Fusion of the *Mycobacterium tuberculosis* Antigen 85A to an Oligomerization Domain Enhances Its Immunogenicity in Both Mice and Non-Human Primates

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**Abstract**

To prevent important infectious diseases such as tuberculosis, malaria and HIV, vaccines inducing greater T cell responses are required. In this study, we investigated whether fusion of the *M. tuberculosis* antigen 85A to recently described adjuvant IMX313, a hybrid avian C4bp oligomerization domain, could increase T cell responses in pre-clinical vaccine model species. In mice, the fused antigen 85A showed consistent increases in CD4+ and CD8+ T cell responses after DNA and MVA vaccination. In rhesus macaques, higher IFN-γ responses were observed in animals vaccinated with MVA-Ag85A IMX313 after both primary and secondary immunizations. In both animal models, fusion to IMX313 induced a quantitative enhancement in the response without altering its quality: multifunctional cytokines were uniformly increased and differentiation into effector and memory T cell subsets was augmented rather than skewed. An extensive *in vivo* characterization suggests that IMX313 improves the initiation of immune responses as an increase in antigen 85A specific cells was observed as early as day 3 after vaccination. This report demonstrates that antigen multimerization using IMX313 is a simple and effective cross-species method to improve vaccine immunogenicity with potentially broad applicability.

**Introduction**

A major challenge in vaccinology is the development of effective vaccines against intracellular pathogens where cell mediated immunity plays an important protective role. Viral vectored vaccines have a remarkable capacity to induce and boost antigen-specific T cells [1], but higher frequency responses will likely be required to achieve useful protective efficacy [2]. There is, therefore, a need for adjuvants in the next generation of vectored vaccines to increase T cell immunogenicity.

Oligomerization is employed by many natural proteins to increase protein valency, binding affinity and structural stability [3], and while a pantameric coiled coil was initially used to improve B-cell responses in mice [4] its sequence is too similar to its human ortholog to be considered safe for use in humans. Recently, a series of homologous oligomerization domain proteins were shown to act as adjuvants in mice, resulting in an augmentation of both B and T cell responses [5,6]. These proteins were derived from the domain encoded by the last exon of the complement 4 binding protein (C4bp) α-chain. This exon encodes the only domain not involved in the complement related functions of C4bp and is both essential and sufficient for oligomerization of the seven C4bp alpha chains [7] and a number of other proteins [8,9,10,11,12]. Fusion of recombinant *Plasmodium yoelii* MSP-119 protein to C4bp oligomerization domains from a variety of mammalian and avian species was shown to improve antibody responses to this weak immunogen [6]; of the domains tested, a chicken C4bp hybrid with less than 20% homology to human C4bp, called IMX313, was shown to induce the highest titers of MSP-119 specific antibodies, buts its effect on T cell responses was not investigated.

Tuberculosis (TB) remains one of the most serious worldwide infectious despite the use of the *M. bovis* strain Bacillus Calmette-Guérin (BCG) as a vaccine since the 1920s. The most advanced sub-unit vaccine in clinical development is a modified vaccinia virus Ankara (MVA) expressing the *M. tuberculosis* protein 85A. Clinical trials in both the UK and Africa have shown the substantial capacity of MVA-Ag85A to boost T cell responses to BCG in healthy individuals [13,14,15], whether this capacity is maintained in HIV-infected individuals is unclear. Although there...
is currently no clear correlate of protection, T cells have been shown to play an important role [16,17,18] and therefore strategies to increase the level of vaccine induced T cells could have a substantial impact.

In this study we have assessed whether fusion to the IMX313 domain could enhance the T cell mediated response to the M. tuberculosis antigen 85A in two pre-clinical animal species, mice and rhesus macaques. In both animal models, IFN-γ ELISpot and multi-parameter flow cytometry were used to investigate the effects of fusion to IMX313 on the overall quality of the immune response in terms of cytokine secretion and generation of effector or memory T cell subsets. Fusion of antigens to the IMX313 domain is a straightforward method and its ability to enhance T cell immune responses could have broad applicability across a variety of animal species and disease settings. In the first instance, we intend to undertake a direct comparison in humans of MVA-Ag85A with MVA-Ag85A IMX313 to confirm the results described here in two very distinct animal species.

Results

Fusion to IMX313 multimerizes antigen 85A

To confirm that fusion of IMX313 resulted in formation of disulphide-linked multimers without affecting protein expression, cell lysates from MVA infected BHK cells were analyzed by Western blots of reducing and non-reducing SDS-PAGE gels (Figure 1). Under reducing conditions, both Ag85A and Ag85A IMX313 proteins migrated at the expected apparent molecular weight of the mature monomeric peptide (35.5 kDa for Ag85A or 40.4 kDa for Ag85A IMX313) (Figure 1d). For both proteins, an additional higher molecular weight band (~45 kDa for Ag85A and ~50 kDa for Ag85A IMX313) was observed, probably representing a glycosylated species, since a predicted N-linked glycosylation site (NNT) is present within Ag85A (http://www.cbs.dtu.dk/services/NetNGlyc/), though this conclusion has not been experimentally verified. More importantly, when the reducing agent was omitted (Figure 1b and 1c), the Ag85A IMX313 fusion protein migrated exclusively in excess of ~200 kDa, consistent with formation of the predicted heptamers (248 kDa). Although other higher-order species were observed, these results indicate that the IMX313 domain was functioning as expected to induce antigen multimerization.

