A recombinant bovine adenoviral mucosal vaccine expressing mycobacterial antigen-85B generates robust protection against tuberculosis in mice

Highlights
- BCG-vaccinated infants need a booster for better protection against tuberculosis
- The BAdv^85C5 vaccine increased protection in BCG-vaccinated mice against tuberculosis
- The BAdv^85C5 vaccine increased antigen presentation and memory T cells responses
- Mucosal vaccination strengthens lung defense against tuberculosis

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In brief
Khan et al. describe a bovine-adenovirus-based BAdv^85C5 mucosal vaccine, which induced a galectin 3/8-dependent autophagy increasing antigen presentation in dendritic cells ex vivo. BAdv^85C5 nasal booster in BCG-vaccinated mice enhanced the protection against tuberculosis and increased antigen-specific CD4 T cells and memory T cells.
A recombinant bovine adenoviral mucosal vaccine expressing mycobacterial antigen-85B generates robust protection against tuberculosis in mice

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SUMMARY

Although the BCG vaccine offers partial protection, tuberculosis remains a leading cause of infectious disease death, killing ~1.5 million people annually. We developed mucosal vaccines expressing the autophagy-inducing peptide C5 and mycobacterial Ag85B-p25 epitope using replication-defective human adenovirus (HAdv85C5) and bovine adenovirus (BAdv85C5) vectors. BAdv85C5-infected dendritic cells (DCs) expressed a robust transcriptome of genes regulating antigen processing compared to HAdv85C5-infected DCs. BAdv85C5-infected DCs showed enhanced galectin-3/8 and autophagy-dependent in vitro Ag85B-p25 epitope presentation to CD4 T cells. BCG-vaccinated mice were intranasally boosted using HAdv85C5 or BAdv85C5 followed by infection using aerosolized Mycobacterium tuberculosis (Mtb). BAdv85C5 protected mice against tuberculosis both as a booster after BCG vaccine (>1.4-log10 reduction in Mtb lung burden) and as a single intranasal dose (>0.5-log10 reduction). Protection was associated with robust CD4 and CD8 effector (TEM), central memory (TCM), and CD103+/CD69+ lung-resident memory (TRM) T cell expansion, revealing BAdv85C5 as a promising mucosal vaccine for tuberculosis.

INTRODUCTION

Mycobacterium tuberculosis (Mtb) is a leading cause of mortality, with 8 million cases and ~1.5 million deaths each year. Bacillus Calmette-Guérin (BCG) is a widely used live-attenuated vaccine for the primary immunization of children worldwide. BCG protects mostly against extrapulmonary tuberculosis (TB), with variable protection against pulmonary disease ranging from 0% to 80%.1 BCG efficacy is influenced by several factors, including population genetics, pre-exposure to environmental mycobacteria, and its ability to induce efficient antigen presentation to T cells.2 We demonstrated earlier that BCG sequesters within immature phagosomes of antigen-presenting cells (APCs) with poor delivery to lysosomes.3,4 This sequestration results in the decreased presentation of the BCG-derived Ag85B-p25 epitope to CD4 T cells in mouse and human macrophages.3,5 We also reported that rapamycin can induce autophagy in APCs to enhance the delivery of BCG to lysosomes, thereby increasing antigen presentation to CD4 T cells in vitro5 and enhancing BCG efficacy.5,7 We have described a second-generation recombinant BCG vaccine overexpressing Ag85B protein (BCG85B) that induces autophagy in APCs and more effectively protects against TB relative to BCG.5 Our more recently described third-generation BCG85BC5 vaccine expresses TLR2-activating and autophagy-inducing peptide C5 (AIP-C5) from Mtb CFP10 protein and is more effective than BCG85B for the prevention of TB in mice.8 These studies suggest that autophagy induction can boost BCG efficacy. An autophagy-inducing recombinant BCG vaccine is under assessment in human clinical trials.9 Although millions of children receive BCG vaccination at birth, a larger number remain susceptible to lung TB.10 Accordingly, childhood TB remains common in many developing countries.
Because the main portal of entry for Mtb is respiratory, strengthening the neonatal lung immune response is a rational approach to prevent both pulmonary and extrapulmonary TB. Multiple studies suggest that viral vector-based vaccine platforms for Mtb are excellent in inducing both systemic and mucosal immunity.11–20 However, respiratory mucosal booster vaccines following BCG immunization may not offer adequate protection from TB in neonates owing to the defective expansion of resident memory T cells (T RM).21 Recent evidence indicates that the neonatal immune system is functional but physiologically immature, requiring vaccine adjuvants to elicit robust T helper (TH) cytokine responses in APCs.22,23 The relevance of an immature neonatal immune system in the context of disease is underscored by the requirement for childhood vaccine boosters for diptheria and tetanus toxoids, pertussis vaccine (DTP), and measles-mumps-rubella (MMR) vaccine. The immature neonatal immune system and the intrinsic defects of the BCG vaccine in APCs combined may explain the occurrence of lung TB and increased Mtb dissemination in neonates despite BCG vaccination.

Because BCG vaccination is continuing and there is an important need to protect children using safer vaccines, we sought to design an efficient vaccine platform capable of enhanced antigen presentation through autophagy. We have developed a bovine adenovirus (BAdv)-based TB vaccine expressing the immunodominant mycobacterial Ag85B-p25 epitope along with AIP-C5. Importantly, pre-existing adenovirus (Adv) antibodies do not interfere with the immunogenicity of BAdv vector-based vaccines.24–26 Our engineered mucosal vaccine augmented the ability of APCs to process and present the Ag85B-p25 epitope to CD4 T cells. Our nasal vaccine protected mice following aerosolized challenge with Mtb, both as a booster after BCG vaccine and alone. Finally, we observed a marked expansion of CD4 and CD8 effector (TEM), central memory (TCM), and CD103+/CD69+ T RM T cells in the lungs of vaccinated mice, suggesting that the vaccine protects mice from TB by inducing a robust pulmonary immune response.

RESULTS

Characterization of BAdv or human Adv (HAdv)-based vectors expressing the Mtb Ag85B-p25 epitope with or without AIP-C5

Adv vector-based vaccines elicit both humoral and cell-mediated immune (CMI) responses26,27 due to the adjuvant-like effect of Adv vectors by activating the innate immune system through both Toll-like receptor (TLR)-dependent and TLR-independent pathways.25,26 Influenza is one of the significant respiratory diseases in humans, animals, and birds. Adv vector-based influenza vaccines have conferred protective efficacy in both animal models28–31 and clinical trials in humans.14,32–35

We expressed the Mtb Ag85B-p25 epitope with or without AIP-C5 in the replication-defective BAdv or HAdv vector system.36 The Mtb Ag85B-p25 epitope (85) or 85 + AIP-C5 (85C5) gene cassette was under the control of the immediate early cytomegalovirus (CMV) promoter and bovine growth hormone (BGH) polyadenylation (PA) signal. The generated vectors BAdv85 (expressing 85), BAdv85C5 (expressing 85C5), HAdv85 (expressing 85), or HAdv85C5 (expressing 85C5) (Figure 1) were assessed for gene cassette expression using sequencing and RT-PCR (Figure S1A). HAdv constructs are shown in Figure S1B and were used in several experiments for comparison.

