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MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

Kashangura R, Jullien S, Garner P, Johnson S

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[Intervention Review]

MVA85A vaccine to enhance BCG for preventing tuberculosis

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ABSTRACT

Background

Tuberculosis causes more deaths than any other infectious disease globally. Bacillus Calmette-Guérin (BCG) is the only available vaccine, but protection is incomplete and variable. The modified Vaccinia Ankara virus expressing antigen 85A (MVA85A) is a viral vector vaccine produced to prevent tuberculosis.

Objectives

To assess and summarize the effects of the MVA85A vaccine boosting BCG in humans.

Search methods

We searched the Cochrane Infectious Diseases Group Specialized Register; Central Register of Controlled Trials (CENTRAL); MEDLINE (PubMed); Embase (Ovid); and four other databases. We searched the WHO ICTRP and ClinicalTrials.gov. All searches were run up to 10 May 2018.

Selection criteria

We evaluated randomized controlled trials of MVA85A vaccine given with BCG in people regardless of age or HIV status.

Data collection and analysis

Two review authors independently assessed the eligibility and risk of bias of trials, and extracted and analyzed data. The primary outcome was active tuberculosis disease. We summarized dichotomous outcomes using risk ratios (RR) and risk differences (RD), with 95% confidence intervals (CI). Where appropriate, we combined data in meta-analyses. Where meta-analysis was inappropriate, we summarized results narratively.

Main results

The search identified six studies relating to four Phase 2 randomized controlled trials enrolling 3838 participants. Funding was by government bodies, charities, and philanthropic donors. Five studies included infants, one of them infants born to HIV-positive mothers. One study included adults living with HIV. All trials included authors from Oxford University who led the laboratory development of the vaccine. Participants received intradermal MVA85A after BCG in some studies, and before selective deferred BCG in HIV-exposed infants.

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The largest trial in 2797 African children was well conducted with low risk of bias for most parameters. Risk of bias was uncertain for selective reporting because there were no precise case definition endpoints for active tuberculosis published prior to the trial analysis.

MVA85A added to BCG compared to BCG alone probably has no effect on the risk of developing microbiologically confirmed tuberculosis (RR 0.97, 95% CI 0.58 to 1.62; 3439 participants, 2 trials; moderate-certainty evidence), or the risk of starting on tuberculosis treatment (RR 1.10, 95% CI 0.92 to 1.33; 3687 participants, 3 trials; moderate-certainty evidence). MVA85A probably has no effect on the risk of developing latent tuberculosis (RR 1.01, 95% CI 0.85 to 1.21; 3831 participants, 4 trials; moderate-certainty evidence). Vaccinating people with MVA85A in addition to BCG did not cause life-threatening serious adverse effects (RD 0.00, 95% CI –0.00 to 0.00; 3692 participants, 3 trials; high-certainty evidence). Vaccination with MVA85A is probably associated with an increased risk of local skin adverse effects (3187 participants, 3 trials; moderate-certainty evidence), but not systemic adverse effect related to vaccination (144 participants, 1 trial; low-certainty evidence). This safety profile is consistent with Phase 1 studies which outlined a transient, superficial reaction local to the injection site and mild short-lived symptoms such as malaise and fever.

Authors' conclusions

MVA85A delivered by intradermal injection in addition to BCG is safe but not effective in reducing the risk of developing tuberculosis.

1 May 2019

Up to date

All studies incorporated from most recent search

All published trials found in the last search (10 May, 2018) were included.

PLAIN LANGUAGE SUMMARY

MVA85A vaccine as a booster to BCG for prevention of tuberculosis

What is the aim of this review?

The aim of this Cochrane review was to evaluate the effectiveness and safety of using MVA85A in addition to BCG compared to using BCG alone for prevention of tuberculosis.

Key messages

MVA85A in addition to BCG showed no added benefit to BCG in prevention of acquiring tuberculosis.

What was studied in the review?

Tuberculosis is an infectious airborne disease which affects the lungs and other organs in the body. It can either be active when a person shows signs and symptoms or has confirmatory tests for tuberculosis or latent when a person has inhaled the bacteria before but does not show signs and symptoms of sickness. Currently, there is only one vaccine licensed for prevention of this disease, which is called BCG. However, the ability for the BCG vaccine to prevent tuberculosis differs in different settings and patient groups resulting in tuberculosis still remaining a problem worldwide despite children being immunized. MVA85A is a vaccine that was investigated for prevention of tuberculosis with the hope that when used in addition to BCG it will improve prevention of people getting tuberculosis.

What are the main results of this review?

After examining the research published up to 10 May 2018, we included six study findings from four randomized controlled trials (clinical trials where people are randomly put into one of two or more treatment groups), enrolling 3838 children and adults. Based on these studies of mostly children and adults living in Africa, MVA85A added to BCG compared to BCG alone probably has no effect on the risk of developing active tuberculosis defined as microbiologically confirmed tuberculosis (moderate-certainty evidence) or the risk of starting on tuberculosis treatment (moderate-certainty evidence). MVA85A has no effect on the risk of developing latent tuberculosis (moderate-certainty evidence). MVA85A does not cause any life-threatening serious side effects (highly-certainty evidence). There were more local skin reactions in people vaccinated with MVA85A, however, there was no increase in overall side effects in people given MVA85A.

How up-to-date is this review?

The review authors searched for studies that have been published up to May 2018.

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SUMMARY OF FINDINGS

Summary of findings for the main comparison. MVA85A compared to placebo for preventing tuberculosis

MVA85A compared to placebo for preventing tuberculosis

Patient or population: HIV-positive and -negative adults and children Setting: South Africa, Senegal

Intervention: MVA85A

Comparison: placebo

Outcomes	Anticipated abso CI)	olute effects* (95%	Relative effect (95% CI)	Number of par- ticipants (trials)	Certainty of the evidence (GRADE)	Comments	
	Risk with placebo	Risk with MVA85A		((1103)			
Active tuberculosis: confirmed by cul- ture or Xpert® MTB/ RIF longest reported follow-up	17 per 1000	16 per 1000 (10 to 28)	RR 0.97 (0.58 to 1.62)	3439 (2 RCTs)	⊕⊕⊕⊝ Moderate ^{a,b,c}	Vaccinating people with MVA85A in addition to BCG probably made little or no difference to the risk of developing active tuberculosis.	
Active tuberculosis: started on tubercu- losis treatment	102 per 1000	112 per 1000 (94 to 136)	RR 1.10 (0.92 to 1.33)	3687 (3 RCTs)	⊕⊕⊕⊝ Moderate ^{a,c,d}	Vaccinating people with MVA85A in addition to BCG probably made little or no difference to the risk of needing to start tuberculosis treatment.	
Latent tuberculosis	114 per 1000	115 per 1000 (97 to 138)	RR 1.01 (0.85 to 1.21)	3831 (4 RCTs)	⊕⊕⊕⊝ Moderate ^{c,d,e}	Vaccinating people with MVA85A in addition to BCG probably made little or no difference to the risk of developing latent tuberculosis.	
Serious adverse ef- fects	1 per 1000	1 per 1000 (0 to 4)	RD 0.00 (-0.00 to 0.00) ^f	3692 (3 RCTs)	⊕⊕⊕⊕ High	Vaccinating people with MVA85A in addition to BCG did not cause life-threatening serious ad- verse effects.	
Adverse effects of any severity (lo- cal reactions of the skin)	Vaccination with ated with more re of the injection.g	MVA85A was associ- eactions at the site	-	3187 (3 RCTs)	⊕⊕⊕⊙ Moderate ^{h,i,j}	Vaccinating people with MVA85A in addition to BCG probably increased the risk of having an ad- verse reaction related to vaccination at the site of the injection.	
Adverse effects of any severity (sys- temic symptoms)	Adverse events re malaise, lethargy iting although dif	eported included r, fever, and vom- ferences between	_	144 (1 RCT)	⊕⊕⊝⊝ Low ^{k,l,m}	Vaccinating people with MVA85A in addition to BCG may not have been associated with an in- crease in adverse effects related to vaccination.	

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groups were not significant at a 95% CI level.g

Adverse events of any severity	808 per 1000	849 per 1000 (824 to 873)	RR 1.05 (1.02 to 1.08)	3836 (4 RCTs)	⊕⊕⊕⊕ High ⁿ	Vaccination with MVA85A alone slightly increased the risk of having an adverse event.

*The risk in the intervention group (and its 95% CI) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

BCG: Bacillus Calmette-Guérin; CI: confidence interval; RCT: randomized controlled trial; RD: risk difference; RR: risk ratio.

GRADE Working Group grades of evidence

High certainty: we are very confident that the true effect lies close to that of the estimate of the effect.

Moderate certainty: we are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

Low certainty: our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect.

Very low certainty: we have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of effect.

^{*a*}Not downgraded for risk of bias. The largest trial was at unclear risk of bias due to selective reporting; however, the outcomes presented were unlikely to be affected by this (Tameris 2013).

^bDowngraded by one level for imprecision. Few events and wide CIs containing clinically appreciable benefit and harm.

^cNot downgraded for indirectness. The only trial in HIV-positive adults was stopped early meaning it was underpowered to detect efficacy (Ndiaye 2015). Therefore, evidence of efficacy is more generalizable to infants; however, results in adults were consistent with little or no effect being seen across all endpoints.

^dDowngraded by one level for imprecision. Broad CI containing little or no effect and clinically appreciable harm.

eNot downgraded for risk of bias. The largest trial was at unclear risk of bias due to selective reporting; however, the outcome of latent tuberculosis was unlikely to be affected by this (Tameris 2013).

^fRisk difference presented as explained in our result section.

gExtensive investigation of the vaccine in Phase 1 studies outlined in the Background of this review outlined "a transient, superficial reaction local to the injection site and mild short-lived viral symptoms" consistent with the findings reported in the Phase 2 trials.

^hDowngraded by one level for imprecision. Broad CIs containing clinically appreciable benefit and harm.

ⁱNot downgraded for risk of bias. The largest study reported local adverse events and defined these as solicited by the vaccine (Tameris 2013).

jNot downgraded for heterogeneity. While there might be some heterogeneity between the included trials in terms of time of outcome collection, the outcomes are consistent in favour to placebo as shown in Analysis 1.5.

^kDowngraded by one level for risk of bias. There were some deficiencies in the trial reporting these outcomes.

Additional safety data from Phase 1 studies in 712 participants did not show any adverse effect signals (see section in Background of this review).

^mDowngraded by one level for imprecision. Few events reported in the largest trial (Tameris 2013), data not disaggregated in the second largest trial (Ndiaye 2015).

ⁿNot downgraded for inconsistency. I² value of 37% judged to be non-significant heterogeneity.

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BACKGROUND

Description of the condition

Tuberculosis is an infectious disease caused by *Mycobacterium tuberculosis*. It was estimated that 10 million people developed tuberculosis in 2017. Tuberculosis now ranks first, followed by HIV, as the leading cause of death from an infectious disease worldwide killing an estimated 1.6 million people in 2017, including 300,000 people living with HIV. Over 95% of these people were living in low-and middle-income countries (WHO 2018).