Fusion of antigen 85A to IMX313 enhances CD4+ and CD8+ T cell responses in mice

Initial screening experiments were performed in mice to assess the ability of IMX313 to enhance antigen 85A-specific responses. Balb/c mice received two intramuscular (i.m.) immunizations with either 50 µg of DNA-Ag85A or DNA-Ag85A-IMX313 two weeks apart and the immune response to Ag85A measured in the blood or spleen by IFN-γ ELISpot. Following a single immunization of 50 µg DNA plasmid vaccine, significant increases in the response to the dominant CD4+ (p15) and CD8+ (p11) T cell epitopes were observed in PBMCs from mice vaccinated with the Ag85A IMX313 compared to Ag85A vaccinated controls (Figure 2A). These increases were enhanced after a second homologous DNA immunization (Figure 2A) and a significant increase in the p11-specific response was observed in CD8+ splenocytes two weeks after the final vaccination (Figure 2B) (2.1-fold increase). A small 1.5-fold increase in the CD4+ p15 response was also detected but did not reach statistical significance (Figure 2B).

For the assessment of IMX313 in an MVA vector, Balb/c mice were vaccinated intramuscularly (i.m.) or intradermally (i.d.) with 10⁶ PFU of either MVA-Ag85A or MVA-Ag85A IMX313 and the immune responses measured in the spleen of mice by IFN-γ ELISpot (Figure 2C and D). One week after i.m. vaccination, significant increases in both p15 (2.3-fold) and p11 (3.3-fold) specific responses were observed in MVA-Ag85A IMX313 vaccinated mice compared to MVA-Ag85A controls (Figure 2C), with a similar enhancement observed in intradermally (i.d.) immunized mice (Figure 2D). MVA vaccines were subsequently compared in a variety of prime-boost combinations to determine the priming versus boosting capacity of the IMX313 fusion in two separate strains of mice: Balb/c and C57BL/6. Mice received an i.m. priming vaccination with either MVA-Ag85A or MVA-Ag85A IMX313 followed two weeks later with either 10⁶ PFU of MVA-Ag85A or MVA-Ag85A IMX313 delivered via the i.d. route (Figure 2E and F). Two weeks after the second immunization, the highest response in either strain of mice was observed after homologous immunization with MVA-Ag85A IMX313 (Figure 2E and 2F). In Balb/c mice, a 2.2-fold increase in the mean CD4+ T cell response (95% CI 1.82–2.60) and 3.3-fold increase in the mean CD8+ T cell response (95% CI 2.52–4.00) was observed when compared to mice homologously vaccinated with MVA-Ag85A (Figure 2E). In summary, fusion of IMX313 to
antigen 85A consistently enhanced T cell responses in the two vaccine delivery systems and in two strains of mice.

Fusion of antigen 85A to IMX313 enhances the immune response in rhesus macaques

While mice have proved to be excellent indicators of immunogenicity, the ability to predict efficacy from murine tuberculosis models has often been disappointing [19] and larger animals may be better predictors of human immunogenicity and efficacy. As the primary focus of this study was to test the capacity of IMX313 to increase T cell responses, we performed further immunological assessments in non-human primates.

Rhesus macaques were immunized with 10^6 PFU of MVA-Ag85A or MVA-Ag85A IMX313 at weeks 0 and 6, and the response to antigen 85A was measured by IFN-γ ELISpot 1 week after vaccination, then every 2 weeks using a single pool of 66

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peptides (Figure 3A) and 7 pools of 20 peptides each (Figure 3B). The peak of the antigen 85A-specific responses was observed 1 week after each vaccination (Figure 3A and B) consistent with previous data in mice and humans [20]. At multiple timepoints, higher antigen 85A responses were observed in the group of animals vaccinated with the MVA-Ag85A IMX313 when compared to those vaccinated with MVA-Ag85A (Figure 3). We conducted two separate analyses to determine if there was a statistically significant effect of IMX313. Firstly, without making distributional assumptions, we noted that over the weeks 1–13, the total number of Ag85A IFN-γ secreting cells in the MVA-Ag85A IMX313 group was greater than those in the MVA-Ag85A control group (Kruskal-Wallis test, $p = 0.02$), this was also true with single peptide pools A, B, C, D, and G ($p = 0.05$, $0.07$, $0.05$, $0.08$, $0.02$, respectively, Figure S1). In a second approach, we constructed a parametric model of the responses to the seven individual peptide pools to derive an estimate of IMX313 induced effect of 1.8-fold (95% CI 1.31–2.41). Despite the increased variance in immune responses to Ag85A detected in non-human primates, the magnitude of the increase was similar in the two species.

To measure whether vaccination with IMX313 induced cross-reactive T cells to C4bp, frozen PBMCs were thawed and the response to 85A peptides, IMX313 peptides and hC4bp peptides measured by IFN-γ ELISpot. In agreement with the data from freshly isolated PBMCs, a statistically significant increase in the antigen 85A response was observed 1 week after the second vaccination in MVA-Ag85A IMX313 animals compared to the controls (Figure 3C left panel). Only 1 animal had a detectable T cell response to IMX313 peptides (Figure 3C central panel) and importantly no detectable response to human C4bp was observed in any of the macaques (Figure 3C right panel). Antibodies to 85A, IMX313 and human C4bp were also assessed but no detectable responses above background to any of the antigens were observed (data not shown).

In summary, consistent with the data observed in mice, fusion to IMX313 resulted in an overall increase in the responses to the antigen 85A induced by MVA vaccination and importantly did not result in the induction of cross-reactive T cells to the oligomerization domain of C4bp.

**Fusion of antigen 85A to IMX313 does not alter the quality of the immune response in mice and macaques**

A number of recent studies have suggested that the quality of the immune response, as determined by cytokine polyfunctionality, can have a substantial effect on the overall protective ability of a vaccine regime [21,22]. While there is no clear correlate of protection to TB, T cells have a protective effect [16,17,18] and any second-generation TB vaccine should ideally augment the level of both CD4+ and CD8+ T cell immunogenicity while maintaining its overall quality.