Replication-defective BAdv85C5 vaccine efficiently infects immature mouse dendritic cells (DCs) and induces a robust transcriptome

To characterize the virus-induced transcriptome, bone marrow-derived CD11c+ immature DCs from wild-type (WT)-C57BL/6 mice and autophagy-deficient ATG7 KO-DC mice37 were infected with BAdv85C5, BAdv vector, HAdv85C5, or HAdv vector followed by RNA sequencing (RNA-seq) analysis (Novogene). Figure 2A illustrates the heatmap of differentially expressed genes (DEGs) (n = 2) of BAdv- and HAdv-derived vaccine or vector-infected WT-DCs and ATG7 KO-DCs. ClusterProfiler analysis (Novogene) showed a relative enrichment of multiple genes in BAdv85C5- versus HAdv85C5-infected WT-DCs. KEGG (Kyoto Encyclopedia of Genes and Genomes) and Gene Ontology (GO; cellular components [CC] and biological processes [BP]) profiles are illustrated in Figures 2B–2E. Intriguingly, heatmaps of transcripts derived from RNA-seq expressed as FPKMs (fragments per kilobase per million mapped reads; n = 2) show that BAdv85C5 upregulated the expression of gene clusters involved in antigen processing and endosome sorting to lysosomes. For example, cathepsins (CTSA, CTSB, CTSK, CTSS, CTSL, and CTSZ) were enriched in BAdv85C5-infected DCs compared to HAdv85C5 (Figures 2F and 2G) (see also Figure S2). CTSSs proteolytically cleave antigens into peptides and help their loading onto the major histocompatibility complex class II (MHC class II), which are in turn, exported to the plasma membrane for activation of CD4 T cells. Furthermore, BAdv85C5 upregulated the
genes of the MHC pathway (H2-D1, B2m), and LAMP1, LAMP2, Hspa8, Hspa9, CD68 (LAMP4), and galectin3 (Gal3; also known as Lgals-3), which participate during the sorting of antigens and pathogens into the lysosomes.38-40 Transcripts for Lgals-8 were also found to be enriched BAdv85C5-infected DCs (mean FPKMs 510 ± 15; BAdv85C5/HAdv85C5, p < 0.001; data not shown). Finally, Gabarap (LC3 family), SQSTM1, and Rab7 were upregulated in BAdv85C5-infected DCs and are key players during autophagolysosome fusion.41,42 Many of these genes were also selectively upregulated by BAdv85C5-infected WT-DCs compared to ATG7KO-DCs (Figure S3), suggesting that BAdv85C5-upregulated genes are associated with autophagy. qPCR validation of the genes involved during antigen sorting and processing is shown in Figure S4 and is discussed in the context of vaccine-induced immunogenicity below.

Replication-defective BAdv85C5 vaccine is rapidly internalized by DCs and localizes to autophagolysosomal compartments

DCs are essential for vaccine-induced immune responses in both mice and humans.43 Earlier studies indicated that Cy3-fluorescent labeled Adv retain >98% infectivity.44 Therefore, we used Cy3-labeled BAdv85C5 or BAdv vector for the infection of mouse CD11c+ immature DCs followed by confocal microscopy. Since Adv can interact with the autophagy pathway and microtubule-associated light chain 3 (LC3) is a known marker of autophagosomes, LC3 was used as a marker for the virus containing endosomes.45 Cy3-BAdv85C5 and Cy3-BAdv vectors were rapidly internalized, and Figure 3 illustrates their uptake; both were found distributed between the cytosol and nucleus after a 4-h infection cycle, similar to a previous report.46 Endosomes colocalizing with the LC3 marker were quantitated, indicating an increased labeling of Cy3-BAdv85C5 endosomes compared to those containing Cy3-BAdv vector (Figure 3A). Similarly, lysosome-associated membrane protein-1 (LAMP1) labeling was higher, suggesting that Cy3-BAdv85C5 localizes more in autophagolysosomes than the vector (Figure 3B). Previous studies indicate that autophagy plays a role during the translocation of Adv capsids from the endosome to the nuclear pore complex, and during this process, lectin-like intracellular receptors Gal3 and Gal8 (Lgals-3/8) are associated with the vesicular transport of Adv.47,48 BAdv85C5-infected DCs showed a stronger enrichment of both Lgals-3 and Lgals-8 (Figures 3C and 3D). Isotype control is shown in Figure S5. It is pertinent to recall here that HAdv infects immature mouse DCs less effectively than a HAdv85C5 vaccine antigen processing (Figure S3). BAdv85C5-infected DCs are associated with the autophagy pathways in DCs, BAdv85C5 shows an enhanced expression of genes correlating with endosome traffic.

Replication-defective BAdv85C5 vaccine enhances autophagy-dependent and -independent antigen presentation in mouse DCs and human macrophages

Previous studies show that HAdv vectors expressing mycobacterial antigens protect mice and macaques against TB variably, although human studies have not been encouraging.13,19,50 Because BAdv85C5-induced robust gene expression in DCs, we sought to determine whether it increases the immunogenicity of DCs compared to HAdv85C5.

A major function of vaccine-ingested DCs is an efficient antigen processing and activation of CD4 and CD8 T cells through the MHC class II and the MHC class I pathway, respectively. Whereas vaccines degraded in lysosomes are routed to the MHC class II pathway, proteasome-digested peptides of vaccines are routed through MHC class I. We demonstrated earlier that autophagy can increase the MHC class I-dependent presentation of mycobacterial Ag85B,5 whereas autophagy was also reported to increase the MHC class I-dependent presentation of antigens in APCs.51 To define DC-mediated antigen-processing mechanisms, we used a well-characterized ex vivo assay wherein BCG- or Mtb-infected APCs rapidly present an Ag85B-derived p25 epitope to BB7 CD4 T cells ex vivo in the presence or absence of autophagy.3,52-55

Autophagy begins with an intracellular membrane vesicle nucleation, vesicle elongation, and autophagophore formation, which encloses pathogens in an autophagosome. The latter fuse with the lysosomes, which in turn, degrade pathogens generating antigenic peptides through CTS proteases. This process, also known as macroautophagy, involves several autophagy-regulating genes (ATGs), of which ATG7 and ATG5 are key genes, although alternative pathways exist.56 To determine whether autophagy plays a role during virus vaccine-induced antigen presentation, we infected WT-DCs and ATG7KO-DCs with BAdv85C5 or HAdv85C5 followed by antigen presentation. ATG7 deficiency in DCs led to a significant reduction in antigen presentation after infection with either BAdv85C5 or HAdv85C5 (Figure 4A). Supporting these data, WT-DCs and ATG7KO-DCs infected with BAdv85C5 showed striking differences in gene expression, indicating that autophagy is important during BAdv85C5 vaccine antigen processing (Figure S3). BAdv85C5 also showed upregulated antigen presentation in macrophages comparable to HAdv85C5 (Figure S4B) and enhanced gene expression compared to HAdv85C5 vaccine (Figure S6).

Because the BAdv85C5 vaccine platform is being developed for human neonates following BCG vaccination, we also treated human CD14+ macrophages with an autophagy inhibitor.
3-methyladenine (50 μM), followed by infection with BAdvBSCs or HAdvBSCs. Autophagy blockade reduced antigen presentation by both BAdvBSCs and HAdvBSCs in human CD14+ macrophages (Figures 4C and 4D). However, BAdvBSCs induced an elevated and sustained increase in antigen presentation in human CD14+ macrophages compared to HAdvBSCs (Figure 4E). Previously, we found that autophagy-mediated delivery of mycobacteria to lysosomes results in their degradation and reduced viability.2 Mouse CD14+ macrophages were infected with vectors or vaccines followed by infection with Mtb and growth assay. BAdvBSCs showed a greater ability to reduce the viability of intracellular Mtb than HAdvBSCs or vectors (Figures 4F and 4G). These data indicate that BAdvBSCs induces a robust autophagy-mediated antigen presentation to CD4 T cells in mouse DCs, mouse macrophages, and human macrophages.