Tuberculosis can be classed as active when people experience signs or symptoms of tuberculosis or have radiological evidence of it. Tuberculosis can also be classified as latent tuberculosis infection (LTBI) where immunological evidence of previous exposure to M tuberculosis exists without clinical or radiological evidence of the disease (CDC 2000). Of healthy adults with immunological evidence of previous exposure to *M* tuberculosis, the overall lifetime risk of progressing to active disease if not treated for the infection is 5% to 10% (Harries 2006). Often this happens months or years after the initial infection in response to a weakening of the body's immune system. The probability of developing active disease is higher in HIV-positive people, people with diabetes, and young children (Baker 2011; Perez-Velez 2012; Tiemersma 2011). Fifty percent of infants with evidence of LTBI will progress to active disease if untreated (Marais 2004). People with LTBI require early diagnosis and treatment to reduce the pool of active tuberculosis cases. This is particularly important in high-risk groups, such as those coinfected with HIV (Sharma 2012). Tuberculosis can be treated with long courses of multiple antibiotics, but the rise of HIV and spread of multidrug-resistant tuberculosis (MDR-TB) means that tuberculosis is still one of the largest threats to public health worldwide (WHO 2018). Structural determinants such as rapid urbanization of populations and economic inequalities, social determinants such as poverty and poor housing, alongside biological factors such as HIV and drug-resistant strains of tuberculosis play a vital role in the spread of tuberculosis through vulnerable populations (Daftary 2012).

The Bacillus Calmette-Guérin (BCG) vaccine is currently the only available vaccine. Epidemiological studies indicate that it has a protective effect against tuberculosis disease in children, particularly against the more severe forms of the disease such as tuberculosis meningitis or miliary tuberculosis (Roy 2014). The effectiveness of BCG differs greatly depending on the site of infection. It has consistent protection against tuberculous meningitis and miliary disease in children but variable protection against pulmonary tuberculosis (Abubakar 2013; Colditz 1995). As a result, despite many areas achieving high coverage of BCG vaccination, the disease remains a problem, and a new tuberculosis vaccine remains an important global research priority (WHO 2018).

Previously it has been impossible to ascertain reliably whether the BCG vaccine protected against active disease or infection with *M tuberculosis*. This was due to the tuberculin skin test being unable to distinguish between cases of LTBI and people who had been vaccinated with BCG (Roy 2014). Therefore, the development and use of interferon γ release assays (IGRA), which can distinguish between tuberculosis infection and vaccination, has proved useful. This has allowed researchers to establish that BCG vaccination reduces the risk of *Mycobacterium* infection in some settings (Eisenhut 2009).

Description of the intervention

Many researchers and policy makers emphasize that a new effective vaccine could be a major contribution to tuberculosis control and elimination as a public health problem (de Cassan 2010). There are 12 vaccine candidates in clinical trials: eight in Phase 2 or Phase 3, and four in Phase 1. They include candidates to prevent the development of tuberculosis, and candidates to help improve the outcomes of treatment for tuberculosis disease (WHO 2018).

The modified Vaccinia Ankara virus-expressing antigen 85A (MVA85A) is a viral vector vaccine based on the modified Vaccinia Ankara (MVA) virus. MVA is an attenuated virus that does not replicate in human tissue and, as such, has been used as a platform to encode multiple antigens and allowing development of multivalent vaccines (Altenburg 2014). In this case, MVA has had pieces of DNA from *M tuberculosis* inserted into it, so that it expresses the antigen 85A. This antigen complex is an enzyme that is involved in the cell wall biosynthesis of *M tuberculosis* and constitutes a vital part of the way in which the bacteria forms its outer mycomembrane. This is important for the viability of the mycobacterium and works as an effective barrier to drug therapies by repelling some antibiotics and preventing them from entering the cell (Favrot 2013).

Immunological studies have shown that a prime boost strategy, where MVA85A is used to boost the effects of BCG, is effective in expanding immune responses specific to *M tuberculosis* (Beveridge 2007). Thus, MVA85A was proposed primarily as a booster to people already vaccinated with BCG (Tameris 2013). Further studies have assessed MVA85A in other regimens including in combination with other viral vector vaccines (Sheehan 2015).

How the intervention might work

MVA85A is the first vaccine since 1968 to be tested in efficacy trials (Tameris 2013). It has been tried with a promise of prolonged antimycobacterial immunity in human UK trials (McShane 2004), and in tuberculosis endemic areas (Hawkridge 2008). The intention is that MVA85A would boost the immune response to tuberculosis above that which is afforded by vaccination with BCG (Roy 2014). MVA85A is administered as a single intradermal dose in people who have already received BCG vaccine (Tameris 2013). Other routes have been studied in animal studies, such as intravenous administration (Romano 2006), and are being considered in humans (Satti 2014).

The researchers who developed the vaccine evaluated its effects in animals and conducted Phase 1 studies in humans. Early literature and reviews by the team noted the vaccine was safe and produced an immune response in several populations (McShane 2004; Rowland 2012).

One independent systematic review of the animal studies, carried out by some members of this Cochrane Review team, raised questions about whether these animal studies provided evidence of efficacy in the various animal models used (Kashangura 2015), when clinical and pathological endpoints were examined in a variety of animal models subjected to challenge studies. This has led to a debate about the reporting of animal studies, in particular the lack of published protocols so that the question being tackled in an animal study is made clear in advance (Cohen 2018). These studies administered BCG, BCG and MVA85A, or

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no vaccine. Afterwards, animals were exposed to tuberculosis challenge. Clearly progression to clinical trial is not solely based on evidence derived from preclinical efficacy studies, and MVA85A was evaluated in a number of trials in humans before proceeding to an efficacy study (McShane 2018). However, preclinical studies remain an important component of the tuberculosis vaccine development paradigm (Barker 2012; McShane 2014).

The systematic review of animal studies pointed out that there was one study in macaques where more monkeys required euthanasia in the MVA85A plus BCG vaccine group than the BCG control group (Kashangura 2015). This led to considerable controversy as to whether the publication of the results were delayed (Cohen 2018). The findings from this study could be the result of chance; or because the vaccine impaired functional immunity; or the result of a separate adverse effect. The vaccine development team then carried out a relatively large number of safety studies in humans; and, in their words, "none of the 14 trials of MVA85A in over 400 humans (the target species) before the infant efficacy trial showed a safety signal" (McShane 2018). The standard approach for Cochrane Reviews within the Cochrane Infectious Diseases Group is to only summarize efficacy trials. However, as the primary concern of the studies included in this review was safety, we summarized the considerable number of Phase 1 studies that the researchers carried out to exclude severe adverse effects attributable to the vaccine in humans in this 'Background' section of the review. We searched registered clinical trial databases (ClinicalTrials.gov, World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP), Pan African Trials Registry, EU Clinical Trials Register) in June 2017 and summarized the Phase 1 studies identified in Table 1. We found 21 separate studies as registered (prospectively and retrospectively) dating from 2003 with the most recent studies scheduled to complete follow-up in 2018. In addition, we found an existing narrative review of Phase 1 studies (Rowland 2012), which summarized Phase 1 safety data relating to selected trials including unpublished data and compared this to selected trials in yellow fever and BCG.

The 21 studies included 712 participants investigated from 2002 with follow-up expected to be completed by 2018. The studies covered a diverse population in the UK, South Africa, Senegal, and The Gambia with HIV-positive and HIV-negative people as well as infants, children, and adults. Intramuscular, intradermal, and aerosolized delivery routes were all investigated. The summary showed most of the adverse effects related to vaccination were mild and were contained locally to the injection site. There were very few serious adverse effects; erythema and mild pain were the most common adverse effects of the vaccine.

Why it is important to do this review

Summarizing the evidence to date will be useful to the public, scientists, and to others interested in innovation in tuberculosis as a case study from laboratory development to field testing. If critical appraisal and systematic review of this vaccine in humans shows no clear effect, this raises questions about any further testing. However, as of November 2017, there were ongoing studies looking at aerosolized delivery of the vaccine (NCT01954563; NCT02532036). In 2017, studies were published that addressed the immunogenicity of the candidate tuberculosis vaccine MVA85A in *Schistosomiasis*-infected teenagers (Wajja 2017), and a further efficacy study in HIV-exposed infants (Nemes 2018).

OBJECTIVES

To assess and summarize the effects of the MVA85A vaccine boosting BCG in humans.

METHODS

Criteria for considering studies for this review

Types of studies

Randomized controlled trials (RCTs) that include measures of clinical efficacy (Phase 2 clinical trials).

Types of participants

Any person regardless of age or HIV status.

Types of interventions

Intervention

MVA85A vaccine regardless of vaccination schedule, dosage, route, or formulation given with BCG.

Control

BCG alone, or Candin[®] (Candida albicans skin test antigen).

Types of outcome measures

Primary outcomes

- Active tuberculosis, defined by:
 - clinical signs and symptoms plus confirmation by microscopy, culture, or Xpert[®] MTB/RIF (an automated nucleic-acid amplification test);
 - * treatment commenced for tuberculosis.

Secondary outcomes

• Latent tuberculosis, diagnosed by IGRA or Mantoux without clinical or radiological evidence of active disease.

Adverse outcomes

- Adverse effects of any severity, defined as "an adverse event for which the causal relation between the intervention and the event is at least a reasonable possibility" (Loke 2011).
- Serious adverse effects, defined as an adverse event attributable to the intervention "leading to death, are life threatening, requires inpatient hospitalisation or prolongation of existing hospitalisation, or result in persistent or significant disability or incapacity" (ICH 1994).
- Adverse events of any severity, defined as "any untoward medical occurrence that may present during treatment with a pharmaceutical product but which does not necessarily have a causal relationship with this treatment" (WHO-ART 2008).
- Abnormal haematological tests during the follow-up period after being vaccinated.
- Abnormal biochemical tests during the follow-up period after being vaccinated.

Search methods for identification of studies

We conducted the literature search up to the 10 May 2018 and identified potential studies regardless of language or publication status (published, unpublished, in press, and in progress).

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



Electronic searches

We searched the following databases using the search terms and strategy described in Appendix 1: the Cochrane Infectious Diseases Group Specialized Register (10 May 2018); the Cochrane Central Register of Controlled Trials (CENTRAL, 2018, Issue 4, published in the Cochrane Library); MEDLINE (PubMed, 1966 to 10 May 2018); Embase (Ovid, 1947 to 10 May 2018); Science Citation Index-Expanded, Social Sciences Citation index, conference proceedings (Web of Science, 1900 to 10 May 2018); and CINAHL (EBSCOHost (1982 to 10 May 2018). We also searched the World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP; www.who.int/ictrp/en/), and ClinicalTrials.gov (clinicaltrials.gov/ct2/home), for trials in progress, up to 10 May 2018, using MVA85A, "modified vaccinia virus Ankara", Ag85A, "Antigen 85A", and tuberculosis OR tuberculosis as search terms.

Searching other resources

We searched the proceedings and abstracts of the following tuberculosis conferences: Union World Conference on Lung Health, European Respiratory Society, and the International Conference of the American Thoracic Society (ATS), from 2012 to 2018. We also handsearched reference lists of relevant papers.

