment the number of effector CD8+ T cells specific for ASP2 immunodominant epitope, whose displayed unique phenotype and resulted in increased survival of vaccinated C3H/He mice challenged with T. cruzi.

Supporting Information

Figure S1 Representative of ancestry gates for flow cytometry experiments. Correspondent ancestry gates for the figure 5 analysis. (TIF)

Figure S2 Histopathological analyses of liver, spleen and heart derived from infected mice. Male C3H/He mice were primed and boosted according different immunization protocols and infected with 500 bloodstream trypomastigotes of Y strain of T. cruzi. Fifteen days after the infection, mice were euthanized and spleen, liver and heart were harvested, fixed and processed for histopathology. The organ sections were stained using hematoxinelin-eosin and the degree of tissue inflammation was evaluated (scale bar - 100 µm). (TIF)

Figure S3 Cellular responses to immunodominant epitope from ASP2 in mice immunized twice using Flu-M-ASP2. ELISPOT of stimulated splenocytes taken from C3H/He mice immunized with the depicted protocols. The prime-boost was performed within an interval of 28 days and the experiment was performed 21 days post boost. The splenocytes were incubated 18 h in the presence of 10 µg of TENVETGQI peptide (n = 5 for all groups except non-immunized group NI/NI, n = 3). (TIF)

Figure S4 Perforin production in splenocytes derived from C3H/He immunized mice. Splenocytes derived from immunized C3H/He mice were ex vivo stimulated or not in the presence of Brefeldin A and Monesin A and the immunodominant peptide TENVETGQI for 12 hours, prepared, labeled and submitted to flow cytometry (n = 4). N.I. Non-immunized/Non-infected. Their staining profiles were analyzed using FlowJo and statistical analysis performed was 2-Way ANOVA with Bonferroni post-test using GraphPad Prism 5.0 Software. (TIF)

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Author Contributions

Conceived and designed the experiments: RPAB DCC MMR RTG AVM. Performed the experiments: RPAB BGF LIS PASJ RVSP PEM AVM. Wrote the paper: RPAB AVM.

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