Replication-defective BAdvBSCs vaccine induces expression of galectins and CTs to enhance antigen presentation in DCs

Previous studies show that autophagy plays a role during the translocation of Adv capsids from the endosome to the nuclear pore complex.35 During this process, galectins, specifically, Gal1 (Lgals-1) and Gal3 (Lgals-3), were found to be associated with the vesicular transport of Adv.36 However, HAdv5 infection of A549 epithelial cells strongly downregulated Gal3 (Lgals-3) expression.37 Transcriptomic studies show that Lgals-3 (Figure 2) and Lgals-8 (not shown; below the level of FPKMs indicated) transcripts were upregulated in BAdvBSCs-infected DCs. qPCR validation showed that BAdvBSCs induced increased mRNA transcripts for Lgals-3 and Lgals-8 relative to HAdvBSCs-infected DCs (Figure 5A). Interestingly, western blot analysis indicated that Lgals-3 was uniformly induced in DCs by both vectors and vaccines, while Lgals-8 was induced more strongly in BAdvBSCs-infected DCs (Figure 5B). qPCR and western blot analysis of BAdvBSCs- and HAdvBSCs-infected DCs also showed that the enrichment of uptake-related genes, including Gabarap (LC3 family), SQSTM1 (a known substrate of autophagy), and Rab7 (a small GTPase), are essential for the fusion of autophagosomes to lysosomes.58

Because Lgals-3 binds the tripartite containing motif protein-16 (TRIM16) to activate autophagy through ATG1/ULK1, we hypothesized that BAdvBSCs may induce autophagy through a Lgals-3- and/or Lgals-8-dependent mechanism.59–61 We reported that autophagy enhances the ability of APCs (macrophages; and DCs) to process and present a mycobacterial antigen to CD4 T cells.6,5 We performed small interfering RNA (siRNA) knockdown of galectins in APCs. BAdvBSCs- or HAdvBSCs-infected DCs or macrophages were subjected to siRNA versus Lgal-3 and Lgal-8 or siRNA versus Lgals-3, respectively. Galectin knockdown reduced antigen presentation by both BAdvBSCs- and HAdvBSCs-infected DCs or macrophages (Figures 5C and 5D). We note here that WT HAdv modulates autophagy through a Nedd4.2 pathway, reducing antigen presentation.48 We suggest that BAdvBSCs may augment autophagy and antigen presentation through the induction of Lgals-3; however, additional studies are required to determine whether Lgals-8 synergizes to activate autophagy.62

DCs degrade mycobacteria in their lysosomes using proteases such as CTs, lipases, and glycosidases. We demonstrated earlier that CTS cleaves Ag85B in mouse macrophages to generate the p25 epitope, which is then rapidly presented to CD4 T cells in vitro.3 CTSs not only digest proteins but also help to load peptides into the groove of MHC class II, and CTS and CTSL play a pivotal role.63,64 DC transcriptome studies indicated that BAdvBSCs enhanced the expression of multiple CTs (Figure 2F). Figure 5E shows that CTS inhibitors reduced antigen presentation in WT-DCs, but only the pan-inhibitor E64 had an inhibitory effect on BAdvBSCs- or HAdvBSCs-infected antigen presentation. All three CTS inhibitors reduced antigen presentation in WT-DCs, but only the pan-inhibitor E64 had an inhibitory effect on BAdvBSCs- or HAdvBSCs-infected Ag85KO-DCs (Figure 5F). To determine whether CTSs mediate the processing of virus-encoded antigens, WT-DCs and ATG7KO-DCs were pharmacologically blocked using pan-specific CTS inhibitors E64 and N-acetyl-L-leucyl-L-leucyl-l-methioninal (NALLM), and the CTS inhibitor calpeptin, followed by infection with BAdvBSCs or HAdvBSCs and antigen presentation. All three CTS inhibitors reduced antigen presentation in WT-DCs, but only the pan-inhibitor E64 had an inhibitory effect on BAdvBSCs- or HAdvBSCs-infected Ag85KO-DCs (Figure 5G). These data indicate that BAdvBSCs increases the immunogenicity of DCs through CTS-dependent antigen processing and presentation.

BAdvBSCs mucosal vaccine protects mice against aerosolized Mtb independently and as a BCG booster

to determine whether BAdvBSCs enhances immunogenicity in vivo, we intranasally inoculated age- and sex-matched inbred C57BL/8 mice, which show a “developing immune system” akin to the neonatal immune system,65 with vaccines or empty vectors. At 3 days post-inoculation, we euthanized the animals and analyzed bronchoalveolar lavage (BAL) cells and lung tissues using qPCR. BAdvBSCs markedly enhanced mRNA transcripts for a panel of genes associated with antigen processing and presentation compared to HAdvBSCs (Figures 6A and S4).

Figure 3. Replication-defective BAdvBSCs vaccine is rapidly internalized by mouse DCs

DC11+ DCs purified from bone marrow of WT-C57BL/6 mice were infected with the Cy3-labeled BAdvBSCs vaccine or control vector (10^7 PFU/10^6 DCs), followed by antibody staining for intracellular localization using confocal microscopy at 4 h post-infection. Cy3-BAdv vector or Cy3-BAdvBSCs vaccine-infected DCs were incubated for a 4-h infection, followed by fixation and staining using specific antibodies to (A) microtubule associate light chain 3 (LC3) autophagosome marker, (B) lysosome associated membrane protein-1 (LAMP1), (C) galectin3 (Lgals-3), and (D) Lgals-8 followed anti-immunoglobulin G (IgG) Alexa Fluor 488 and DAPI nuclear stain. Panels show Cy3 (red), Alexa Fluor 488 (green), and merged images with or without DAPI nuclear stain. Colocalization (inset) analyzed using MetaView software. Cytosolic (white arrow) and nuclear localization (yellow arrow) of Cy3-labeled vector or vaccines are indicated. Bar graphs to the right show percentage of virus-containing endosomes colocalizing with antibodies calculated per DC and averaged for 20 DCs in triplicates per experiments done twice. *p < 0.01 t test. Isotype stains are shown in Figure S5.
Figure 4. Replication-defective BAdv<sup>85</sup>C5 vaccine enhances autophagy-dependent and -independent antigen presentation in mouse DCs and human macrophages

(A) WT-C57BL/6- or ATG7 KO-C57BL/6-derived DCs were infected with vaccines or control vectors (10<sup>7</sup> plaque-forming units [PFUs]/10<sup>6</sup> DCs) followed by overlay using BB7 CD4 T cells specific for the Ag85B-p25 epitope in the context of MHC class II. IL-2 in supernatants at 18 h post-infection was determined using sandwich ELISA.

(B) Antigen presentation in vaccine- or vector-infected mouse macrophages (10<sup>7</sup> PFU/10<sup>6</sup> macrophages) is shown.

(C and D) Human peripheral blood mononuclear cell (PBMC)-derived CD14<sup>+</sup> macrophages were infected with vaccines or control vectors followed by an overlay with F9A6 T cells specific for an Ag85B epitope in the context of human leukocyte antigen-DR1 (HLA-DR1) for antigen presentation. Macrophages were untreated or treated with 3-methyladenine (50 μM) before infection to inhibit autophagy.

(E) Time-dependent antigen presentation by human macrophages infected with BAdv- or HAdv-derived vaccines. Horizontal line shows IL-2 levels of T cells incubated over naive macrophages.

(F and G) CD14<sup>+</sup> mouse bone marrow-derived macrophages were infected with 10<sup>6</sup> PFUs of vectors and vaccines indicated for 4 h, followed by co-infection with Mtb (Erdman strain; MOI = 1), washing, and incubation. Macrophage lysates were plated for CFU counts at indicated time points. For all of the panels, 1 of 2–3 separate experiments shown.

*p < 0.01; **p < 0.009 1-way ANOVA with Tukey’s test.
Figure 5. Replication-defective BAdv85C5 vaccine enhances galectin and cathepsin-dependent antigen presentation in mouse DCs and human macrophages.

CS7BL/6 mouse-derived DCs were infected with vaccines or control vectors, washed, and processed.

(A) At 18 h post-infection, lysates collected in Trizol were used for qPCR using primers (Method details) for endosome trafficking genes enriched during transcriptomics (Figure 2) (*R 1.4-fold increase in mRNA expression, n = 2, 2 experiments).