We therefore investigated whether fusion to IMX313 altered the quality of the immune response in terms of cytokine secretion and differentiation into effector and memory T cells. In the MVA prime-boost experiments described above (Figure 2E), splenocytes were stimulated with a single pool of antigen 85A analyzed for the intracellular production of IFN-γ, IL-2, TNF-α and simultaneously stained for CD62L and CD127 to discriminate between the three main population of T effector and memory cells [23,24]. In Balb/c mice, CD4+ T cells were shown to produce predominately all of the three cytokines measured (IL-2, IFN-γ and TNF-α: triple positive cells) or IL-2 alone (Figure 4A) and homologous prime-boost with MVA-Ag85A IMX313 was shown to induce the highest frequency of both of these subsets when compared to the control MVA-Ag85A vaccinated group (Figure 4A). This increase in frequency did not come at a cost to other cytokine producing subsets, as there was no change in the proportion of cytokine producing cells that secreted either 1, 2 or all 3 cytokines (Figure 4A pie-charts). CD8+ T cells demonstrated a difference in the cytokine profile compared to CD4+ T cells, the predominant cytokine producing population observed was CD8+ T cells that simultaneously produced IFN-γ and TNF-α (double positive) or IFN-γ only (Figure 4B). An overall increase in the frequency of IFN-γ and TNF-α or IFN-γ only producing CD8+ T cells in MVA-Ag85A IMX313 vaccinated mice did not alter the other subsets, as equal proportions of cells producing 1, 2 or all 3 cytokines were observed between all groups of mice (Figure 4B pie-charts). To analyze the differentiation of antigen-specific cells into effector and memory T cell subsets, any cell producing at least one cytokine (i.e. IFN-γ or TNF-α or IL-2) was analyzed for the expression of surface markers of CD127 and CD62L and separated into CD127+ CD62L+ T central memory (Tcm) cells, CD127−, CD62L− T effector memory (Tem) cells or CD127−, CD62L+ T effector (Teff) cells. A higher frequency of both CD4+ Tcm cells (Figure 4C) and CD8+ Tcm cells (Figure 4D) were observed in MVA-Ag85A IMX313 vaccinated mice when compared to MVA-Ag85A controls (Figure 4C). In all groups of animals, around 90% of either CD4+ or CD8+ cytokine producing T cells displayed a Tem phenotype, indicating that the fusion of antigen 85A to IMX313 did not skew the distribution of measured phenotypes.

A similar analysis of quantity versus quality was performed on the non-human primate samples one week after the second MVA vaccination. Frozen PBMCs were restimulated with a single pool of antigen 85A peptides and cells analyzed for upregulation of CD107α (a marker of degranulation) [25], intracellular production of IFN-γ, IL-2 and TNF-α and surface expression of CD95 and CD28. The predominant CD4+ T cell sub-population seen was cells secreting TNF-α alone or in combination with IFN-γ, IL-2 or both (i.e. triple positive cells) (Figure 5A). Consistent with the data in mice, MVA-Ag85A IMX313 vaccinated animals displayed an overall increase in the frequency of these populations, but this did not result in a change in the proportion of cells producing either 1, 2 or all 3 cytokines (Figure 5A pie-charts). Substantially lower overall responses were observed in CD8+ T cells (Figure 5B graphs) and in the MVA-Ag85A IMX313 group, two strong responders drove a shift towards a higher proportion of cells producing all three cytokines and upregulating CD107α (Figure 5B graphs). Importantly, the weaker responding animals in this group displayed an identical proportional distribution to animals vaccinated with MVA-Ag85A (Figure 5B pie-charts), suggesting it is the overall size of the immune response and not an effect of IMX313 that was driving an increase in CD8+ T cell multi-functionality. At the same time, any cell producing one or more cytokines or CD107α was analyzed for expression of CD28 and CD95 and further subdivided into CD28−CD95− Tcm, CD28+CD95− Tcm or CD28+CD95+ Tcm cells [26,27,28]. One week after a second vaccination with MVA, Tcm was the predominant phenotype of antigen 85A specific CD4+ T cells observed (Figure 4C), consistent with previous data from macaques [28]. An increase in the total frequency of CD4+ Tcm cells was seen in MVA-Ag85A IMX313 vaccinated animals (Figure 5C left panel) yet this did not alter the differentiation of cells into either Teff Tcm or Tcm (Figure 5C middle panel). This trend was less clearly observed for CD8+ T cells due to the overall lower responses (Figure 5D). However it is clear that strong CD8+ T cell responding animals displayed a predominately Tcm phenotype (Figure 5D right panel), while weaker responders displayed equal distribution of cells into Teff Tcm and Tcm cells.
regardless of the vaccine regime (Figure 5D). In summary, the data from both mice and macaques demonstrate that fusion to IMX313 increased the frequency of antigen 85A specific cells without altering the overall quality of the immune response generated in terms of multi-functionality or differentiation into effector or memory T cell subsets.

IMX313 has a very early effect on the immune response

An increase in antigen persistence, stability and/or uptake as a result of multimerization or binding of IMX313 to a specific (in vivo) target receptor, are all conceivable mechanism of IMX313 induced immune response enhancement. As all of these mechanisms could have an early effect on the immune response, we characterized the in vivo response as early as 3 days after MVA vaccination.