Data collection and analysis

Selection of studies

Two review authors independently screened all abstracts retrieved by the search strategy above using predefined eligibility criteria designed and piloted by the review authors. We excluded clearly irrelevant studies. We searched for multiple publications using studies from the same data set. We retrieved full-text copies for all trials thought to be potentially relevant. Two review authors (SoJ and SaJ) independently assessed all identified trials for inclusion in the review using the predefined inclusion criteria.

We resolved any disagreements in assessment through discussion. In cases of unresolved differences, a third review author adjudicated. We kept records of the initial results and the changes after discussion. We also kept a list all studies excluded after fulltext assessment in the Characteristics of excluded studies table. We illustrated the study selection process in a PRISMA diagram (Figure 1).



Figure 1. Study flow diagram.



Data extraction and management

We designed and piloted data extraction forms. Two review authors independently performed data extraction. We gathered information from each included trial separately on trial characteristics. These included:

- study setting, design, study duration, population sample size, and power calculations;
- baseline characteristics of study population including age, sex, weight, prematurity, HIV, other comorbidity, whether

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breastfeeding, race, HIV status, antiretroviral therapy (ART), CD4 count, and viral load;

- intervention and control group vaccine dosages, routes of administration, and times of vaccination;
- time of outcome measure after administering MVA85A;
- duration of follow-up, withdrawals from the study, and reasons for withdrawal.

All outcomes were dichotomous, so we tabulated the numbers of participants who developed tuberculosis or an adverse event (n) with the total sample size number (N) in each comparison group. We documented the different definitions of outcomes in the trials for further consideration and only combined data from endpoints that were similar across studies.

Assessment of risk of bias in included studies

We assessed risk of bias for RCTs using the Cochrane 'Risk of bias' tool (Higgins 2011). Two review authors independently assessed studies for risk of bias. We resolved any disagreement through discussion and, where necessary, through consultation with a third review author.

We assessed sequence generation (if predictable method used) and allocation concealment for selection bias and detection bias by looking at blinding methods. We also considered both the intention of blinding and the success of blinding for each outcome. If there was no description of the procedure, for example how randomization was done, we marked it as unclear.

In addition, we examined the objectivity of outcome measures, use of intention-to-treat (ITT) analysis, loss to follow-up, and selective outcome reporting to assess the risk of bias in included studies. We assessed whether outcome measures were specified a priori and whether the published endpoints matched those specified in study protocols.

We assessed incomplete outcome data in each included trial to determine the proportion of missing results and whether it affected the results in terms of event risk and effect size. We assessed if reasons for missing data were related to adverse events or death from MVA85A and if missing data were balanced in the two experimental groups to have an overall decision on risk associated with incomplete outcome data.

We assessed other dimensions to risk of bias, including conflicts of interest, large differences in baseline characteristics, and early cessation of the trial.

We assessed the included trials for risk of bias of adverse events by examining if monitoring was active or passive; whether participants and outcome assessors were blinded; whether the outcome data reporting was complete; whether all participants were included; and whether data analysis was independent of pharmaceutical companies (Table 2; Bukirwa 2014). We also looked at the times when data were collected in comparison to when they were reported. All this information was included under overall study assessment of blinding, selective outcome reporting, incomplete outcome data, or other biases.

Measures of treatment effect

We analysed all data using Review Manager 5 (Review Manager 2014). We pooled dichotomous data using risk ratios (RR) with their

corresponding 95% confidence intervals (CI). When inappropriate due to a small number of events in each group, we presented the pooled data using risk difference (RD) with their 95% CI.

Unit of analysis issues

For included studies that had multiple intervention arms, we included data from these studies by splitting the control group so that participants were only included in the meta-analysis once.

Dealing with missing data

In our protocol, we anticipated that if the amount of incomplete outcome data was such that the trials were thought to be at a high risk of bias, we may have used imputation and perform sensitivity analyses to investigate the impact of these missing data. However, we identified no studies where missing data affected our ability to measure outcomes. Therefore, we used available-case analysis, as planned in our protocol.

Assessment of heterogeneity

We assessed extracted data from included trials to find key differences in population groups, study setting, intervention and control groups, dosages and route of vaccine administration, or timing between BCG and boosting. We assessed degree of risk of bias, when and how the outcome was measured, and variation in treatment effects.

We determined the level of heterogeneity by inspecting forest plots for overlapping CIs. We judged a Chi² P value significance level of 0.1 or less as likely heterogeneity. An I² statistic value of less than 40% was regarded as not showing any significant heterogeneity.

Assessment of reporting biases

There was an insufficient number of trials included and so we were unable to assess for publication bias using funnel plots or Egger regression.

Data synthesis

We used the fixed-effect Mantel-Haenszel model for meta-analysis where there was little heterogeneity. The intention for metaanalysis of adverse outcomes was limited to three to five of the most frequent adverse effects and all those that were considered to be serious. However, due to different methods of monitoring adverse effects that in turn lead to different results where meta-analysis could not be performed, we gave a narrative report.

Subgroup analysis and investigation of heterogeneity

We intended to explore heterogeneity by: subgroup by children and adults; background prevalence of tuberculosis (or tuberculosis incidence in the control group); HIV status; and geographical location. However, there were not enough trials to explore such subgroups when we found high heterogeneity.

We considered random-effects meta-analysis if subgroup analysis did not explain the heterogeneity. We applied the l^2 statistic according to guidance of: less than 40% as not significant heterogeneity; 30% to 60% representing moderate heterogeneity; 50% to 90% representing substantial heterogeneity; and 75% to 100% considerable heterogeneity (Higgins 2011). We regarded a Chi² P value significance level of 0.1 or less and an l^2 statistic greater than 40% as showing significant heterogeneity, in which case we

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either considered a random-effects model or did not perform metaanalysis.

Sensitivity analysis

We did not perform sensitivity analysis for imputed data, risk of bias, or any other peculiarities between the trials identified during the review process.

Certainty of the evidence

We assessed the certainty of the evidence using the GRADE approach (Schünemann 2013). We constructed a 'Summary of findings' table, which outlines the main review findings alongside the certainty of the evidence.

RESULTS

Description of studies

Results of the search

We identified 153 records, with 152 records remaining after removing duplicates. We excluded 118 records based on title and abstract and assessed the full text of 34 articles. We excluded 28 full-text articles. Six articles fulfilled the eligibility criteria and were included in the review. See Figure 1 for the flow diagram of inclusion and exclusion of studies in the review.

Included studies

Six studies (3838 participants) that met our inclusion criteria reported findings from four Phase 2 clinical trials (Ndiaye 2015; Nemes 2018; Scriba 2011; Tameris 2013). Andrews 2017 and Bunyasi 2017 presented data based on the Tameris 2013 clinical trial. The six included studies are described in the Characteristics of included studies table.

Setting and time

All took place in South Africa involving rural and urban areas between 2008 and 2015, with one trial that took place at two sites: South Africa and Senegal (Ndiaye 2015).

Source of funding

Aeras sponsored five trials (Andrews 2017; Bunyasi 2017; Ndiaye 2015; Nemes 2018; Tameris 2013). The University of Oxford sponsored one trial (Scriba 2011). The Wellcome Trust funded all the trials. Other funders were Oxford Emergent Tuberculosis Consortium (OETC) for Ndiaye 2015 and Tameris 2013, the European and Developing Countries Clinical Trials Partnership and the Bill and Melinda Gates Foundation for Ndiaye 2015, the UK Medical Research Council for Nemes 2018, and the EuropeAID European Commission for Scriba 2011. Andrews 2017 and Bunyasi 2017 conducted further follow-up based on the participants enrolled in Tameris 2013, and mentioned that there was no specific additional funding for the analysis performed.

Participants

Five trials included infants (Andrews 2017; Bunyasi 2017; Nemes 2018; Scriba 2011; Tameris 2013). One trial assessed the efficacy and safety of the vaccine in adults with HIV (Ndiaye 2015). Tameris 2013 and Scriba 2011 recruited infants who were HIV-negative, while Nemes 2018 assessed the vaccine in newborns of HIV-positive mothers. None of the trials reported other morbidities. In Tameris

2013, 412 (29.4%) participants in the intervention group and 268 (26.4%) participants in the control group were preterm.

Interventions

Intervention

All the infants in the intervention groups received a single dose of intradermal MVA85A. In the trial recruiting adults, the 324 adults allocated in the intervention group received a second dose (booster) of intradermal vaccine six months after the first dose (Ndiaye 2015). The vaccine was given at a dose of 1 × 10⁸ plaqueforming units (pfu) in Ndiaye 2015, Nemes 2018, and Tameris 2013. Scriba 2011 assessed three different doses of the vaccine by giving a dose of 2.5x10⁷ pfu, 5x10⁷ pfu and 1x 10⁸ pfu to 36 participants in each of the three groups. All the infants in Scriba 2011 and Tameris 2013 received the BCG vaccine in the first four weeks of life, prior to receiving the MVA85A vaccine, as an inclusion criteria. Nemes 2018 gave the MVA85A vaccine to the neonates in the first 96 hours of life, with no prior administration of BCG, and gave BCG at eight weeks of age only to HIV-negative infants. Ndiaye 2015 did not mention whether the adults they recruited received BCG.

Comparator

Five trials gave Candida skin test antigen (Candin[®]) as a placebo, using the same route (intradermal) and schedule (one or two doses) as for the intervention group in each of the trial (Andrews 2017; Bunyasi 2017; Ndiaye 2015; Nemes 2018; Tameris 2013). Scriba 2011 gave the infants in the comparator group one dose of pneumococcal 7-valent conjugate vaccine by the intramuscular route.

Outcomes

Three studies reported different endpoints as measures of tuberculosis disease (Ndiaye 2015; Nemes 2018; Tameris 2013). These are compared in Table 3.

All the included studies reported data on latent tuberculosis (or tuberculosis infection) to assess either efficacy or safety outcomes. Four trials looked at safety outcomes, including adverse effects of any severity, serious adverse effects, and adverse events of any severity (Ndiaye 2015; Nemes 2018; Scriba 2011; Tameris 2013). Tameris 2013 collected data on biochemical or haematological blood test findings but did not report this element of their primary outcome. Ndiaye 2015 collected data on blood tests but did not report disaggregated findings. Only Scriba 2011 and Nemes 2018 reported on blood test data collected.

Length and method of follow-up

Scriba 2011 followed up participants for 24 weeks, Nemes 2018 for 52 weeks, Ndiaye 2015 for at least six months after the last participant was enrolled, and Tameris 2013 for up to 39 months. Andrews 2017 was an observational follow-up study of the participants enrolled in Tameris 2013; authors analysed the data collected at day 336 after the intervention and at the end of the study, which ranged from six to 24 months after day 336. Bunyasi 2017 followed the participants recruited in Tameris 2013 for a median of five years.

Investigators of five studies used diary cards to record adverse events during the seven days following vaccination (Andrews 2017; Bunyasi 2017; Ndiaye 2015; Scriba 2011; Tameris 2013); Nemes 2018

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did not mention this. Researchers performed blood investigations at several intervals in all trials, to detect adverse events and to assess immunogenicity. Ndiaye 2015 and Tameris 2013 performed active follow-up every three months to identify signs, symptoms, or exposure to tuberculosis that merited further investigation, while this was done at irregular but planned intervals in Scriba 2011 and Nemes 2018. The long-term follow-up study was based on passive surveillance based on the electronic tuberculosis register database (Bunyasi 2017).