(B) At 18 h post-infection, lysates collected in anti-protease buffer analyzed using specific antibodies and Wes Simple blot protocol (Method details). Densitometry panels shown.

(C and D) DCs and macrophages were subjected to siRNA versus Lgals-3 and Lgals-8 or scrambled controls and at 18 h, infected using vaccines or vectors for 4 h. Washed DCs were overlaid using BB7 CD4 T cells, and 18 h later, supernatants assayed for IL-2 using sandwich ELISA (triplicate wells of DCs per group and 2 independent experiments).

(E and F) Lysates collected at 18 h post-infection were used for qPCR using primers (Method details) for cathepsins enriched during transcriptomics (Figure 2) (*R 1.4-fold increase in mRNA expression, n = 2, 2 experiments).

(G) WT-DCs or ATG7KO-DCs DCs were left untreated or treated with cathepsin inhibitors as indicated, followed by vaccine or vector infection and antigen presentation. NALLM, N-acetyl-L-leucyl-L-leucyl-L-methionyl; calpeptin; E64, each at 30 μM; triplicate wells of DCs per group and 2 independent experiments.

*p < 0.01, **p < 0.006, 1-way ANOVA with Tukey’s post-test.
We mock-inoculated or subcutaneously vaccinated mice with BCG and administered a single intranasal booster vaccination with BAdv, BAdvSC, or BAdv vector control 7 days post-immunization (Figure 6B). After 3 weeks, we administered an aerosolized Mtb Erdman challenge. At 4 weeks post-challenge, we euthanized the mice and collected lungs and spleen for Mtb bacterial counts and T cell profiling. Compared to vector, BAdvSC induced robust protection against Mtb as a booster following primary BCG vaccination, reducing lung Mtb burden by >1.4 log_{10} and spleen load by ~0.8 log_{10} (Figure 6C). BAdv generated a comparable decrease in Mtb colony-forming units (CFUs) in the lungs and spleen. However, only the autophagy-inducing BAdvSC dramatically reduced spleen CFUs (≥1.0 log_{10}) as an intranasal treatment alone. To confirm the specificity of the Ag85B-p25 immunogenic epitope, we stained lung- and spleen-derived T cells with an Ag85B-specific tetramer. Flow cytometry analysis showed that BAdvSC induced a robust expansion of Ag85B-p25-specific CD4 T cells in the lungs and spleen, consistent with reports that the Ag85B-p25 epitope protects mice against TB (Figure 6D).^{8,16} Finally, co-expression of the AIP-C5 peptide in BAdvSC led to a robust tetramer-positive CD4 T cell...
response in the lungs compared to the BAdv85 vaccine (Figure 6D).

We note here that BAdv85C5 induced a dramatic expansion of tetramer-positive CD4 T cells compared to mice vaccinated using recombinant BCG85C5 and Mtb-challenged mice in our previous study.5 This augmentation can be explained by the immunodominance of the p25 epitope and its ability to activate naïve T cells toward the Th1 pathway.67 Notably, unlike BCG85C5 given intradermally in our recent study, BAdv85C5 was given nasally, enabling a robust activation of lung APCs to induce tetramer-positive cells. Supporting this observation, co-expression of the AIP-CS peptide in BAdv85C5 led to a robust tetramer-positive CD4 T cell response in the lungs compared to BAdv85 vaccine (Figure 6D). These data suggest that BAdv85C5 expressing one major antigenic epitope of Mtb (Ag85B) can serve as a robust mucosal vaccine, even when administered as a single dose.

A BAdv85C5 booster induces cytokine-positive T cells, TEM cells, and TCM cells in BCG-vaccinated mice

Because Tn1 immunity-mediating T cell-secreted cytokines such as interferon-γ (IFN-γ) and interleukin-2 (IL-2) play a major defensive role against TB in mice and humans,58 we investigated whether BAdv85, BAdv85C5, or BAdv vectors affected these cytokines when administered as a single dose or as a booster. BAdv85C5 vaccine significantly enhanced IFN-γ and IL-2-expressing CD4 and CD8 T cells compared to either BAdv85 or BAdv vector (Figures 7A–7D). Importantly, BAdv85C5 induced an expansion of IFN-γ+ and IL-2+ T cells in lung and spleen when given as an intranasal vaccine, suggesting that irrespective of BCG vaccination, it can elicit immune responses and adults can also be nasally vaccinated against TB.

Respiratory infections induce a TEM cell response, which enables an immediate containment of infection, followed by their transition into a long-lasting TCM cell response. In many viral infection models and LCMV (lymphocytic choriomeningitis virus) infection of mice, TCM cells mediate long-term immunity.68–71 Since the lungs are not lymphoid organs, TEM and TCM are thought to arise in the lymph nodes and home back to the lungs for the containment of infection recruited by the chemokines secreted from the Mtb-infected macrophages. For example, chemokine receptors CCR2, CCR5, CCR6, and CXCR6 enhance lung infiltration with Tn1-type immune cells, and CCR2-deficient mice are highly susceptible to TB.72,73 Figures 7E–7H demonstrate that both BAdv85 and BAdv85C5 induced an expansion of TEM cells in the lungs and spleens of the BCG-vaccinated mice, although only BAdv85C5 induced a significant TCM response. BCG induces a robust TEM response, but a reduced TCM response in mice.74 Therefore, we propose that using BAdv85C5 as a booster in BCG-vaccinated mice can augment long-lived memory T cells.

**DISCUSSION**

This study presents an alternative BAdv vaccine platform approach to HAdv-based TB vaccines, which have shown promise in mouse and macaque models,13,19 but not in human studies.50 We have previously demonstrated that BAdv vector-based vaccines are effective in the presence of exceptionally high levels of preexisting HAdv vector immunity24; thus, preexisting HAdv vector immunity should not affect vaccine efficacy. Furthermore, BAdv3 internalization is independent of HAdv5 receptors (Coxsackievirus-Adv receptor [CAR] and αvβ3 or αvβ5 integrin),78 and instead uses α(2,3)-linked as well as α(2,6)-linked sialic acid-containing proteins as major receptors.79 Preexisting HAdv-neutralizing antibodies in humans do not cross-neutralize BAdv3,80 and HAdv-specific immune response does not affect BAdv.81 Notably, unlike HAdv5, BAdv3 is a strong inducer of TLR4 and does not deplete Kupffer cells of the liver.82 Our previous biodistribution study using a BAdv3 vector in mice showed that BAdv3 efficiently transduces the heart, kidney, lung, liver, and spleen, and persists longer than HAdv5, particularly in the heart, kidney, and lung.82 Sequential administration of HAdv5 and BAdv3 vectors also overcomes vector immunity in an immunocompetent mouse model of breast cancer,83 and BAdv3 and HAdv5 vector genomes have shown similar persistence in human and nonhuman cell lines.82 Thus, BAdv3 vectors are an attractive alternative to HAdv vectors to effectively immunize individuals with high levels of preexisting HAdv immunity. Because sialic acid is the primary receptor for BAdv, this vector-based vaccine platform should be ideal for intranasal immunization.36 Here, we demonstrate that the BAdv nasal vaccine platform expressing the Mtb Ag85B-p25 epitope in combination with
AIP-C5 generates significant protection against Mtb in mice. We have previously shown that APC processing and presentation of mycobacterial antigens is critical for vaccine-mediated anti-TB immunity.5,6,8,12 While BCG and WT-Mtb evade phagolysosome fusion, BCG overexpressing Ag85B (BCG85B), Ag85B plus AIP-C5 (BCG85B:CS), or sapM/tbpA gene knockouts of Mtb (ΔbtpA-ΔsapM-Mtb) are delivered to mouse APC lysosomes to generate peptides that activate CD4 T cells ex vivo.14,15 Both BCG85B and BCG85BS are induced robust activation of Ag85-p25 epitope-specific CD4 T cells in mice after vaccination and Mtb challenge.5,8 However, a high risk of lung inflammation precludes intranasal administration of recombinant BCG vaccines in infants.