We then characterized the cellular composition of the lymph node to determine whether IMX313 was mediating an effect on APC recruitment and activation. Three days after MVA vaccination, draining auricular lymph nodes were harvested to investigate the frequency and number of CD4+ T cells, CD8+ T cells, B cells, macrophages and more specifically a number of

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**Figure 3. Macaque IFN-γ ELISpot responses.** Male rhesus macaques were immunized i.d. at weeks 0 and 6 with either 10^8 PFU MVA-Ag85A or MVA-Ag85A IMX313 and the response to antigen 85A measured in the blood by IFN-γ ELISpot before vaccination (pre) and every 2 weeks from week 1 onwards. Graphs represent the response of each individual animal to a single pool containing all antigen 85A peptides (A) or the sum of 7 separate peptide pools (B, C) Graphs represent the ELISpot responses of frozen PBMCs to a pool of antigen 85A peptides, a pool containing IMX313 peptides and a pool containing hC4bp peptides, pre-vaccination and at week 1 or week 7 post-vaccination. Bars represent the median group response with each animal displayed as a single point. Data was analyzed with a 2-way analysis of variance with a post-hoc Bonferroni test; the asterisks denote the level of statistical significance (*p* < 0.05).

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different dendritic cell (DC) subsets (Figure S2A). At this timepoint, there were significant changes in the composition of draining lymph nodes in terms of the frequency of each cell population (Figure S2B), but all lymphocyte populations were shown to increase in total number (Figure S2C). The most dramatic increase was observed for CD11c⁺ DCs (Figure S2C) primarily due to an influx in CD11b⁺ MHC IIintermediate (int) DCs (Figure S2D), although other DC subsets did also increase in number (Figure S2D, E). Recruitment of these DCs was antigen 85A independent, as a similar frequency and number were present after vaccination with an irrelevant MVA (Figure S2F, G). IMX313 did not however alter the activation state of CD11b⁺ MHC II⁺ DCs, as no change in CD80 (Figure S2H) or MHC class II expression (Figure S2I) was observed.

While we observed no prolonged antigen 85A persistence or specific effect of IMX313 on lymph node cell populations, these
experiments did not rule out an early effect in the initial in vivo antigen dose, which could also be affected by stability and uptake of antigen 85A. As changes in T cell frequencies and activation marker expression has been observed in vivo with alterations in antigen dose [29,30], we investigated both of these factors as a potential surrogate marker of in vivo antigen dose. Balb/c mice were immunized with MVA-Ag85A or MVA-Ag85A IMX313 and draining lymph nodes harvested 3 days later. Using intracellular IFN-γ staining to identify antigen 85A-specific cells, we observed an early increase (2-fold) in the frequency of both CD4+ (Figure 6A)

Figure 5. Quality of the response to antigen 85A in rhesus macaques 1 week after a homologous MVA-Ag85A boost. A & B) Graphs represent the frequency of each cytokine producing subset as a percentage of all CD4+ (A) or CD8+ (B) T cells, while the pie charts represent the proportion of cytokine producing cells positive for 1, 2, 3 or all 4 markers. C & D) Any cell positive for either CD107a, IFN-γ, TNF-α or IL-2 was further subdivided into a CD28+CD95- Tem, CD28-CD95- Tem or CD28+CD95+ Tem cell. Graphs on the left represent the absolute frequency of each memory population relative to the total number of CD4+ (C) or CD8+ (D) T cells, centre graphs represent the percentage of each population relative to cytokine producing cells, and graphs on the right represent total percentage of cytokine+ cells compared to the percentage of each individual T cell population. For each graph the bars represent the median group response, with each animal displayed as a single point. Data were analyzed with a 2-way analysis of variance, no statistical difference between the two groups of vaccinated animals was observed.

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and CD8+ (Figure 6B) T cells in MVA-Ag85A IMX313 vaccinated mice compared to MVA-Ag85A controls. Despite this increase, we observed no statistically significant changes in the levels of a number of T cell activation markers investigated (Figure S3), demonstrating once again that IMX313 was not skewing the induction of the T cell responses in any particular way.

In summary, while this in vivo characterization did not suggest a specific receptor for IMX313, we observed an increase in antigen specific cells as early as three days after MVA vaccination, showing that the effect of IMX313 occurs very early in the immune response.

Discussion

In this study we have used the M. tuberculosis antigen 85A as a model to demonstrate the potent cross-species adjuvant effect of IMX313. Straightforward procedures were used to incorporate this adjuvant into two vectored vaccine systems, and an immune-potentiating effect was subsequently demonstrated in both mice and non-human primates. In both animal models the fusion of antigen 85A to IMX313 resulted in an augmentation of the response without skewing it towards a particular cytokine-producing population as has been observed with some traditional adjuvants [31,32].

Our detailed in vivo characterization of the response three days after vaccination did not reveal a specific activation effect of IMX313. However at this timepoint, we observed a significant increase in the frequency of antigen 85A specific CD4+ and CD8+ T cells (Figure 6). The two-fold increase at day 3 was equivalent to the fold increase observed at day 7, and since this fold increase was maintained over the course of the response, it would suggest that IMX313 does not affect the rate of T cell division. Therefore the increase in antigen specific cells occurs as a result of an early event in T cell activation, most likely an increase in the effective in vivo antigen dose, i.e. amount of antigen presented. This is supported by our data showing no shift in the kinetics or quality of the 85A immune response by fusion to IMX313. In addition, the consistent effect of IMX313 in DNA and MVA and in vivo different vaccination routes would support a mechanism that relies on an early antigenic effect, as opposed to a specific recruitment of cells or receptors that may not be accessed by different vectors or vaccination routes.

It is not clear how multimerization enhances the effective in vivo antigen dose. Fusion to C4bp was originally proposed to increase the serum half-life of proteins and their immunogenicity [6], but multimerization may itself result in an increase the uptake of antigen 85A by APCs due to increased avidity. Antigen multimerization may also lead to prolonged antigen presentation by making antigen 85A more difficult to process [35]. In an elegant study, Delmarre et al demonstrated that by increasing the lysosomal susceptibility, both T and B cell immunogenicity was reduced in vivo. Bins et al [34] subsequently demonstrated that increasing in vivo luciferase stability resulted in increased levels of T cell immunogenicity induced by DNA vaccination without altering the kinetics of the immune response, supporting the data observed in this study. Fusion of antigen 85A to IMX313 may either increase serum half-life, improve antigen uptake, prolong antigen processing or all of the above and further experiments are underway to try and address some of these possibilities.