Excluded studies

We excluded 28 studies from the review, with the reasons for exclusion listed in the Characteristics of excluded studies table.

Studies awaiting classification

We did not identify any studies that are awaiting classification.

Ongoing studies

We did not identify any ongoing studies.

Risk of bias in included studies

See Characteristics of included studies table for the assessment of the risk of bias for each included study. See Figure 2 and Figure 3 for the risk of bias summaries.

Figure 2. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.



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Allocation

Five trials were at low risk of selection bias (Andrews 2017; Bunyasi 2017; Ndiaye 2015; Nemes 2018; Tameris 2013). They

reported adequate sequence generation and methods of allocation concealment. Scriba 2011 used systematic allocation at a 3:1 ratio allowing predictability of the sequence (high risk of bias).

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Blinding

Three studies had adequate blinding of participants, study personnel, laboratory assessors, and clinical assessors and were at low risk for performance and detection bias in all domains (Ndiaye 2015; Nemes 2018; Tameris 2013). Five studies reported blinding of participants and study personnel (Andrews 2017; Bunyasi 2017; Ndiaye 2015; Nemes 2018; Tameris 2013). Scriba 2011, an open-label trial with different routes of administration for placebo and vaccine, had low risk of detection bias for laboratory assessors as outcomes were objective and high risk of detection bias for subjective assessments by clinicians. Two studies were at unclear risk of detection bias for laboratory assessors and clinicians (Andrews 2017; Bunyasi 2017). Andrews 2017 did not provide any details on blinding, while Bunyasi 2017 reported on post-trial data and had no information on how data was collected from registers.

Incomplete outcome data

Four trials reported details of all randomized participants (Ndiaye 2015; Nemes 2018; Scriba 2011; Tameris 2013). Only a few participants randomized were not included in the analysis, without resulting in a disbalance between the intervention and control groups. Indeed, three participants were randomized in the control group in Tameris 2013, but not included in the efficacy analysis (two of them were not included either in the safety analysis), while five participants were randomized (four in the intervention group and one in the control group), but not included in the efficacy analysis in Ndiaye 2015. As a result we considered these studies at low risk of attrition bias. There were no details of how many of each group came from the 119 participants excluded from Tameris 2013 for analysis in Bunyasi 2017. Andrews 2017 and Bunyasi 2017 had an unclear risk of attrition bias as these were follow-up studies from Tameris 2013, and there were unclear discrepancies with those reported previously.

Selective reporting

Nemes 2018 was prospectively registered and appeared free of selective outcome reporting as ascertained from data in trial registers and reports of trials. We also judged Scriba 2011 at low risk of reporting bias, with all the outcomes reported in their methods section presented in the results.

Four studies were at unclear risk of bias due to selective reporting (Andrews 2017; Bunyasi 2017; Ndiaye 2015; Tameris 2013). There were multiple instances where predefined endpoints were poorly defined or were deviated from in the final reported results as laid out in Table 4.

Description of Tameris 2013 published prior to commencement of the trial (NCT00953927) stated that the authors intended to report endpoints of clinical disease based on "observational cohort studies." This was subsequently changed following the publication of the trial in October 2013 to include "clinicallyderived tuberculosis diagnostic criteria." The main trial reports adapting the primary elements proposed in a consensus statement (Graham 2012). There was no record of the change in approach from empirically derived endpoints to endpoints developed by the investigators in the study protocol.

Tameris and colleagues reported on three outcomes with complex definitions (Table 3).

- Endpoint one, described as "primary efficacy endpoint," comprising nine criteria, which included a binary measure of quantiFERON conversion.
- Endpoint two, described as "exploratory efficacy endpoint," comprising nine criteria.
- Endpoint three, described as "exploratory efficacy endpoint," which was defined as "all participants placed on treatment for tuberculosis."

The difference between endpoints one and two, which varied in the direction of the point estimate of the effect, was 5 mm on a tuberculin skin test or household contact with acid-fast bacilli (AFB) smear-positive person (Table 3). The process of defining these three endpoints was unexplained, and it is unclear why these specific definitions were used. These endpoint definitions were only used in this trial and not in subsequent studies.

In a subsequent critique, Behr and colleagues noted that the outcomes reported in the trial did not include the simple measure of a positive microbiological endpoint (Behr 2013). The endpoint used in the abstract was endpoint one, which authors have settled as primary efficacy outcome, while endpoints two and three were reported as exploratory outcomes. The complexity of the definitions and the analysis in Behr's paper pointed to the risk of selective reporting. This may not have been intentional, but arose with post-hoc approaches with different approaches to expressing the results, but could be excluded if outcomes were precisely and clearly defined a priori. The only information publicly available prior to the trial commencing were broad descriptions of the outcome. Hence for selective reporting the classification was unclear.

Andrews 2017 was at unclear risk of reporting bias as this was a nested observational study and there was no prespecified study protocol. Ndiaye 2015 was at unclear risk of reporting bias as the authors commented that there were no differences between biological and haematological tests; however, no data or how these data were analysed to come to this conclusion were reported.

Other potential sources of bias

We considered that the risk of other potential biases was unclear in all included studies. We were concerned as a number of the authors were involved in the private company manufacturing the vaccine or were patent holders for MVA85A. In these circumstances, it would be good practice for this to be declared in the publication. Only one study declared no conflicts in relation to patent holding (Scriba 2011).

Two trials reported a role of funders in design, data analysis, and manuscript writing (Ndiaye 2015; Tameris 2013), and one study had employees of the funder involved in manuscript writing (Andrews 2017). Ndiaye 2015 calculated incident tuberculosis cases from day 28 after vaccination versus from day 0 in Tameris 2013. This was likely to be due to the risk of pre-existing undiagnosed tuberculosis being inappropriately counted as developing following the intervention. If participants are not followed from the start of the intervention then a period of follow-up has been excluded, and participants who experienced the outcome soon after intervention will be missing from analyses. We considered this to be of unclear risk of bias as it is unclear if this impacted on outcomes.

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Adverse events

For adverse events, we conducted additional assessments on adequacy of safety monitoring and completeness of reporting for participant-reported outcomes and laboratory tests taken (Table 5). Four trials reported on safety outcomes (Ndiaye 2015; Nemes 2018; Scriba 2011; Tameris 2013). Monitoring of participant-reported outcomes was active in all trials and blinding was adequate in two trials (Nemes 2018; Tameris 2013). All trials reported specified timing of data collection but only one study reported under some of the days (Scriba 2011). None of the trials completely reported outcomes on prespecified time points including for laboratory results. All trials reported all participants who received intervention per-protocol. Timing of taking laboratory tests was inadequate in Scriba 2011 and Tameris 2013 as there was no clear indication of tests being taken at the end of the study.

Effects of interventions

See: Summary of findings for the main comparison MVA85A compared to placebo for preventing tuberculosis

See Summary of findings for the main comparison.

Active tuberculosis

Studies varies in the way they defined active tuberculosis (see section "description of studies" (Table 3)). Tameris 2013 and Ndiaye 2015 reported hierarchical endpoints including microbiologically confirmed tuberculosis, composite clinical definitions, and participants starting on tuberculosis treatment, with no significant effect consistently seen across endpoints (Analysis 2.1; Analysis 2.2; Table 3; Table 6).

Tameris 2013 reported three endpoints in their main manuscript, with endpoint one described as their primary efficacy endpoint (RR 0.82, 95% CI 0.52 to 1.30, point estimate favouring MVA85A). A fourth endpoint was described in the supplementary material, taking into account the microbiologically confirmed cases of tuberculosis. Other outcomes (endpoint two, endpoint three, and endpoint four of microbiologically confirmed cases) were not statistically different, although their point estimate favoured placebo (endpoint two: RR 1.05, 95% CI 0.73 to 1.53; endpoint 3: RR 1.10, 95% CI 0.91 to 1.33; endpoint four (microbiologically confirmed): RR 1.10, 95% CI 0.60 to 2.00; Analysis 2.1; Figure 4).

Figure 4. Forest plot of comparison: 2 Comparison of endpoints, outcome: 2.1 Tameris 2013: incidence of tuberculosis according to post-hoc endpoints.

	MVA8	5A	Place	bo	Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Tameris 2013 (1)	32	1399	39	1395	0.82 [0.52, 1.30]	
Tameris 2013 (2)	55	1399	52	1395	1.05 [0.73, 1.53]	
Tameris 2013 (3)	196	1399	177	1395	1.10 [0.91, 1.33]	- ++
Tameris 2013 (4)	22	1399	20	1395	1.10 [0.60, 2.00]	
						0.5 0.7 1 1.5 2 Favours MVA85A Favours placebo
<u>Footnotes</u> (1) Endpoint 1 compo	site clinio	alend	point			

(2) Endpoint 2 composite clinical endpoint

(3) Endpoint 3 composite clinical endpoint

(4) Microbiologically confirmed

Two studies reported no effect of MVA85A on cases of active tuberculosis confirmed by culture or Xpert® MTB/RIF (RR 0.97, 95%

CI 0.58 to 1.62; 3439 participants, two trials) (Analysis 1.1; Figure 5; Ndiaye 2015; Tameris 2013).

Figure 5. Forest plot of comparison: 1 MVA85A Vs Placebo, outcome: 1.1 Tuberculosis confirmed by culture or Xpert® MTB/RIF longest reported follow-up.

	MVA8	5A	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Ndiaye 2015 (1)	6	320	9	325	30.8%	0.68 [0.24, 1.88]	
Tameris 2013 (2)	22	1399	20	1395	69.2%	1.10 [0.60, 2.00]	
Total (95% CI)		1719		1720	100.0%	0.97 [0.58, 1.62]	+
Total events	28		29				
Heterogeneity: Chi ² = 0.64, df = 1 (P = 0.42); l ² = 0% Test for overall effect: Z = 0.13 (P = 0.90)							0.01 0.1 1 10 100 Favours MVA85A Favours placebo
<u>Footnotes</u>							

At least 6 months' follow-up

(2) At least 15 months' follow-up

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Three studies (Ndiaye 2015; Nemes 2018; Tameris 2013) reported no effect of MVA85A on cases of active tuberculosis when considering

patients started on tuberculosis treatment (RR 1.10, 95% CI 0.92 to 1.33; 3687 participants, 3 trials; Analysis 1.2; Figure 6).

Figure 6. Forest plot of comparison: 1 MVA85A versus placebo, outcome: 1.2 Active tuberculosis: started on tuberculosis treatment.

	MVA8	5A	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Ndiaye 2015	8	320	9	325	4.7%	0.90 [0.35, 2.31]	
Nemes 2018	5	123	3	125	1.6%	1.69 [0.41, 6.93]	<u> </u>
Tameris 2013	196	1399	177	1395	93.7%	1.10 [0.91, 1.33]	—
Total (95% CI)		1842		1845	100.0%	1.10 [0.92, 1.33]	+
Total events	209		189				
Heterogeneity: Chi ² =	0.53, df=	2 (P =	0.77); I ≊ =	= 0%			
Test for overall effect:	Z=1.05	(P = 0.2	29)				Favours MVA85A Favours BCG alone

Nemes 2018 reported active tuberculosis as defined by participants starting tuberculosis treatment. One participant in this trial was diagnosed by culture; however, the authors did not report what intervention this participant received.