Because DCs play a pivotal role during vaccination, we evaluated gene expression in BAdv85C5-infected DCs in comparison with HAdv85C5. Interestingly, BAdv85C5 induced significant upregulation of multiple genes involved in the sorting of vaccine-containing endosomes to lysosomes and genes regulating antigen processing (Figures 2F and 2G). These genes included the galectins Lgals-3 and Lgals-8, which are associated with the vesicular transport of Adv from the endosome to the nuclear pore complex during autophagy;16 the endosome trafficking proteins Gabarap, Rab7, LAMP1, LAMP2, CD63, and CD53; and the antigen-presentation molecules B2m, H2-D1, and CTSs. These findings indicate that BAdv85C5 enhances the ability of DCs to present the antigen to T cells, a key event during the expansion of Th1 cell responses.

Based on these findings, we further investigated the mechanism(s) underlying BAdv85BSCS vaccine-induced immunogenicity. Previous studies have shown that Lgals-8 plays a role during WT-HAdv modulation of autophagy, reducing antigen presentation in human epithelial cells.46 Galectins participate at various levels of autophagy; Lgals-8 inhibits mammalian target of rapamycin (mTOR)-inducing autophagy;47 Lgals-3 regulates lysosome stability through lysosome-associated membrane protein (LAMP)-1,48 and WT HAdv modulates autophagy through a Nedd4.2 pathway, reducing antigen presentation.49 BAdv85C5 may augment autophagy and antigen presentation through the induction of both Lgals-3 and Lgals-8; however, additional studies are required to determine whether Lgals-8 synergizes to activate autophagy. Our initial studies using BAdv85C5 and HAdv85C5 revealed reduced antigen presentation to CD4 T cells when ATG7KO-DCs were used (Figure 4A). Inhibition of autophagy in human macrophages also reduced antigen presentation (Figure 4D). Although HAdv85-infected DC antigen presentation was demonstrable, the addition of AIP-C5 in HAdv85C5 significantly increased antigen presentation. Thus, co-expression of AIP-C5 with the Ag85B-p25 epitope not only bypassed the ability of HAdv to interfere with autophagy but it also enhanced the antigen presentation ability of BAdv85C5-infected DCs. BAdv85C5 induced robust Lgals-3 and Lgals-8 expression compared to HAdv85C5 (Figures 5A and 5B). Because siRNA blockade led to the downregulation of antigen presentation, BAdv85C5 induction of Lgals-3 and Lgals-8 ostensibly facilitates antigen presentation (Figures 5C and 5D).

Consistent with our previous observation that the induction of autophagy enhances MHC class II-dependent antigen presentation by mycobacteria-infected DCs,5 Adv85BSCS-infected DCs upregulated several genes, which likely facilitated autophagy-dependent antigen presentation. Different genes are induced for specific types of autophagy; LAMP2 participates during chaperone-mediated autophagy when soluble proteins of cytosol are internalized into lysosomes,86,87 and Gabarap (LC3 family) and Rab7 are involved during selective autophagy88 and autophagolysosome fusion, respectively. Because multiple immunoregulatory genes were expressed at lower levels following the infection of ATG7KO-DCs with BAdv85C5 compared to WT-DCs (Figure S3), the signaling pathways that deliver BAdv85C5 into autophagolysosomes should be investigated further. Our findings define a CTS-dependent mechanism through which BAdv85C5-infected DCs demonstrate increased immunogenicity relative to HAdv85C5.

BAdv85C5 enhanced CTS gene expression and protein synthesis compared to HAdv85C5 (Figures S5E and S5F). CTSs play a major role during antigenic peptide production due to the acidic pH of the lysosomes. For example, CTSD cleaves mycobacterial Ag85B to produce the p25 epitope, which is then presented to CD4 T cells.3 CTSB, CTSS, and CTSL play a pivotal role in digesting antigens and loading microbial peptides into the groove of MHC class II.64 Our pharmacological blockade of CTSs led to a near-abrogation of antigen presentation by BAdv85C5 and HAdv85C5 in infected DCs, validating the observed gene expression profiles (Figure 5G). Earlier studies have indicated a potential bactericidal effect of CTSs. CTSG induces the degradation of Mtb within THP-1 macrophages, and CTSB, CTSS, or CTSL blockade enhances Mtb growth in human macrophages.89,90 Both BAdv85 and HAdv85C5 exerted a bactericidal effect against Mtb in mouse macrophages, suggesting that perhaps CTSs mediate this effect (Figures 4F and 4G).

These data indicate that relative to HAdv85C5, BAdv85C5 more robustly enhances the lysosomal production of the Ag85B-p25 epitope to boost DC immunogenicity. mTOR is a major negative regulator of autophagy that controls the accumulation of DCs in the lungs and governs the programming of their metabolomes.91 Because BAdv85C5 induced an upregulation of genes in DCs mainly in autophagy pathways (Figures 2, 3, 4, and 5), we propose that BAdv85C5 may confer superior antigen-mediated activation of T cells to lung DC populations, particularly when administered intranasally. Notably, the modified vaccinia Ankara (MVA) vaccine induces apoptosis of DCs in draining lymph nodes,92 which is a possible reason for the failure of MVA85A to protect against TB in children. In our studies, BAdv85C5 was rapidly internalized into DCs (Figure 3), facilitating sustained and robust antigen presentation, with DCs remaining fully viable during in vitro assays (Figure 4E).

Consistent with the in vitro immunogenicity data, BAdv85C5 showed a marked protective effect in mice exposed to Mtb and induced a marked expansion of tetramer-positive CD4 T cells. When given as a single nasal vaccine, BAdv85C5 induced robust gene expression in the lungs and BAL cells of mice compared to HAdv85C5 (Figure 6A). When given as a booster following BCG, the vaccine also induced a ~1.4-log reduction of Mtb burden in the lungs compared to BCG alone (Figures 6B and 6C). For a single dose of a single epitope with a nasal booster vaccine, this appears to be the best level of protection in mice reported for any Adv vaccine for tuberculosis. Furthermore, because a single dose of a nasal vaccine alone reduced Mtb burden in both the lungs...
and spleen, this vaccine strategy may also protect adults at risk for TB.

Immunity against TB is multifactorial, including Th1-immunity facilitated by cytokine-secreting CD4 and CD8 T cells, a variety of other innate T cells, neutrophils, myeloid cells, and antibodies. Figure 7 illustrates that the BAdv85C5 vaccine induced an expansion of IFN-γ- and IL-2-secreting CD4 and CD8 T cells in both BCG-vaccinated mice and those receiving only the BAdv85C5 vaccine. Most TB vaccines induce memory T cells, which can be classified into an early TCM response followed by a longer-lasting TEM response. Interestingly, BCG vaccine is a poor inducer of TCM response in mice, although recombinant BCG vaccines induce a stronger TCM response in mice correlating with superior protection against TB.8,34 Although the role of TCM response in protecting against primary exposure to Mtb is debated, it appears that TCM cells play a major role in defending against reinfection and reactivation of Mtb. For example, HIV-induced CD4 T cell depletion predisposes Mtbc to reactivation, implying that at least CD4 TCM cells persist in a resting stage in those vaccinated with BCG or exposed to Mtb. These resting TCM cells can rapidly expand into TEM cells to contain Mtb after exposure. In our investigation, BAdv85C5 vaccination led to an expansion of both TEM and TCM cells in BCG-vaccinated mice (Figure 7). Curiously, BAdv85C5 given alone expanded TCM cells in the spleen (Figure 7), indicating that the BAdv85C5 vaccine can be used even among adults to activate TCM cells. This is significant because, at least in the mouse model, BCG seems to be a poor inducer of TCM response.74 Intranasal BCG vaccination of neonates followed by a nasal booster with BAdv85C5 may stimulate longer-lasting immunity, while adults exposed to index cases of TB may be protected by the BAdv85C5 vaccine alone.