While many molecular adjuvants fall victim to species differences in signaling and activation pathways, multimerization takes advantage of the immune system’s natural response to an increase in effective in vivo antigen dose, so that its effect is consistently observed through different vaccine platforms and in two very different animals species. Thus multimerization as an adjuvant platform should have broad applicability to a variety of diseases where cell-mediated immunity plays an important role, e.g. malaria, TB, HIV and influenza. In summary, this relatively simple technology of antigen multimerization could be useful in a wide range of vaccine programs, both veterinary and human, as well as for some non-communicable diseases.

Materials and Methods

Plasmids, viruses and cells

A 229 bp DNA fragment encoding the IMX313 domain was PCR amplified from the plasmid pIMX313 [6] using the oligonucleotides 5'-GAAGCCCGACCTGCAACGTGGATCCAAGAAGC ATGCTGATG-GAGCGGCCGCTTATTACTCCTTGCTCAGTC- and 3'-AAAGCCCTCCCTAGATGCTC-GAGCGGCGCTTATTACTCCTTGCTCAGTCCTTGCG- 3' and inserted into the DNA vaccine vector pSG2-Ag85A [35] using site-directed mutagenesis [36] #. This replaced the 9 amino acid PK tag at the C-terminus of antigen 85A protein with the 55 amino acid IMX313 domain. Ag85A IMX313 was conventionally subcloned into the poxvirus shuttle vector pMVA-GFP-TD which places it under the control of the vaccinia virus p7.5 promoter and provides flanks for recombination at the TK locus of MVA. Markerless recombinant viruses were isolated following transfection of this plasmid into infected primary chick embryo fibroblasts (CEFs) (Day 9 of gestation) (Institute for Animal Health) by transient dominant selection with GFP. For immunizations, viruses were amplified in CEFs, purified over sucrose cushions, and titered twice in duplicate by plaque assay according to standard methods. Virus identity and purity was confirmed by PCR.

Ethics Statement

Mice were used in accordance with the UK Animals (Scientific Procedures) Act under project license number 30/2414 granted by the UK Home Office.

Male rhesus macaques (Macaca mulatta) aged between 2 1/2 to 6 years were maintained and used at the Institute of Neurobiology and Molecular Medicine, National Research Council of Italy, in accordance with approval granted by the ethics committee of the Italian Ministry of Health (Decree by low n°: 2007 114 B and n°: 2007 115 C). The animal care routine and experimental procedures were in compliance with national and international laws and policies (EEC Council Directive 86/609; Italian
Legislative Decree 116/92). Animal welfare was ensured and steps taken to ameliorate suffering in accordance with the recommendations of the Weatherall report, “The use of non-human primates in research.” To minimize distress and suffering related to vaccination or blood withdrawal, animals were anesthetized by i.m. injection of 10 mg/kg ketamine hydrochloride for all procedures and no animals were sacrificed at the end of the experiment.

Animals and Immunizations
Female Balb/c or C57BL/6 mice of 6 weeks of age or older (Harlan, UK) were immunized i.m. in the musculus tibialis or intradermally (i.d.) in the ear with a total volume of 50 µl of vaccine diluted in PBS. Mice received either 50 µg of DNA or 10⁶ PFU of MVA per vaccination. Rhesus Macaques were assigned into each group to ensure homogeneous distribution of age and body weight. The control group (MVA-Ag85A) included a total of 4 animals with an average weight of 5 kg (range 3.9–7.5 kg) and age of 4 years (range 2.8–5.9); the test group (MVA-Ag85A IMX313) included 5 animals with an average weight of 4.90 kg (range 4–7.55) and average age of 4 years (range 3.4–6). Animals received two intradermal immunizations, at weeks 0 and 6, with of 10⁶ PFU of MVA, with arms switched between vaccinations. Blood samples were kept at room temperature and shipped overnight to Oxford where all processing and assays were performed.

Western Blots
Monolayers of BHK-21 cells (European Collection of Cell Cultures) were infected for 24 hours with MVA-Ag85A, MVA-Ag85A IMX313 or an unrelated recombinant MVA at 4 PFU per cell. Cells were washed in PBS and harvested into non-reducing Laemmli’s sample buffer. After addition of β-mercaptoethanol to half of each sample, the lysates were processed for SDS-PAGE and Western blotting by standard methods. Lanes were loaded with volumes of lysate corresponding to approximately 10⁵ infected cells. Blots were labeled with chicken anti-Ag85A (AbCam) or a hyperimmunun serum from rabbits immunized with IMX313 protein, followed by alkaline phosphatase conjugated secondary antibodies (Jackson) and colorimetric visualization.

Peptides
Murine spleenocytes were stimulated for a total of 6 hours with a single pool containing all antigen 85A peptides (2 µg/ml final concentration) or media only (unstimulated control), with the addition of Golgi-Plug (BD) (1 µg/ml) for the final 4 hours. Following surface staining with CD4+APCAlexa780, CD8 PerCP-Cy5.5, CD62L-PECy7 and CD127 Pacific Blue, cells were fixed with 4% paraformaldehyde and stained intracellularly with TNF-α Alexa488, IL-2 PE and IFN-γ Alexa647 diluted in Perm-Wash buffer (BD).