Latent tuberculosis

Four studies reported no effect of MVA85A on cases of latent tuberculosis (RR 1.01, 95% CI 0.85 to 1.21; 3831 participants, four trials; Analysis 1.3).

Scriba 2011 was underpowered and not designed to detect measures of efficacy. However, they reported latent tuberculosis, presumably as a measure of safety, as this outcome was poorly defined a priori.

Adverse effects

Four studies reported effects of any severity (Table 7). We presented the effect of the estimates for adverse effects of any severity

with disaggregated (Analysis 1.4; Figure 7) and aggregated data (Analysis 1.5) to provide detailed information as provided by the study authors. However, we did not perform meta-analysis of the estimates due to high heterogeneity. Local reactions of the skin at the injection site was the most common adverse effect associated with the vaccine MVA85A, this was reported in three studies, with the three studies showing direction towards more adverse effects in the intervention group (3187 participants; Nemes 2018; Scriba 2011; Tameris 2013). However, only one study reported systemic symptoms defined as fever, lethargy, malaise, and vomiting (144 participants; Scriba 2011). Therefore, we chose to report adverse effects of any severity disaggregated by local reactions of the skin and systemic symptoms in our Summary of findings for the main comparison as different amount of information is provided for each group (Scriba 2011).

Figure 7. Forest plot of comparison: 1 MVA85A versus placebo, outcome: 1.4 Adverse effects of any severity.

	MVA8	5A	Place	bo	Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
1.4.1 Local: skin						
Nemes 2018	121	123	118	125	1.04 [0.99, 1.09]	+
Scriba 2011	106	108	6	36	5.89 [2.84, 12.23]	│ _
Tameris 2013	1251	1399	628	1396	1.99 [1.87, 2.11]	+
1.4.2 Malaise						
Scriba 2011	6	108	1	36	2.00 [0.25, 16.06]	
1.4.3 Lethargy						
Scriba 2011	6	108	2	36	1.00 [0.21, 4.74]	
1.4.4 Any fever						
Scriba 2011	18	108	2	36	3.00 [0.73, 12.30]	
1.4.5 Vomiting						
Scriba 2011	6	108	2	36	1.00 [0.21, 4.74]	

Favours MVA85A Favours placebo

Three studies reported no increase in the risk of experiencing a serious adverse effect attributable to MVA85A (3692 participants; Analysis 1.6). Nemes 2018 reported serious adverse events and specified that none of them were related to the investigational product. Therefore, we classified this as no serious adverse effects following the definition of our review.

Adverse events of any severity

Four studies reported a small increase in the risk of experiencing an adverse event of any severity following vaccination with MVA85A (RR 1.05, 95% CI 1.02 to 1.08; 3836 participants; Analysis 1.7; Table 8). Adverse effects related to the vaccine and adverse events not attributed to the vaccine were conflated in the largest trial. No disaggregated data were available.

Abnormal haematological and biochemical tests

Three studies reported abnormal haematological or biochemical laboratory tests. The percentage of those with elevated liver enzymes ranged from 2.8% to 25% in the three different groups reported in Scriba 2011 and there was a dose-response effect of MVA85A. However, none of the doses showed a significant increase at a 95% CI. Ndiaye 2015 reported that routine haematological and biochemical test results did not differ between study groups but disaggregated data were not reported. Nemes 2018 reported no difference between groups in the percentage of people with abnormal biochemical tests (11.4% versus 10.4%), but disaggregated data were not reported. The largest study performed haematological and biochemical tests but did not report data (Tameris 2013). We summarized the report and findings of abnormal haematological and biochemical tests in Table 9, and presented the effect of estimate for abnormal biochemical tests only (Analysis 1.8), as only one study reported disaggregated data for abnormal haematological tests.

DISCUSSION

Summary of main results

Vaccinating people with MVA85A in addition to BCG:

- probably makes little or no difference to the risk of developing active tuberculosis (moderate-certainty evidence);
- probably makes little or no difference to the risk of needing to start tuberculosis treatment (moderate-certainty evidence);
- probably does not have an important effect on the risk of developing latent tuberculosis (moderate-certainty evidence);
- does not cause life-threatening serious adverse effects (highcertainty evidence);
- probably increases the risk of having an adverse reaction related to vaccination at the site of the injection (moderate-certainty evidence);
- may not be associated with an increase in systemic adverse effects related to vaccination (low-certainty evidence).

Vaccination with MVA85A alone slightly increases the risk of having an adverse event (high-certainty evidence).

Overall completeness and applicability of evidence

This review included trials from two countries in Africa. No studies that measured efficacy of the MVA85A vaccine have been carried out elsewhere. The review included studies on HIV-positive adults, HIV-negative infants, and infants exposed to HIV. It would be reasonable to generalize the results of these findings to other populations of HIV-negative infants. The early cessation of the only trial in HIV-positive adults, resulting in reduced follow-up from two years to minimum six months and a reduction of study sample size from 1200 to 625, led this study to be underpowered for evaluation of efficacy (Ndiaye 2015). This may have limited the certainty of any inferences made to adults with HIV at high risk of contracting tuberculosis in terms of efficacy of MVA85A in

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this population. The effect of tuberculosis vaccination would be very similar regardless of geographical variation. Data from this review consistently showed no effect of the vaccine. As such, it is reasonable to generalize these findings to broader populations. For safety outcomes, the Phase 1 studies that we summarized in the Background section and Table 1, included adults, children, and infants from the UK and three African countries. Most of the adverse effects related to vaccination were mild and were contained locally to the injection site. This supports the trial findings summarized in this review.

Certainty of the evidence

Overall the included studies were well-conducted. For most of our outcomes, there were few events and broad CIs for the pooled estimates of effect which contained clinically appreciable benefit and harm or no effect (see Summary of findings for the main comparison).

In the largest trial, the main reported endpoint (endpoint one) point estimate was in the direction of benefit of the vaccine on tuberculosis disease (Analysis 2.1; Tameris 2013). Whether this was due to the definition of endpoints or due to statistical heterogeneity was unclear. To minimize the impact of this inconsistency we presented results for cases diagnosed microbiologically and cases defined by being started on treatment. This was felt to reflect the most specific measure of efficacy and a measure of the real-world situation. As a result of this, the methodological uncertainties surrounding case definition did not reduce our confidence in the effect estimates.

Failure to follow-up participants from the start of intervention for efficacy measures in Ndiaye 2015 risked biasing outcomes. While it is plausible that participants with undiagnosed active tuberculosis would be inappropriately picked up, it is also plausible that participants who hypothetically could have developed tuberculosis immediately after vaccination would be excluded from analysis. However, the potential impact of this was unclear and as such we did not downgrade due to risk of bias for efficacy outcomes including this study.

In terms of latent tuberculosis, using the online calculator at www.sealedenvelope.com/power/binary-noninferior/ at a significance level of 5% and with 80% power at a failure rate of 11% and a non-inferiority limit of 5% a sample size per group of 484 would be sufficient to demonstrate non-inferiority. Therefore, in terms of risk of developing latent tuberculosis where we had high certainty evidence that MVA85A had no important effect in reducing risk, we are confident that future trials are unlikely to change this result as we had 3831 participants in the analysis versus a minimum number of 484 participants required in each group.

Regarding the safety outcomes, the summary of findings from the Phase I trials for MVA85A performed in adults, adolescents, and infants with 712 participants showed that most of the adverse effects related to vaccination were mild and were contained locally to the injection site, and none of the trials reported a serious adverse event attributable to the vaccine. This supports the certainty of the evidence found in this review.

Potential biases in the review process

We followed standard methods in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). The Cochrane Infectious Disease Group Information Specialist performed a comprehensive literature search with no restriction in language to identify all eligible studies, thus it is unlikely that we missed any large studies. We were unable to formally assess publication bias as fewer than 10 studies met our inclusion criteria.

Agreements and disagreements with other studies or reviews

No previous systematic reviews have been undertaken looking at the effects of MVA85A.

There has been much debate over the contribution of animal studies to the progression of MVA85A vaccine to trial (Cohen 2018; McShane 2018). We systematically assessed Phase 1 and 2 data and we found no difference in tuberculosis incidence in any population, and no increase in the risk of serious adverse effects attributable to the vaccine. There was a small increase in the risk of experiencing any adverse event.

The findings of this review are consistent in that MVA85A is not efficacious for preventing tuberculosis and that there is no evidence that the MVA85A vaccine caused any serious harm to participants in the trials during its investigation.

AUTHORS' CONCLUSIONS

Implications for practice

MVA85A in conjunction with Bacillus Calmette-Guérin (BCG) has no effect on the risk of developing active or latent tuberculosis.

Implications for research

Researchers should define outcomes precisely before starting the trial. If composite outcomes are developed during the trial, this process needs to be transparent, clearly reported, and published prior to breaking the randomized code. Standardization of outcome measures for tuberculosis vaccine efficacy may make it easier for future researchers in the field and allow easy comparison and meta-analysis of different study outcomes.

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REFERENCES

References to studies included in this review

Andrews 2017 {published data only}

Andrews JR, Nemes E, Tameris M, Landry BS, Mahomed H, McClain JB, et al. Serial QuantiFERON testing and tuberculosis disease risk among young children: an observational cohort study. *Lancet Respiratory Medicine* 2017;**5**(4):282-90.

Bunyasi 2017 {published data only}

Bunyasi EW, Luabeya AK, Tameris M, Geldenhuys H, Mulenga H, Landry BS, et al. Impact of isoniazid preventive therapy on the evaluation of long-term effectiveness of infant MVA85A vaccination. *International Journal Tuberculosis and Lung Disease* 2017;**21**(7):778-83. [DOI: dx.doi.org/10.5588/ijtld.16.0709]

Ndiaye 2015 {published data only}

Ndiaye BP, Thienemann F, Ota M, Landry BS, Camara M, Dièye S, et al. Safety, immunogenicity, and efficacy of the candidate tuberculosis vaccine MVA85A in healthy adults infected with HIV-1: a randomised, placebo-controlled, phase 2 trial. *Lancet Respiratory Medicine* 2015;**3**(3):190-200.

Nemes 2018 {published data only}

Nemes E, Hesseling A, Tameris M, Mauff K, Downing K, Mulenga H, et al. Safety and immunogenicity of newborn MVA85A vaccination and selective, delayed Bacille Calmette-Guerin (BCG) for infants of HIV infected mothers: a Phase 2 randomized controlled trial. *Clinical Infectious Diseases* 2018;**66**(4):554-63. [DOI: 10.1093/cid/cix834]

Scriba 2011 {published data only}

Scriba TJ, Tameris M, Mansoor N, Smit E, van der Merwe L, Mauff K, et al. Dose-finding study of the novel tuberculosis vaccine, MVA85A, in healthy BCG-vaccinated infants. *Journal of Infectious Diseases* 2011;**203**(12):1832-43. [DOI: 10.1093/infdis/ jir195]

Tameris 2013 {published data only}

Tameris MD, Hatherill M, Landry BS, Scriba TJ, Snowden MA, Lockhart S, et al. Safety and efficacy of MVA85A, a new tuberculosis vaccine, in infants previously vaccinated with BCG: a randomised, placebo-controlled phase 2b trial. *Lancet* 2013;**381**(9871):1021-8.