BCG revaccination of healthy volunteers reduces skin test conversion, but whether it affects TB susceptibility remains unclear. Intranasal BCG is not feasible, but we propose that a BAdv85C5 nasal vaccine can strengthen the lung compartment to resist aerosol infection with Mtb. Supporting this concept, the BAdv85C5 booster enhanced TEM responses in the lungs of BCG-vaccinated mice before and after Mtb exposure (Figure S7). This is a clinically significant finding, as human and mouse neonates both show a reduced expansion of TEM cells after influenza infection.21 Others have suggested that during RSV vaccination, TLR activation may help in overcoming the hypo-responsiveness of TEM cells in the lungs.34 In this context, we recall that AIP-C5 activates TLR-2 in both mouse and human APCs, inducing autophagy.8 Although additional studies are required, we consider that the TLR2-dependent adjuvant effect of AIP-C5 may contribute to the robust TEM responses in the lungs following BAdv85C5 vaccination, which will likely augment neonatal lung immunity when given as a nasal vaccine.

In summary, the BAdv vaccine platform enables the expression of an Mtb immunogenic peptide and activates Lgals-3 and Lgals-8-dependent autophagy. This platform led to robust antigen presentation by both mouse DCs and human macrophages, resulting in excellent protection against TB. Autophagy-mediated antigen presentation through a BAdv vaccine platform expressing multivalent immunogenic proteins of Mtb may facilitate future designs of more potent mucosal TB vaccines for children and adults.

Limitations of the study
We have demonstrated the efficacy of BAdv85C5 vaccine using 4- to 6-week-old C57BL/6 mice that have an immune system resembling that of human infants. However, a neonatal mouse model would be ideal for an additional evaluation of the vaccine, given that BCG is provided after birth. We also need to test nasal booster vaccine months after BCG vaccine using neonatal mice and evaluate gene expression studies post-TB challenges. The potential of the full 85B antigen with or without other immunogenic antigens of Mtb in the BAdv vaccine platform should be explored.

STAR METHODS
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SUPPLEMENTAL INFORMATION
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AUTHOR CONTRIBUTIONS
A.K., E.E.S., V.K.S., and A.M. performed the experiments; S.D.-E., S.N., and K.J.S. analyzed the flow cytometry data; D.H.C., M.C., and J.W. provided the research materials; C.J. and S.K.M. designed the experiments and wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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## STAR★METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| CD3                 | BD Biosciences | 564380     |
| CD62L               | BD Biosciences | 553150     |
| CD8                 | BD Biosciences | 551162     |
| CD44                | BD Biosciences | 562464     |
| CD4                 | BD Biosciences | 561025     |
| CCR7                | eBioscience   | 12-1971-80 |
| IFN-γ               | BD Biosciences | 564336     |
| IL-2                | BD Biosciences | 560547     |
| CD103               | BD Biosciences | 562772     |
| CD69                | BD Biosciences | 740220     |
| CD11a               | BD Biosciences | 740849     |
| Rab7                | Cell Signaling | 9367T      |
| LC3B                | Cell Signaling | 2775s      |
| GAPDH               | Cell Signaling | 5174s      |
| Gbp1                | Novus Biologicals | NBP-1-31560 |
| **Bacterial and virus strains** |        |            |
| Mycobacterium tuberculosis Erdman | ATCC | ATCC 25177 |
| Bovine adenovirus   | ATCC | ATCC VR-639 |
| Human Adenovirus    | ATCC | ATCC VR-5  |
| Mycobacterium bovis BCG Pasteur | ATCC | ATCC 35732 |
| **Biological samples** |        |            |
| Peripheral blood from Human subjects | Gulf coast regional Blood Center, Houston | NA |
| **Chemicals, peptides, and recombinant proteins** |        |            |
| IMDM Media          | GE Healthcare HyClone | SH30228.01 |
| FBS                 | GE Healthcare HyClone | SH30070.01 |
| Penicillin and gentamycin | GibCO | 15140148 |
| HEPES               | GibCO | 15630080 |
| 2-Mercapto ethanol  | GibCO | 21885023 |
| h-GM-CSF            | BioLegend | 572904 |
| m-GM-CSF            | BioLegend | 576306 |
| Histopaque 1077     | Sigma | 1077500ML |
| ACK buffer          | Lonza | 10548E |
| RPMI 1640 Medium    | Sigma | R8758 |
| RIPA buffer         | Sigma | R0278 |
| SDS                 | Sigma | L3771 |
| Collagenase         | Thermo Fisher Scientific | 17101015 |
| Human AB serum      | Fisher | BP2525100 |
| Kanamycin           | Fisher | BP906-5 |
| Lysotracker red     | Invitrogen | L7528 |
| Lysotracker green   | Invitrogen | L7526 |
| PMA                 | Sigma | P1585 |
| Ionomycin           | Sigma | I0634 |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Chinnaswamy Jagannath (cjagannath@houstonmethodist.org).

Materials availability
Viral strains generated in this study will be made available upon request. A provisional patent application for BAdv85C5 has been filed by Purdue University and HMRI. Vaccines related to this study will be made available on request, but we may require a payment and/or a completed Materials Transfer Agreement if there is a potential for commercial application.

Data and code availability
The RNaseq dataset supporting the current study are available in NCBI database: NCBI Bioproject #PRJNA745965 (SRR15107864; SRR15107868; SRR15107866; SRR15107869; SRR15107865; SRR15103113; SRR15103107; SRR15103114; SRR15103108; SRR15103112; SRR15103117; SRR15103116; SRR15103106; SRR15103109; SRR15103111; SRR15103115; SRR15103110). https://www.ncbi.nlm.nih.gov/sra/PRJNA745957

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice
The age- and sex-matched (4-6 weeks old; M/F) C57BL/6 mice were purchased from Jackson laboratory, Bar Harbor, ME, USA, and housed under specific pathogen-free conditions in the animal facility of the University of Texas Health Sciences Center, Houston, where the experiments on mice were performed before the relocation of Dr. Jagannath to HMRI. Mouse in ABSL3 vivarium maintenance was in accordance with ethical and animal welfare committee protocols complying with the current PHS policy of USA.
Ethics statement
All animal experiments in this study were performed as per animal protocols verified and approved by the Institutional Biosafety Committee of University of Texas Health Science Center, Houston (where part of this study was performed). The animal study was carried out in accordance with the US government PHS policy on Humane Care and Use of Laboratory Animals and the protocol for the same was approved by the UTHSC Animal Welfare Committee of under protocol number # AWC – 16-0136. HMRI approval number for similar protocols is AUP-0620-0037.

Cell lines
The CD4 T cell hybridoma BB7 specific for the p25-epitope of Mtbg85B was a gift of Dr. Clifford V. Harding, Case Western Reserve University, OH. The F9A6 cell line is specific for a human epitope of Ag85B and donated by Dr. DH. Canaday, coauthor. Both were maintained as frozen stocks at –80°C and thawed cells culture in DMEM medium with β-ME.

Primary cells
CD14 beads (Miltenyi Biotec, USA) were used to purify primary human and mouse macrophages, which were then cultured in GM-CSF containing medium as described under methods.