ELISPots
Mice spleens or peripheral blood samples were treated with ACK lysis buffer to remove RBCs prior to stimulation with the relevant peptides (final concentration of 2 µg/ml) on IPVH-membrane plates (Millipore) coated with 5 µg/ml anti-mouse IFN-γ (AN18). Macaque PBMCs were isolated over a lymphoprep gradient (Sigma) while frozen PBMCs were thawed and rested overnight in media containing 10 U/ml Benzonase (Sigma). 250 000 PBMCs per well were stimulated with relevant peptides at a final concentration of 2 µg/ml on IPVH-membranes coated with 15 µg/ml anti-human IFN-γ (GZ4).

After 18–20 hours of stimulation, IFN-γ spot forming cells (SFC) were detected by staining membranes with either anti-mouse IFN-γ biotin (1 µg/ml) (R46A2) or anti-human IFN-γ (7B6-1) followed by streptavidin-Alkaline Phosphatase (1 µg/ml) and development with AP conjugate substrate kit (BioRad, UK).

Intracellular cytokine staining
Murine splenocytes were stimulated for a total of 6 hours with a single pool containing all antigen 85A peptides (2 µg/ml final concentration) or media only (unstimulated control), with the addition of Golgi-Plug (BD) (1 µg/ml) for the final 4 hours. Following surface staining with CD4+APCAlexa780, CD8 PerCP-Cy5.5, CD62L-PECy7 and CD127 Pacific Blue, cells were fixed with 4% paraformaldehyde and stained intracellularly with TNF-α Alexa488, IL-2 PE and IFN-γ Alexa647 diluted in Perm-Wash buffer (BD).

Frozen macaque PBMCs were thawed and rested overnight in media containing 10 U/ml Benzonase (Sigma). 2×10⁶ PBMCs were restimulated for 6 hours in 96 well U-bottom plates (total volume of 200 µl) with 1 µg/ml anti-human CD49d (9F10), 20 µl anti-human CD28-APC (CD28.2) and 20 µl of anti-human CD107a-PECy5 (eBIOH4A3) in addition to 2 µg/ml antigen 85A peptide pool or DMSO (unstimulated control) with the addition of Golgi-Plug (1 µg/ml) and Golgi-Stop (0.8 µg/ml) for the final 4 hours. Cells were surface stained for CD95-biotin (eBioscience) followed by strepavidin qDot 565, CD4 eFluor605 (OKT4), Live-Dead Violet (Invitrogen), CD14 eFluor 450 (61D3) and CD20 eFluor 450 (2H7). Cells were fixed with 4% paraformaldehyde and stained intracellularly with TNF-α FITC (MAb11), IL-2 PE (MQ1-17H12), IFN-γ PECy7 (48.B3). CD3 Alexa700 (SP34-2) and CD8-APC Alexa780 (RPA-T8) diluted in Perm-Wash buffer.

Sample acquisition was performed on a LSR II and data analyzed in FlowJo (TreeStar), Pestle v.1.6 and Spice v.4.3 [41] with all final figures made in Canvas X (ACD Systems).

Statistical methods
All graphs and statistical analysis were created in Prism v5.0c (Graphpad), data were log transformed and a one-way or two-way analysis of variance performed; figure legends denote the test performed for each experiment. Where IMX313 was shown to induce a significant effect, a Kruskal Wallace (one-way ANOVA) or Bonferroni test (two-way ANOVA) was performed and p values stated in the figure legend.

For analysis of macaque peptide pool data, Kruskal-Wallis tests were used to compare the responses from animals at baseline (pre-immunization) with those at all timepoints after immunization. Parametric models estimating the effect of IMX313 modeled log(number of IFN-γ secreting cells, detected by ELISPOT) as an outcome variable, because of the approximate log-normal distribution of ELISPOT counts in these animals (not shown). Conditions with no spots were scored as having 1 spot, which is the limit of detection of the assay. Modeling of Log(ELISPOT-)

T−Effect of IMX313+effect of Timepoint+effect of peptide pool allowed estimation of an overall effect of IMX313. Assumptions include (i) the responses to the peptide pools are independent, (ii) the effect IMX313 is similar over all peptide pools, (iii) the effect of IMX313 is similar over timepoints, (iv) effect of IMX313 is similar across animals. Analyses used R2.1.1 for Windows and Graph Pad Prism software.
Supporting Information

Figure S1 Immune responses to MVA-Ag85A and MVA-Ag85A IMX313 in macaques. Shown are individual responses to seven peptide pools (A–G) and a 6 peptide pool (‘Pool’) in five animals receiving MVA-Ag85A IMX313 (+) and four animals receiving MVA-Ag85A control (o) before vaccination (shown at t = 0), and at 1,3,5,7,9 and 13 weeks after the first vaccination (at t = 0, dashed vertical line); a second vaccination was given at week 6 (dashed vertical line). Geometric means are shown at each time point, joined by a solid line (Ag85A) or a dashed line (Ag85A IMX313). Conditions with no detectable response are shown as having 1 spot (see Methods). p values test the hypothesis that responses in the Ag85A-IMX313 post vaccination exceed those in the Ag85A group (Kruskal-Wallis test).