References to studies excluded from this review

Brookes 2008 {published data only}

Brookes RH, Hill PC, Owiafe PK, Ibanga HB, Jeffries DJ, Donkor SA, et al. Safety and immunogenicity of the candidate tuberculosis vaccine MVA85A in West Africa. *PLOS One* 2008;**3**(8):e2921.

Bunyasi 2015 {published data only}

Bunyasi EW, Tameris M, Geldenhuys H, Schmidt BM, Luabeya AK, Mulenga H, et al. Evaluation of Xpert(R) MTB/ RIF assay in induced sputum and gastric lavage samples from young children with suspected tuberculosis from the MVA85A TB vaccine trial. *PLOS One* 2015;**10**(11):e0141623.

Dieye 2013 {published data only}

Dieye TN, Ndiaye BP, Dieng AB, Fall M, Britain N, Vermaak S, et al. Two doses of candidate TB vaccine MVA85A in antiretroviral therapy (ART) naive subjects gives comparable immunogenicity to one dose in ART plus subjects. *PLOS One* 2013;**8**(6):e67177.

Harris 2011 {published data only}

Harris S, Meyer J, Satti I, Kosotv M, Rowland R, Poulton ID, et al. Comparison of the safety and immunogenicity of a candidate TB vaccine, MVA85A, given by the intramuscular or intradermal route in BCG-vaccinated adults. Annual Congress of the British-Society-for-Immunology. 2011; Vol. 135:55.

Harris 2014a {published data only}

Harris SA, Meyer J, Satti I, Marsay L, Poulton ID, Tanner R, et al. Evaluation of a human BCG challenge model to assess antimycobacterial immunity induced by BCG and a candidate tuberculosis vaccine, MVA85A, alone and in combination. *Journal of Infectious Diseases* 2014;**209**(8):1259-68.

Hawkridge 2008 {published data only}

Hawkridge T, Scriba TJ, Gelderbloem S, Smit E, Tameris M, Moyo M, et al. Safety and immunogenicity of a new tuberculosis vaccine, MVA85A, in healthy adults in South Africa. Journal of Infectious disease 2008; Vol. 198, issue 4:544-52.

Matsumiya 2014a {published data only}

Matsumiya M. Immune response to MVA85A or placebo in BCG-vaccinated South African infants at days 0 and 28 postvaccination. Arrayexpress-repository, V1 2014. [https:// www.ebi.ac.uk/arrayexpress/experiments/E-GEOD-56561]

Matsumiya 2014b {published data only}

Matsumiya M, Harris SA, Satti I, Stockdale L, Tanner R, O'Shea MK, et al. Inflammatory and myeloid-associated gene expression before and one day after infant vaccination with MVA85A correlates with induction of a T cell response. Biomed Central Infectious Diseases 2014; Vol. 14:314.

Matsumiya 2014c {published data only}

Matsumiya M. GSE56559: immune response to MVA85A or placebo in BCG-vaccinated South African infants at day 1 post-vaccination. Arrayexpress-repository, V1 2014. [https:// www.ebi.ac.uk/arrayexpress/experiments/E-GEOD-56559]

McShane 2004 {published data only}

McShane H, Pathan AA, Sander CR, Keating SM, Gilbert SC, Huygen K, et al. Recombinant modified vaccinia virus Ankara expressing antigen 85A boosts BCG-primed and naturally acquired antimycobacterial immunity in humans. Nature Medicine 2004; Vol. 10:1240-4.

Meyer 2013 {published data only}

Meyer, J, Harris, S. A, Satti I, Poulton ID, Poyintz HC, Tanner R, et al. Comparing the safety and immunogenicity of a candidate TB vaccine MVA85A administered by intramuscular and intradermal delivery. Vaccine 2013; Vol. 31, issue 2013:1026-33.

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



Minassian 2011 {published data only}

Minassian, AM, Rowland R, Beveridge NE, Poulton ID, Satti I, Harris S, et al. A Phase I study evaluating the safety and immunogenicity of MVA85A, a candidate TB vaccine, in HIVinfected adults. British Medical Journal 2015; Vol. 1:e000223.

Minhinnick 2016 {published data only}

Minhinnick A, Satti I, Harris S, Wilkie M, Sheehan S, Stockdale L, et al. A first-in-human phase 1 trial to evaluate the safety and immunogenicity of the candidate tuberculosis vaccine MVA85A-IMX313, administered to BCG-vaccinated adults. Vaccine Vol. 34, issue 2016:1412-21.

Mulenga 2015 {published data only}

MulengaH, Tameris MD, Luabeya KK, Geldenhuys H, Scriba TJ, Hussey GD, et al. The Role of Clinical Symptoms in the Diagnosis of Intrathoracic Tuberculosis in Young Children. Pediatric Infectious Diseases Journal Vol. 34:1157-62.

Odutola 2012 {published data only}

Odutola AA, Owolabi OA, Owiafe PK, Mcshane H, Ota MO. A new TB vaccine, MVA85A, induces durable antigen-specific responses 14 months after vaccination in African infants. Vaccine 2012; Vol. 30:5591-4.

Ota 2011 {published data only}

Ota MO, Odutola AA, Owiafe PK, Donkor S, Owolabi OA, Brittain NJ, et al. Immunogenicity of the tuberculosis vaccine MVA85A is reduced by coadministration with EPI vaccines in a randomized controlled trial in Gambian infants. Sci. Transl. Med. 2011; Vol. 3:88ra56.

Pathan 2007 {published data only}

Pathan AA, Sander CR, Fletcher HA, Poulton I, Alder NC, Beveridge NE, et al. Boosting BCG with recombinant modified vaccinia ankara expressing antigen 85A: different boosting intervals and implications for efficacy trials. *PLOS One* 2007;**2**(10):e1052.

Pathan 2012 {published data only}

Pathan AA, Minassian AM, Sander CR, Rowland R, Porter DW, Poulton ID, et al. Effect of vaccine dose on the safety and immunogenicity of a candidate TB vaccine, MVA85A, in BCG vaccinated UK adults. *Vaccine* 2012;**30**(38):5616-24.

Rowland 2012 {published data only}

Rowland R, Brittain N, Poulton ID, Minnassain AM, Sander C, Porter DW, et al. A review of the tolerability of the candidate TB vaccine, MVA85A compared with BCG and Yellow Fever vaccines, and correlation between MVA85A vaccine reactogenicity and cellular immunogenicity. *Trials in vaccinology* 2012;**1**:27-35.

Rowland 2013 {published data only}

Rowland R, Pathan AA, Satti I, Poulton ID, Matsimuya MM, Whitaker M, et al. Safety and immunogenicity of an FP9vectored candidate tuberculosis vaccine (FP85A), alone and with candidate vaccine MVA85A in BCG-vaccinated healthy adults: a phase I clinical trial. *Human Vaccines and Immunotherapeutics* 2013;**9**(1):50-62.

Sander 2009 {published data only}

Sander CR, Pathan AA, Beveridge NE, Poulton I, Minassian A, Alder N, et al. Safety and immunogenicity of a new tuberculosis vaccine, MVA85A, in Mycobacterium tuberculosis-infected individuals. *American Journal of Respiratory and Critical Care Medicine* 2009;**179**:724-33.

Satti 2014 {published data only}

Satti I, Meyer J, Harris SA, Thomas ZM, Griffiths K, Antrobuset RD, et al. Safety and immunogenicity of a candidate tuberculosis vaccine MVA85A delivered by aerosol in BCGvaccinated healthy adults: a phase 1, double-blind, randomised controlled trial. *Lancet Infectious diseases* 2014;**14**:939-46.

Scriba 2010 {published data only}

Scriba TJ, Tameris M, Mansoor N, Thomas ZM, Griffiths K, Antrobus RD, et al. Modified vaccinia Ankara-expressing Ag85A, a novel tuberculosis vaccine, is safe in adolescents and children, and induces polyfunctional CD4+ T cells. *European Journal of Immunology* 2010;**40**:279-90.

Scriba 2012 {published data only}

Scriba TJ, Tameris M, Smit E, Van der Merwe I, Jane Hughes E, Kadira B, et al. A phase IIa trial of the new tuberculosis vaccine, MVA85A, in HIV- and/or Mycobacterium tuberculosis-infected adults. *American Journal of Respiratory and Critical Care Medicine* 2012;**185**(7):769-78.

Sheehan 2015 {published data only}

Sheehan S, Harris SA, Satti I, Hokey DA, Dheenadhayalan V, Stockdale L, et al. A Phase I, Open-Label Trial, Evaluating the Safety and Immunogenicity of Candidate Tuberculosis Vaccines AERAS-402 and MVA85A, Administered by Prime-Boost Regime in BCG-Vaccinated Healthy Adults. *PLOS One* 2015;**10**(11):e0141687.

Tameris 2014 {published data only}

Tameris M, Geldenhuys H, Luabeya AK, Smit E, Hughes JE, Vermaak S, et al. The candidate TB vaccine, MVA85A, induces highly durable Th1 responses. *PLOS One* 2014;**9**(2):e87340.

Tanner 2014 {published data only}

Tanner R, Kakalacheva K, Miller E, Pathan AA, Chalk R, Sander CR, et al. Serum indoleamine 2,3-dioxygenase activity is associated with reduced immunogenicity following vaccination with MVA85A. *Biomed Central Infectious Diseases* 2014;**14**:660.

Whelan 2009 {published data only}

Whelan KT, Pathan AA, Sander CR, Fletcher HA, Poulton I, Alder NC, et al. Safety and immunogenicity of boosting BCG vaccinated subjects with BCG: comparison with boosting with a new TB vaccine, MVA85A. *PLOS One* 2009;**10**(11):e5934.

Additional references

Abubakar 2013

Abubakar I, Pimpin L, Ariti C, Beynon R, Mangtani P, Sterne JA, et al. Systematic review and meta-analysis of the current evidence on the duration of protection by bacillus Calmette-

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



Guérin vaccination against tuberculosis. *Health Technology Assessment* 2013;**17**(37):1-372, v-vi.

Altenburg 2014

Altenburg AF, Kreijtz JH, de Vries RD, Song F, Fux R, Rimmelzwaan GF, et al. Modified vaccinia virus ankara (MVA) as production platform for vaccines against influenza and other viral respiratory diseases. *Viruses* 2014;**6**(7):2735-61.

Baker 2011

Baker MA, Harries AD, Jeon CY, Hart JE, Kapur A, Lönnroth K, et al. The impact of diabetes on tuberculosis treatment outcomes: a systematic review. *BMC Medicine* 2011;**9**:81.

Barker 2012

Barker L, Hessel L, Walker B. Rational approach to selection and clinical development of TB vaccine candidates. *Tuberculosis* 2012;**92**(Suppl 1):S25-9.

Behr 2013

Behr MA, Schwartzman K, Pai M. Tuberculosis vaccine trials. *Lancet* 2013;**381**(9885):2252-3.