METHOD DETAILS

Isolation and cultivation of primary mouse macrophages and DCs (APCs)
The bone marrows from 4-6 week-old C57BL/6 mice (M/F) (Jackson research Labs, USA) were obtained by flushing both the femur and tibia. The pooled bone marrows from 2 mice at a time was centrifuged for five minutes at 1,000 rpm. The cells were washed two times in ACK lysing buffer (Fisher BW10-548E), followed by 1 mL of phosphate buffered saline (PBS), passed through a 36-gauge needle, and suspended in Iscove’s Modified Dulbecco’s Medium (IMDM) with 10% percent fetal bovine serum (FBS), 20 ng per milliliter granulocyte macrophage colony stimulating factor (GM-CSF) and antibiotics (penicillin and gentamicin). The cells were then plated into six-well plates and incubated at 37°C in a CO2 incubator for one week, while spiking the wells with 2 mL of fresh medium every two days. After seven days, the wells are flushed and the pooled were transferred to a 50 mL tube. The cells were pelleted and resuspended in MACS buffer (PBS containing 0.5% FBS) at a volume of 400 μL per 10^8 cells. CD11c microbeads (Miltenyi 130-052-001) were then added to the cells, 100 μL per 10^8 cells. The cells were incubated at 4°C for 15 minutes, washed once in 1 mL of MACS buffer and resuspended into 500 μL of MACS buffer. During this time, the Miltenyi separation columns (Miltenyi 130-042-201) were prepared by rinsing once with 500 μL MACS buffer, one column per 10^8 cells and placed onto a magnetic holder. The cells are added to the columns and the pass through was collected. The macrophages and other cells pass through since they lack CD11c on their surface. The columns were washed three times with 500 μL MACS buffer and then removed from the magnetic holders. One mL of dissociation buffer was then plunged through the column, detaching the dendritic cells off the column. Wash off was repeated once. The flow-through macrophages were then run through another CD14 column repeating the procedure to purify macrophages. The cells (DCs and macrophages) were counted and plated into 24-well plates with 10^6 cells per well in IMDM supplemented with 10% FBS and GM-CSF, and allowed to differentiate for 4-7 days. The medium was replaced with a GM-CSF-free medium rested and cells were used for various assays.

APCs were used to conduct mycobacterial infection, antigen presentation assay, and confocal imaging studies. They were used as monolayers in 8-well slide cultures for antibody stains and in 24-well plates for antigen presentation assays (IL-2 assays) or Mtbg CFU assays.

Bacterial strains and culture conditions
Log phase organisms of wild-type M. tuberculosis (Erdenman strain) and BCG vaccine (ATCC; Pasteur strain) cultured in 7H9 broth for 7 days were frozen in aliquots. Before use, aliquots were thawed, washed three times in PBS (12,000 rpm; 15 mins), sonicated at 4 W using a sonication probe and dispersed suspension matched with McFarland #1 in turbidity (10^6 CFU/mL). A single cell colony forming unit (CFU) suspension of Mtbg/BCG were made for infections by gentle centrifugation at 500 rpm for 5 min and the supernatants containing well dispersed organisms were used 10^7 CFU/mL.

Adenoviral vectors and culture conditions
BHH3 (bovine-human hybrid clone 3), BHH2C (bovine-human hybrid clone 2C), 293 (human embryonic kidney cells expressing HAdV E1 proteins), and 293Cre (293 cells expressing Cre recombinase) were propagated as monolayer cultures in minimum essential medium (MEM) (Life Technologies, Thermo Fisher Scientific, Waltham, MA) enriched with either 10% reconstituted fetal bovine serum or fetal calf serum (HyClone, Thermo Fisher Scientific) and gentamycin (50 μg/mL). The production and characterization of BAdv (BAdv-3 E1 and E3 deleted empty vector), and HAdv (HAdv-5 E1 and E3 deleted empty vector) have been described previously. Gene constructs containing the immunogenic epitope Ag85B-p25 of Mtbg without (85) or with AIP-C5 (85C5) were codon-optimized for rodent expression and synthesized commercially (GenScript Biotech Corporation, Piscataway, NJ). We adapted the technology of homologous recombination in bacteria for the generation of BAdv vectors expressing the Ag85B-p25 epitope of Mtbg without (BAdv85) or with AIP-C5 (BAdv85C5) (Figure 1). For developing HAdv vectors
expressing the Ag85B-p25 epitope of Mtb without (HAdv85) or with AIP-C5 (HAdv85C5) (Figure 1), we applied the technology of Cre recombinase-mediated homologous recombination. The gene cassette in BAdv or HAdv vectors was under the control of the cytomegalovirus (CMV) immediate early promoter and the bovine growth hormone (BGH) polyadenylation signal. The presence of the foreign gene cassette in the vector was identified initially by restriction analysis followed by sequencing the region containing the gene cassette. The expression of the immunogenic epitope/s in vector-infected cells was confirmed by RT-PCR. BAdv, BAdv85 and BAdv85C5 vectors were grown and titrated in BH3 cells as described. Whereas HAdv, HAdv85 and HAdv85C5 vectors were replicated in 293 cells and titrated in BH2C cells as described elsewhere. All vectors were purified by cesium chloride density-gradient ultracentrifugation as described.

Isolation and cultivation of primary human macrophages from peripheral blood samples
After obtaining informed consent, peripheral blood samples from healthy HLA-DR1 positive donors were obtained from Gulf Coast Regional Blood center. CD14 magnetic beads (Miltenyi, USA) were used to purify monocytes which were plated in 24 tissue culture plate wells for all experiments at 1x10^6 cells per well. CD14 bead purified monocytes were grown in Iscove’s medium (IMDM) with 10% FBS and 10 ng/mL GM-CSF for 7 days and then plated in GM-CSF free medium for 24 hr before assay.

Infection of APCs with Adv vectors and vaccines
APCs were infected with BAdv or HAdv vectors (10^7 PFU per million APCs) followed by 4 hr incubation and washing three times with 10% FBS and 10 ng/mL GM-CSF for 7 days and then plated in GM-CSF free medium for 24 hr before assay.

RNaseq analysis
Naive or BAdv/HAdv vector-infected MΦs or dendritic cells (2 X 10^6 cells per pellet; n = 2) were collected and mixed with Trizol buffer and snap frozen in liquid nitrogen. RNAsseq analysis, data interpretation and pathway analysis were done by Novogene (USA). Additional bioinformatic analysis was done in-house. Targeted genes were validated using RT-PCR using probes as described below.

qPCR assay for gene expression in mouse bone marrow derived dendritic cells
Total RNA was extracted using RNAeasy mini kit (QIAGEN, Germany) from mouse bone marrow derived dendritic cells (DCs). RNA concentration and purity ratios (OD260/280, OD260/230) were measured using the NanoDrop ND-1000 spectrophotometer (Thermo Fisher Scientific, USA). cDNA synthesis was performed on a CFX96 Real-Time PCR System (Bio-Rad, USA) using the 2X OneStep qRT-PCR Mastermix Kit (Applied Biosystems, USA) according to manufacturer’s instruction. Quantitative PCR (qPCR) was performed using SYBR green probe and gene specific primers (Table-primers). Threshold cycle numbers were transformed to ΔΔCt values, and the results were expressed relative to the reference gene, β-actin and GAPDH. Gene expression data was performed using GraphPad Prism ver. 6.0 suite (GraphPad Software). Student’s t test was used for means comparison between control or vaccine infected DCs. Significance was set at the 0.05 level.

Bronchoalveolar macrophages were aspirated from mice and pelleted in trizol for qPCR; Lungs were dissected, and equal amounts of fragments suspended in trizol. Primers for both DCs and mouse tissues are described in Figure S8.

Ex vivo Mtb Ag85B antigen presentation to CD4 T cells
The Ag85B epitope (241-256)-specific T cell hybridoma (BB7) was a kind gift from Drs. Cliff Harding (Case Western Reserve University, OH). The ex vivo antigen presentation assay has been described in detail by us, and the original method described by the Harding lab has been extensively used by us and others for in vitro antigen presentation by APCs (references cited in main text). Briefly, APCs infected with vaccines and vectors were washed 4 hr post-infection and overlaid with the BB7 CD4 T cell hybridoma which recognizes an Ag85B-p25 epitope in the context of mouse MHC-II (H2-d). IL-2 secreted from hybridoma T using a sandwich ELISA kit (eBiosciences). For human macrophages, the F9A6 cell line was used, which detects another Ag85B epitope in the context of human HLA-DR1.

siRNA knockdown of BAdv/HAdv vectors-infected APCs
The kits for various siRNAs (mixture of duplexes for Gal3, Gal8), were purchased from Origene as listed in the Key resources table. APCs were treated with siRNA and the scrambled control according to the manufacturers’ instructions, and this was followed by an overlay with BB7 CD4 T cells and evaluation of antigen presentation.