(TIF)

Figure S2 IMX313 does not induce activation of APCs. Balb/c mice were immunized i.d. with 10^6 PFU MVA-Ag85A (circles) or MVA-Ag85A IMX313 (squares) or unvaccinated (diamonds) and draining auricular lymph nodes harvested 3 days later for analysis by flow cytometry. Organs were diced into small pieces and digested at room temperature for 30 minutes in PBS containing 0.3 mg/ml collagenase-dispase (Roche) and 0.02 mg/ml DNase. For initial characterization of MVA vaccinated mice, LNs were surface stained for CD80-FITC, CD11c-PE, CD8-PerCPCy5.5, B220-PECy7, F4/80 Pacific Blue, CD103-biotin, streptavidin qDot 565, CD4-qDot 650, CD11b Alexa-700, MHC Class II APC-Alexa780 and Live-Dead Aqua prior to fixation and intracellular staining with CD207-APC (eBioscience) in perm-wash buffer. A) FACS plots demonstrate the gating strategy to classify the population of CD11c^+ B220^− CD11b^+ MHC II^int relative to plasmacytoid DCs (pDCs), lymph node (LN) resident DCs, or skin migrating DCs isolated from the draining auricular nodes (top panel) or non-draining inguinal and popliteal nodes (bottom panel) three days after id vaccination. CD11b^+ MHC II^int DCs were overlaid over the population of LN resident DCs to investigate expression of CD4 and CD8, or overlaid over the population of skin migrating DCs to investigate the expression of CD103 and CD207. Histograms represent the expression of either F4/80 (left) or CD80 (right) on LN resident (grey filled line), skin migrating (black dashed line) or CD11b^+ MHC II^int (black bold line) DCs. Graphs represent the percentage of live cells (B) or total number (C) of B cells (B220^+ CD11c^−), CD4^+ CD8^− (CD4^+ CD8^−), CD8^− (CD8^− CD4^−), DCs (CD11c^+) and CD11c^+ CD11b^+. CD11c^+ MHC II^int cells were further subdivided into plasmacytoid DCs (pDCs) (B220^+ ), CD11b^+ MHC II^int, LN resident (CD11b^+, MHC II^int) or skin migrating (MHC II^int) with graphs representing the percentage (D) or total number of each population (E). In a separate experiment, auricular LNs from Balb/c mice immunized id with 10^6 PFU of either MVA-GFP (triangle), MVA-Ag85A (circles), MVA-Ag85A IMX313 (square) or unvaccinated (diamonds) were analyzed at day 3 days by flow cytometry. Digested lymph nodes were surface stained for CD80-FITC, CD11c-PE, CD8-PerCPCy5.5, B220-PECy7, CD4-qDot650, CD11b-Alexa700, MHC Class II APC with the addition of DAPI prior to acquisition of samples on the LSR II. Graphs represent the percentage (F) and total number (G) of CD11b^+ MHC II^int cells or surface expression level of CD80 (H) or MHC Class II fluorescence (I) on CD11b^+ MHC II^int cells. In all graphs, bars represent the mean per group with each individual animal displayed as a single point.

(TIF)

Figure S3 Expression of activation markers on IFN-γ^+ CD4^+ or CD8^+ T cells 3 days after immunization. Balb/c mice were vaccinated i.d. with 10^6 PFU MVA85A or MVA85A-IMX313 and the draining auricular lymph node harvested 3 days later. After 6 hours of restimulation with the total 85A peptide pool, cells were surface stained with CD69-FITC, CD44-PE, CD8-PerCPCy5.5, CD62L-PECy7, CD127 Pacific Blue, CD122-bi followed by streptavidin dQaot655 (Invitrogen), CD4-eFluro 650 and CD25-Alexa700 prior to fixation and intracellular staining for IFN-γ Alexa647. Cells were subdivided into either CD4^+ T cells (A) or CD8^+ T cells (B) prior to analysis of IFN-γ and surface marker expression. Data was analyzed with a two-way analysis of variance and no statistically significant difference was observed for any marker investigated.

(TIF)

Table S1 Ag85A peptide pools. The table lists the separation of the 66 individual 20 mer peptide (overlapping by 10) spanning the length of Ag85A into the 7 peptide pools labelled A to G.

(DOCX)

Table S2 Amino acid sequences of C4bp oligomerization domains. Using a Clustal V Method of alignment (MegaAlign, DNASTAR Lasergene software), bold letters represents homologous amino acids between chicken, IMX313 and human protein sequences. Grey letters represent the amino acid changes between human and the predicted rhesus macaque sequence (accession no. XP_002801994). Italic letters represent the major sequence change made to chicken version 1 in construction of IMX313, to reduce sequence homology to the human sequence.

(DOCX)

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Author Contributions

Conceived and designed the experiments: AJ S FH MGC CSR JF KCS. Performed the experiments: AJ S JDH MGC CSR JF KCS. Analyzed the data: AJ JDH MGC DHW. Contributed reagents/materials/analysis tools: FH SJD SCG. Wrote the paper: AJ S FH MGC AVSH.

References

1. Draper SJ, Heaney JL (2010) Viruses as vaccine vectors for infectious diseases and cancer. Nat Rev Microbiol 8: 62–73.
2. Hill AV (2006) Pre-erythrocytic malaria vaccines: towards greater efficacy. Nat Rev Immunol 7: 381–389.
3. Engler J, Kammerer RA (2000) What are oligomerization domains good for? Matrix Biol 19: 293–300.
4. Chambers RS, Johnston SA (2003) High-level generation of polyclonal antibodies by genetic immunization. Nat Biotechnol 21: 1088–1092.
5. Draper SJ, Moore AC, Goodman AL, Long GA, Holder AA, et al. (2006) Effective induction of high-titer antibodies by viral vector vaccines. Nat Med 14: 819–821.
6. Oguin NA, Dumon-Seignonert L, Marchand JB, Holder AA, Hill F (2008) The oligomerization domain of C4-binding protein (C4bp) acts as an adjuvant, and the fusion protein comprised of the 19-kilodalton merosin surface protein 1 fused with the murine C4bp domain protects mice against malaria. Infect Immun 76: 3817–3823.
7. Kask L, Hillarp A, Ramesh B, Dahlback B, Blom AM (2002) Structural requirements for the intracellular subunit polymerization of the complement inhibitor C8binding protein. Biochemistry 41: 9349–9357.

8. Christiansen D, Devaux P, Revel P, Echalier A, Horvat B, et al. (2006) Octamerization enables soluble CD46 receptor to neutralize measles virus in vitro and in vivo. J Virol 74: 4672–4676.