Beveridge 2007

Beveridge NE, Price DA, Casazza JP, Pathan AA, Sander CR, Asher TE, et al. Immunisation with BCG and recombinant MVA85A induces long-lasting, polyfunctional Mycobacterium tuberculosis-specific CD4+ memory T lymphocyte populations. *European Journal of Immunology* 2007;**37**(11):3089-100.

Bukirwa 2014

Bukirwa H, Unnikrishnan B, Kramer CV, Sinclair D, Nair S, Tharyan P. Artesunate plus pyronaridine for treating uncomplicated Plasmodium falciparum malaria. *Cochrane Database of Systematic Reviews* 2014, Issue 3. [DOI: 10.1002/14651858.CD006404.pub2]

CDC 2000

Centers for Disease Control and Prevention. Diagnostic standards and classification of tuberculosis in adults and children. This official statement of the American Thoracic Society and the Centers for Disease Control and Prevention was adopted by the ATS Board of Directors, July 1999. This statement was endorsed by the Council of the Infectious Disease Society of America. *American Journal of Respiratory and Critical Care Medicine* 2000;**161**(4 Pt 1):1376-95.

Cohen 2018

Cohen D. Oxford TB vaccine study calls into question selective use of animal data. *BMJ* 2018;**360**:j5845.

Colditz 1995

Colditz GA, Berkey CS, Mosteller F, Brewer TF, Wilson ME, Burdick E, et al. The efficacy of bacillus Calmette-Guérin vaccination of newborns and infants in the prevention of tuberculosis: meta-analyses of the published literature. *Paediatrics* 1995;**96**(1 Pt 1):29-35.

Daftary 2012

Daftary A, Padayatchi N. Social constraints to TB/HIV healthcare: accounts from coinfected patients in South Africa. *AIDS Care* 2012;**24**(12):1480-6.

de Cassan 2010

de Cassan SC, Pathan AA, Sander CR, Minassian A, Rowland R, Hill AV, et al. Investigating the induction of vaccine-induced Th17 and regulatory T cells in healthy, Mycobacterium bovis BCG-immunized adults vaccinated with a new tuberculosis vaccine, MVA85A. *Clinical and Vaccine Immunology* 2010;**17**(7):1066-73.

Eisenhut 2009

Eisenhut M, Paranjothy S, Abubakar I, Bracebridge S, Lilley M, Mulla R, et al. BCG vaccination reduces risk of infection with Mycobacterium tuberculosis as detected by gamma interferon release assay. *Vaccine* 2009;**27**(44):6116-20.

Favrot 2013

Favrot L, Grzegorzewicz AE, Lajiness DH, Marvin RK, Boucau J, Isailovic D, et al. Mechanism of inhibition of Mycobacterium tuberculosis antigen 85 by ebselen. *Nature Communications* 2013;**4**:2748. [DOI: 10.1038/ncomms3748]

Graham 2012

Graham SM, Ahmed T, Amanullah F, Browning R, Cardenas V, Casenghi M, et al. Evaluation of tuberculosis diagnostics in children: 1. Proposed clinical case definitions for classification of intrathoracic tuberculosis disease. Consensus from an expert panel. *Journal of Infectious Diseases* 2012;**205**(2):S199-208.

Griffiths 2011

Griffiths KL, Pathan AA, Minassian AM, Sander CR, Beveridge NE, Hill AV, et al. Th1/Th17 cell induction and corresponding reduction in ATP consumption following vaccination with the novel Mycobacterium tuberculosis vaccine MVA85A. *PLOS One* 2011;**6**(8):e23463.

Harries 2006

Harries AD, Dye C. Tuberculosis. *Annals of Tropical Medicine and Parasitology* 2006;**100**(5-6):415-31.

Harris 2014b

Harris SA, Satti I, Matsumiya M, Stockdale L, Chomka A, Tanner R, et al. Process of assay selection and optimization for the study of case and control samples from a phase IIb efficacy trial of a candidate tuberculosis vaccine, MVA85A. *Clinical and Vaccine Immunology* 2014;**21**(7):1005-11.

Higgins 2011

Higgins JP, Green S, editor(s). Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0 (updated March 2011). The Cochrane Collaboration, 2011. Available from handbook.cochrane.org. Chichester (UK): Available from www.handbook.cochrane.org.

Ibanga 2006

Ibanga HB, Brookes RH, Hill PC, Owiafe PK, Fletcher HA, Lienhardt C, et al. Early clinical trials with a new tuberculosis

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



vaccine, MVA85A, in tuberculosis-endemic countries: issues in study design. *Lancet Infectious Diseases* 2006;**6**(8):522-8.

ICH 1994

International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH). Clinical safety data management: definitions and standards for expedited reporting E2A.1994. ICH harmonised tripartite guideline. Current Step 4 version dated 27 October 1994. www.ich.org/fileadmin/Public_Web_Site/ICH_Products/ Guidelines/Efficacy/E2A/Step4/E2A_Guideline.pdf (accessed 23 August 2017).

Kashangura 2015

Kashangura R, Sena ES, Young T, Garner P. Effects of MVA85A vaccine on tuberculosis challenge in animals: systematic review. *International Journal of Epidemiology* 2015;**44**(6):1970-81. [DOI: 10.1093/ije/dyv142]

Loke 2011

Loke YK, Price D, Herxheimer A. Chapter 14: Adverse effects. In: Higgins JP, Green S, editor(s). Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0 (updated March 2011). The Cochrane Collaboration, 2011. Available from handbook.cochrane.org.

Manjaly 2016

Manjaly TZ, Satti I, Wilkie M, Harris S, Riste M, Hamidi A, et al. A phase I trial evaluating aerosol administration of a candidate TB vaccine, MVA85A, as a way to induce potent local cellular immune responses and avoid anti-vector immunity. *American Journal of Respiratory and Critical Care Medicine* 2016;**193**:A5487.

Marais 2004

Marais BJ, Gie RP, Schaaf HS, Hesseling AC, Obihara CC, Starke JJ, et al. The natural history of childhood intra-thoracic tuberculosis: a critical review of literature from the prechemotherapy era. *International Journal of Tuberculosis and Lung Disease* 2004;**8**(4):392-402.

Matsumiya 2013

Matsumiya M, Stylianou E, Griffiths K, Lang Z, Meyer J, Harris SA, et al. Roles for Treg expansion and HMGB1 signaling through the TLR1-2-6 axis in determining the magnitude of the antigen-specific immune response to MVA85A. *PLOS One* 2013;**8**(7):e67922.

McShane 2014

McShane H, Williams A. A review of preclinical animal models utilised for TB vaccine evaluation in the context of recent human efficacy data. *Tuberculosis* 2014;**94**(2):105-10.

McShane 2018

McShane H, Hill A, Hatherill M, Tameris M, Shea J, Ginsberg A. Helen McShane and colleagues reply to Deborah Cohen. *BMJ* 2018;**360**:k236.

NCT00395720

NCT00395720. The safety and immunogenicity of a TB vaccine; MVA85A, in healthy volunteers who are infected with HIV. clinicaltrials.gov/ct2/show/NCT00395720 (first received 25 August 2017).

NCT00423566

NCT00423566. A phase I study of the safety and immunogenicity of a recombinant MVA vaccine encoding a secreted antigen from *M. tuberculosis*, antigen 85A, delivered intradermally by a needle injection in healthy volunteers. clinicaltrials.gov/ct2/ show/NCT00423566 (first received 25 August 2017).

NCT00423839

NCT00423839. A phase I study of the safety and immunogenicity of MVA85A in healthy Gambian volunteers. clinicaltrials.gov/ct2/ show/NCT00423839 (first received 25 August 2017).

NCT00427453

NCT00427453. A phase I study of the safety and immunogenicity of a recombinant MVA vaccine encoding a secreted antigen from *M. tuberculosis*, antigen 85A, delivered intradermally by a needle injection in healthy volunteers who have received BCG immunisation 1 month previously. clinicaltrials.gov/ct2/show/ NCT00427453 (first received 25 August 2017).

NCT00427830

NCT00427830. A phase I study of the safety and immunogenicity of a recombinant MVA vaccine encoding a secreted antigen from *M. tuberculosis*, antigen 85A, delivered intradermally by a needle injection in healthy volunteers who have previously received BCG. clinicaltrials.gov/ct2/show/NCT00427830 (first received 25 August 2017).

NCT00456183

NCT00456183. Safety and immunogenicity of MVA85A in volunteers latently infected with TB. clinicaltrials.gov/ct2/show/ NCT00456183 (first received 25 August 2017).

NCT00460590

NCT00460590. Safety and immunogenicity of MVA85A, in healthy volunteers in Cape Town. clinicaltrials.gov/ct2/show/ NCT00460590 (first received 25 August 2017).

NCT00465465

NCT00465465. A study of 2 doses of a new TB vaccine, MVA85A, in healthy volunteers previously vaccinated with BCG. clinicaltrials.gov/ct2/show/NCT00465465 (first received 25 August 2017).

NCT00480454

NCT00480454. Safety, immunogenicity, and impact of MVA85A, on the immunogenicity of the EPI vaccines. clinicaltrials.gov/ ct2/show/NCT00480454 (first received 25 August 2017).

NCT00480558

NCT00480558. A study of MVA85A, in asymptomatic volunteers infected with TB, HIV or both. clinicaltrials.gov/ct2/show/ NCT00480558 (first received 25 August 2017).

NCT00548444

NCT00548444. T-cell turnover following vaccination with MVA85A. clinicaltrials.gov/ct2/show/NCT00548444 (first received 25 August 2017).

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



NCT00653770

NCT00653770. A phase I study to assess the safety and immunogenicity of tuberculosis (TB) vaccine candidates FP85A and MVA85A. clinicaltrials.gov/ct2/show/NCT00653770 (first received 25 August 2017).

NCT00731471

NCT00731471. A phase I study of a new tuberculosis (TB) vaccine, MVA85A, in healthy volunteers with HIV. clinicaltrials.gov/ct2/show/NCT00731471 (first received 25 August 2017).

NCT00953927

NCT00953927. A study of MVA85A in healthy infants. clinicaltrials.gov/ct2/show/NCT00953927 (first received 25 August 2017).

NCT01181856

NCT01181856. Safety of tuberculosis vaccine, MVA85A, administered by the intramuscular route and the intradermal route. clinicaltrials.gov/ct2/show/NCT01181856 (first received 25 August 2017).

NCT01194180

NCT01194180. A BCG challenge model study to assess antimycobacterial immunity induced by BCG and a candidate TB vaccine, MVA85A. clinicaltrials.gov/ct2/show/NCT01194180 (first received 25 August 2017).

NCT01497769

NCT01497769. Safety of tuberculosis vaccine, MVA85A, administered by the aerosol route and the intradermal route. clinicaltrials.gov/ct2/show/NCT01497769 (first received 25 August 2017).

NCT01683773

NCT01683773. Safety study of tuberculosis vaccines AERAS-402 and MVA85A. clinicaltrials.gov/ct2/show/NCT01683773 (first received 25 August 2017).

NCT01829490

NCT01829490. Safety study of ChAdOx185A vaccination with and without MVA85A boost in healthy adults. clinicaltrials.gov/ show/NCT01829490 (first received 11 April 2013).