Evaluation of virus containing endosomes and their colocalization with autophagolysosomes using antibodies and Cy3 labeled BAdv vector and vaccine in DCs
To visualize BAdv containing endosomes within DCs, BAdv vector and BAdv85C5 were labeled through covalent linking of the fluorescent dye, FluoroLink Cy3 (Amersham, Inc.), essentially as described by Leopold et al. Briefly, BAdv stocks were adjusted to a concentration of 10^10 particles/mL and then mixed in a 1:9 ratio with Cy3 previously reconstituted in 0.1 M sodium carbonate, pH 9.3. After 30 min, the reaction mixture was transferred to a dialysis chamber (6,000-8,000 MW cutoff, Thermo Fisher) and dialyzed at 4°C for 24 hr against two changes of 10% glycerol, 50 mM Tris-HCl pH 7.5, 250 mM MgCl2. Finally, the dialyzed mixture was then brought to 30% glycerol and stored at −20°C until use. For microscopy, mouse DCs were cultured on tissue culture chambered
slides or coverslips. Primary DCs were seeded at 10^4 cells per chamber and incubated at 37°C in a humidified incubator with 5% CO2, and used 24 hr after plating. DCs were infected using Cy3 labeled BAdv vector and BAdv\textsuperscript{BCS5} at 10^6 PFU per chamber for 4 hr. DCs were washed, fixed with 4% paraformaldehyde, permeabilized with a perm buffer (PBS with 0.05% tween 80, 0.01% SDS and 0.05% BSA) and labeled with LC3, LAMP1, Lgals-3 or Lgals-8 specific primary antibodies with an IgG isotype control overnight at 4°C. Alexa Fluor 488 (Thermo Fisher) tagged anti-IgG was used as secondary conjugate. Fluorescence microscopy was performed with an inverted Nikon N90 microscope equipped with 60-NA and a MetaView software which enables 3D-deconvolution of fluorescent images. Virus containing endosomes were determined counting individual puncta ≥ 200 nm labeling either red (C3 only) or labeling yellow after colocalization with specific antibodies. Noncolocalizing (red) or colocalizing (yellow) endosome puncta (Figure 3) were counted in individual DCs were averaged and repeated for 20 different DCs for duplicate chambers. Data were expressed as percent endosomes labeling for antibodies and analyzed for significance using t test and Graphpad prism software. To avoid visual bias, pixel based colocalization was used.

**Mouse vaccine experiments**

The main mouse vaccine validation protocol is identical to the NIH mouse model that has been described by us previously.\textsuperscript{8} Age and sex matched mice (4-6 weeks) were tested naive or vaccinated with BCG via s.c. route to the hind legs of mice (confirmed as 1 × 10^6 CFUs through plating on 7H11 agar). Intranasal vaccination with BAd and HAd viral vectors were then done after 7 days. Thirty days after prime boost vaccination, mice were aerosol challenged with ~100 CFU of virulent Mtb Erdman using a Glas-Col (Indiana, USA) aerosol apparatus. Four weeks (day 60) after challenge or at indicated times, organs were harvested for CFUs and cytometry. Significance of differences in the CFU counts was calculated using 2-way ANOVA. 5 mice per vaccine strain were used and three-four individual mice per time point for flow cytometry analysis. Two independent experiments were carried out, each with 5 mice per group and Figure 6C shows the combined CFU data. Challenge Mtb organisms were differentiated from BCG vaccine by culturing organ homogenates in 7H11 agar containing Thiophene-2-carboxylic acid hydrazide (10 μg/mL), which inhibits BCG but not wild-type Mtb.

**Post Mtb challenge flow cytometry for TRMs**

Mice were sacrificed on day 60 post Mtb challenge to determine the numbers of lung resident TRMs. 5 mice per group were individually analyzed as follows. a) Lungs perfused with sterile phosphate-buffered saline were teased in buffer with 1 mg/mL collagenase and 1 mg/mL elastase (Sigma Biologicals, USA) to break down the fibrous tissue material. Single-cell suspensions from lungs and red cell lysed spleens were prepared in PBS with 10% FCS and 2 mM EDTA (Sigma-Aldrich) by straining through 40-μm cell strainer using a plunger of 5-mL syringe (BD Biosciences). Lung- and spleen-derived cells were stained for CD4 and CD8 T cells using antibodies to surface markers and intracellular cytokines as described before.\textsuperscript{8}

**Flow cytometry analysis**

Results were reported as absolute numbers of T cells per organs after performing an initial organ cell count using trypan blue and staining and acquiring a fixed number of cells. One million splenocytes or lung cells were stained using antibodies to CD3 (564380), CD4 (561025), CD62L (553150), CD44 (562464), CCR7 (12-1971-80), CD103 (562772), CD69 (740220), and 1 mg/mL elastase (Sigma Biologicals, USA) to break down the fibrous tissue material. Single-cell suspensions from lungs and red cell lysed spleens were prepared in PBS with 10% FCS and 2 mM EDTA (Sigma-Aldrich) by straining through 40-μm cell strainer using a plunger of 5-mL syringe (BD Biosciences). Lung- and spleen-derived cells were stained for CD4 and CD8 T cells using antibodies to surface markers and intracellular cytokines as described before.\textsuperscript{8}

**Post vaccination and post Mtb challenge flow cytometry for TRMs**

Mice were sacrificed on day 21 post vaccination and day 60 post Mtb challenge to determine the numbers of lung resident TRMs using CD103 and CD69 markers. 3 mice per group were individually analyzed.

**Western blots: ‘Wes Protein Simple’ western assay / Wes analysis of proteins in lysates of DCs**

For the analysis of protein levels in DCs, the quantitative Wes capillary immunoassay was used, in which the lysates were separated and detected using Wes separation capillary cartridge 12-230 kDa along with Wes Anti-Rabbit Detection Module (Simple Western system and Compass Software, Protein Simple). In brief, glass microcapillaries were loaded with stacking and separation matrices followed by sample loading. During capillary electrophoresis, proteins were separated by size and then immobilized to the capillary wall. Samples were loaded at 1 mg/mL dilution and the primary rabbit antibodies and GAPDH were used at 1:50 dilution. Data were analyzed with the Compass software (version 2.6.7). The area under the curve (AUC), which represents the signal intensity of the chemiluminescent reaction was analyzed for all the antibodies and β-actin. Values given for protein expression were normalized to β-actin. Quantitation of protein levels (area under each peak; arbitrary units [A.U.]) were performed using the Compass software.
All the primary antibodies used for the Simple Western are listed in the Reagents Table. All the antibodies were purchased from Cell signaling or Novus biologicals.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

All experiments were performed at least 2-3 times at different times. *Ex vivo* assays contained triplicate wells per group or combination. Statistical parameters including the definition of central value and spread (mean ± SD) and the exact number (n) of DCs or macrophages or mice per group are annotated in the corresponding figure legends. Statistical analysis was performed with GraphPad Prism version 8.04 for Windows (GraphPad) by using, unless otherwise stated, unpaired t tests with 95% confidence intervals. For statistical analysis of the microscopic quantification and intracellular survival of *M. tuberculosis*, the one-way ANOVA with Tukey’s multiple comparisons test was applied. (* p value < 0.05; ** p value < 0.01; *** p value < 0.001; ns, not statistically significant). All CFU calculations in mouse experiments were done using 2-way ANOVA using Dunnett's posttest.