9. Shinya E, Dervilless X, Edwards-Levy F, Duric V, Brison E, et al. (1999) In vivo delivery of therapeutic proteins by genetically-modified cells: comparison of organs and human serum albumin alginic-coated beads. Biomed Pharmacother 53: 471–483.

10. Oudin S, Libby MT, Goossens D, Dervilless X, Philibert F, et al. (2000) A soluble recombinant multimeric anti-Rh(D) single-chain Fv/CR1 molecule restores the immune complex binding ability of CR1-deficient erythrocytes. J Immunol 164: 1515–1513.

11. Libby MT, Goossens D, Oudin S, Gupta N, Dervilless X, et al. (1997) A recombinant human sFc anti-Rh(D) antibody with multiple valences using a C-terminal fragment of CR1-binding protein. Blood 90: 3978–3983.

12. Dervilless X, Huther A, Schuhmacher J, Griesinger C, Cohen JH, et al. (2006) Stable expression of soluble therapeutic peptides in eukaryotic cells by multimerisation application to the HIV-1 fusion inhibitory peptide C66g. ChemMedChem 1: 330–339.

13. Ilanga HB, Brooks RH, Hill PC, Osıaıe PK, Fletcher HA, et al. (2006) Early clinical trials with a new tuberculosis vaccine, MVA85A, in tuberculosis-endemic countries: issues in study design. Lancet Infect Dis 6: 522–528.

14. McShane H, Pathan AA, Sander CR, Keating SM, Gilbert SC, et al. (2004) Recombinant modified vaccinia virus Ankara expressing antigen 85A boosts BCG-primed and naturally acquired antityphocellular immunity in humans. Nat Med 10: 1240–1244.

15. Haweledge T, Scriba TJ, Gelderblom S, Smit E, Tameris M, et al. (2002) Safety and immunogenicity of a new tuberculosis vaccine, MVA85A, in healthy adults in South Africa. J Infect Dis 188: 544–552.

16. Brookes RH, Pathan AA, McShane H, Henemann M, Price DA, et al. (2003) CD8+ T cell-mediated suppression of intracellular Mycobacterium tuberculosis growth in activated human macrophages. Eur J Immunol 33: 3293–3302.

17. Caruso AM, Serbina N, Klein E, Triebold K, Bloom BR, et al. (1999) Mice deficient in CD4 T cells have only transiently diminished levels of IFN-gamma, yet succumb to tuberculosis. J Immunol 162: 5417–5426.

18. van Pinxteren LA, Cassidy JP, Smedegaard BH, Agger EM, Andersen P (2000) Multifunctional TH1 cells define a correlate of vaccine-mediated protection against Leishmania major. Nat Med 13: 843–850.

19. McShane H, Pathan AA, Sander CR, Goonetilleke N, Brookes R, et al. (2002) Protective immunity against Mycobacterium tuberculosis H37Rv infection. Infect Immun 66: 1527–1533.

20. Ulanova M, Tarkowski A, Hahn-Zoric M, Hanson LA (2001) The Common vaccine adjuvant aluminium hydroxide up-regulates accessory properties of human monocytes via an interleukin-1-dependent mechanism. Infect Immun 69: 1151–1159.

21. Delamarre L, Couture R, Mollman I, Trombeeta ES (2006) Enhancing immunogenicity by limiting susceptibility to lysosomal proteolysis. J Exp Med 203: 2049–2053.

22. Bins AD, Wolkers MC, van den Boom MD, Haanen JB, Schumacher TN (2007) In vivo antigen stability affects DNA vaccine immunogenicity. J Immunol 179: 2126–2133.

23. Taracha EL, Bishop R, Musoke AJ, Hill AV, Gilbert SC (2006) Heterologous priming-boosting immunization of cattle with Mycobacterium tuberculosis 85A vaccine adjuvant aluminum hydroxide up-regulates accessory properties of human monocytes via an interleukin-1-dependent mechanism. Infect Immun 69: 1151–1159.

24. Geiser M, Cebé R, Drewello D, Schmitt R (2001) Integration of PCR fragments at any specific site within cloning vectors without the use of restriction enzymes and DNA ligase. Biotechniques 31: 88–90, 92.

25. Denis O, Tanghe A, Pailliet K, Junier F, van den Berg TP, et al. (1996) Vaccination with plasmid DNA encoding mycobacterial antigen 85A stimulates a CD4+ and CD8+ T-cell epitopic repertoire broader than that stimulated by Mycobacterium tuberculosis H37Rv infection. Infect Immun 66: 1527–1533.

26. McShane H, Belkoubou S, Goonetilleke N, Brookes R, Hill AV (2002) Protective immunity against Mycobacterium tuberculosis induced by dendritic cells pulsed with both CD8(+)- and CD4(+)-T-cell epitopes from antigen 85A. Infect Immun 70: 1623–1626.

27. D’Souza S, Rosseels V, Romano M, Tanghe A, Denis O, et al. (2003) Mapping of murine Th1 helper T-cell epitopes of mycobacterial Ag85A, Ag85B, and Ag85C from Mycobacterium tuberculosis. Infect Immun 71: 485–493.

28. Goonetilleke NP, McShane H, Hanna CM, Anderson RJ, Brookes RH, et al. (2003) Enhanced immunogenicity and protective efficacy against Mycobacterium tuberculosis of bacille Calmette-Guérin vaccine using mucosal administration and boosting with a recombinant modified vaccinia virus Ankara. J Infect Dis 171: 1602–1609.

29. Roederer M, Nozai JL, Nason MC (2011) SPICE: exploration and analysis of postcryptic complex multivariate datasets. Cytometry A 79: 167–174.