NCT01879163

NCT01879163. Phase I trial evaluating safety and immunogenicity of MVA85A-IMX313 compared to MVA85A in BCG vaccinated adults. clinicaltrials.gov/ct2/show/ NCT01879163 (first received 25 August 2017).

NCT01954563

NCT01954563. Study evaluating aerosol and intradermal administration of a candidate tuberculosis (TB) vaccine, MVA85A, as a way to increase immune response and avoid antivector immunity. clinicaltrials.gov/ct2/show/NCT01954563 (first received 25 August 2017).

NCT02532036

NCT02532036. MVA85A aerosol versus intramuscular vaccination in adults with latent *Mycobacterium tuberculosis*

(M. tb) Infection. clinicaltrials.gov/show/NCT02532036 (first received 25 August 2015).

Owiafe 2012

Owiafe P, Hill P, Ibanga HB, Brookes RH, McShane H, Sutherland JS, et al. Differential cytokine levels in adults induced by a novel candidate TB boost vaccine, MVA85Aaccording to previous BCG vaccination status. *Journal of Vaccines & Vaccination* 2012;**3**(7):158.

Perez-Velez 2012

Perez-Velez CM, Marais BJ. Tuberculosis in children. *New England Journal of Medicine* 2012;**367**(4):348-61.

Review Manager 2014 [Computer program]

Nordic Cochrane Centre, The Cochrane Collaboration. Review Manager 5 (RevMan 5). Version 5.3. Copenhagen: Nordic Cochrane Centre, The Cochrane Collaboration, 2014.

Romano 2006

Romano M, D'Souza S, Adnet PY, Laali R, Jurion F, Palfiet K, et al. Priming but not boosting with plasmid DNA encoding mycolyltransferase Ag85a from *Mycobacterium tuberculosis* increases the survival time of *Mycobacterium bovis* BCG vaccinated mice against low dose intravenous challenge with *M. Tuberculosis* H37Rv. *Vaccine* 2006;**24**:3353-64.

Roy 2014

Roy A, Eisenhut M, Harris RJ, Rodrigues LC, Sridhar S, Habermann S, et al. Effect of BCG vaccination against *Mycobacterium tuberculosis* infection in children: systematic review and meta-analysis. *BMJ* 2014;**349**:g4643. [DOI: 10.1136/ bmj.g4643]

Schünemann 2013

Schünemann H, Brożek J, Guyatt G, Oxman A. GRADE handbook for grading quality of evidence and strength of recommendations. Available from guidelinedevelopment.org/ handbook October 2013.

Sharma 2012

Sharma SK, Mohanan S, Sharma A. Relevance of latent TB infection in areas of high TB prevalence. *Chest* 2012;**142**(3):761-73.

Tiemersma 2011

Tiemersma EW, van der Werf MJ, Borgdorff MW, Williams BG, Nagelkerke NJ. Natural history of tuberculosis: duration and fatality of untreated pulmonary tuberculosis in HIV negative patients: a systematic review. *PLOS One* 2011;**6**(4):e17601.

Wajja 2017

Wajja A, Kizito D, Nassanga B, Nalwoga A, Kabagenyi J, Kimuda S, et al. The effect of current Schistosoma mansoni infection on the immunogenicity of a candidate TB vaccine, MVA85A, in BCG-vaccinated adolescents: an open-label trial. *PLOS Neglected Tropical Diseases* 2017;**11**(5):e0005440.

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



WHO 2018

World Health Organization. Global tuberculosis report 2018. www.who.int/tb/publications/global_report/en/ (accessed 3 January 2018).

WHO-ART 2008

Uppsala Monitoring Centre. WHO adverse reaction terminology (WHO-ART). www.who-umc.org/vigibase/services/learn-moreabout-who-art/ (accessed 23 August 2017).

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

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References to other published versions of this review

Kashangura 2018

Kashangura R, Jullien S, Garner P, Young T, Johnson S. MVA85A vaccine to enhance BCG for preventing tuberculosis. *Cochrane Database of Systematic Reviews* 2018, Issue 1. [DOI: 10.1002/14651858.CD012915]

Methods	Study objective: to investigate the relation between QFT conversion interferon-γ values and risk of sub- sequent active TB disease and of QFT reversion.
	This is a follow-up study of the Tameris 2013 trial.
	Study design: observational follow-up study based on a parallel-group, randomized, placebo-con- trolled double-blind Phase 2b trial.
	Study duration: 41 months
	Length of follow-up: ≥ 15 months after enrolment, and up to 41 months (based on the Tameris 2013 tri- al)
	Follow-up method: no additional data for this study than described in the Tameris 2013 trial.
	Losses to follow-up: 285/2797 children from Tameris 2013 to enrolment for this study analysis at day 336; 467/2512 children from day 336 until the end of the study.
	Power calculation: not relevant for this observational follow-up study.
Participants	Number: 2512/2797 participants enrolled in Tameris 2013 were quantiFERON-negative at enrolment and had another quantiFERON done at day 336 and were therefore enrolled for this study analysis. No disaggregated data on age and sex between intervention and control groups among these 2512 participants.
	Target group: infants aged 4–6 months
	Inclusion criteria
	Healthy infants aged 4–6 months
	Received BCG vaccination within 7 days of birth
	 Received all age-appropriate routine immunizations, and 2 doses of pneumococcal conjugate vaccine at least 28 days before study vaccination (amended to 14 days during enrolment) HIV ELISA-negative
	OuantiFERON-negative
	No substantial exposure to a person with known TB
	Written informed consent obtained from parents/guardian
	• Weight: by chart > 3rd percentile on study day 0 or, if < 3rd percentile, infant had stable growth pattern
	Ability to complete follow-up period as required by the protocol
	Completed simultaneous enrolment in the Aeras Vaccine Development Registry protocol
	Exclusion criteria
	Acute illness on study day 0
	 Fever ≥ 37.5 °C on study day 0

• Evidence of significant active infection on study day 0

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	 Received a EPI immunization within 14 days prior to study day 0 Historical or virological evidence of individual or maternal HIV-1 infection History of allergic disease or reactions likely to be exacerbated by any component of the study vaccine Previous medical history, or evidence, of an intercurrent illness that may compromise the safety of the infant in the study Evidence of chronic hepatitis from any cause History or evidence of any systemic disease on physical examination or any acute, chronic or intercurrent illness that, in the opinion of the investigator, may have interfered with the evaluation of the safety or immunogenicity of the vaccine History of or known TB or treatment for TB Shared residence since birth with a person with active TB or on ATT for < 2 months
	HIV status: negative
	Other comorbidities: none reported
	Preterms:
	 Intervention group: 412 (29.4%) Control group: 368 (26.4%)
Interventions	Intervention group
	 Vaccine: MVA85A/AERAS-485 Dosage: 1 × 10⁸ pfu in 0.06 mL Route: intradermal Schedule: at day 1, 1 dose Timing after BCG: inclusion criteria request BCG given during the first 7 days of life.
	Control group
	 Vaccine: Candida skin test antigen (Candin, AllerMed, USA) Dosage: 0.06 mL Route: intradermal Schedule: at day 1, 1 dose Timing after BCG: inclusion criteria request BCG given during the first 7 days of life.
Outcomes	Outcomes included in this review
	Active TB
	Outcomes not included in this review:
	QFT converters
Notes	Country: South Africa
	Setting: rural, near Cape Town
	Background prevalence of TB: extremely high. The overall incidence of TB in South Africa in 2011 was estimated to be almost 1%, and the incidence of TB in children aged < 2 years was about 3% at the trial site.
	Study dates: enrolment 15 July 2009 to 4 May 2011 and follow-up until 60 days after the 25 October 2012
	Study sponsor: Aeras

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Andrews 2017 (Continued)

Other funders: Wellcome Trust and Oxford Emergent TB Consortium (OETC). No additional funding than from the Tameris 2013 trial was obtained for the analysis of these data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote from report: "Young children were randomly assigned (1:1) using inde- pendently generated sequences with block sizes of four to receive one dose of the vaccine MVA85A or Candida spp skin test antigen (placebo control)."
		Comment: an independent statistician prepared the randomization sched- ule as reported in the trial where the data came from that is referenced above (Tameris 2013).
Allocation concealment (selection bias)	Low risk	Comment: voice response system adequately concealed allocation of inter- vention as reported in Tameris 2013.
Blinding of participants	Low risk	Comment: same as in Tameris 2013. It did not affect long-term follow-up.
mance bias) All outcomes		Quote from Tameris 2013: "Parents or legal guardians of study participants, study staff administering vaccine or undertaking follow up clinical assess- ments and laboratory staff were masked to intervention group assignment."
		"Doses were prepared and labelled in masked syringes by an unmasked study pharmacist."
Blinding of outcome as- sessment (detection bias): laboratory assessors	Unclear risk	Comment: no information on whether laboratory assessors were blinded.
Blinding of outcome as- sessment (detection bias):	Unclear risk	Quote from report: "Study clinicians were not masked to QFT values, but strict case definitions were used that excluded QFT results."
clinical assessors All outcomes		Comment: although clinicians were not masked to QFT values, relevant out- come of conversion is objective and authors used strict case definitions. May not necessarily affect incidence in the two groups as there were no QFT dif- ferences between placebo and MVA85A at 336 days (baseline). No details on whether they were masked to group (MVA85A or placebo).
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote from report: "Among the 2797 young children enrolled in the MVA85A trial (Tameris 2013) 2772 (99%) young children had a negative QFT at enrol- ment, five (<1%) had no quantitative results available, and 20 (1%) had an in- determinate result. 1399 young children were allocated to MVA85A and 1398 were allocated to placebo. Among those 2772 young children with a negative QFT at baseline, 2512 (91%) had a QFT done at the day 336 visit."
		Comment: no imputation
		Of above 2512, 172 positive and 13 indeterminate. Numbers of negative and converted did not add up to the initial study group.
Selective reporting (re- porting bias)	Unclear risk	Comment: outcome/objective of this study not seen in protocol for trial (MVA85A 020 TRIAL). Could not find separate protocol for Andrews trial.
Other bias	Unclear risk	Comment: employees and beneficiaries of funders were involved in design, analysis, and manuscript writing. This study was a follow-up of children enrolled in Tameris 2013 trial.

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



Bunyasi 2017

Methods	Study objective: to evaluate the long-term effectiveness of infant MVA85A vaccination against TB.									
	This is a long-term follow-up study of the Tameris 2013 trial.									
	Study design: retrospective passive follow-up of the randomized controlled trial									
	Study duration: 22 months for enrolment in the original trial									
	Length of follow-up: median of 5 years' follow-up									
	Follow-up method: passive surveillance based on the electronic TB register database.									
	Losses to follow-up: there was some inconsistency between the number of participants included for this long-term follow-up study and the number of participants who were lost to follow-up at an early point in the original trial (Tameris 2013).									
	Power calculation: not relevant for this observational follow-up study									
Participants	Number: 2794 in the Tameris 2013 trial, 2678 included in this long-term follow-up analysis									
	Median age: 4.8 years (IQR 4.4 to 5.2) at the end of the extended follow-up period, comparable across intervention and control groups with no detailed data given in the manuscript.									
	Target group: infants aged 4–6 months									
	Inclusion criteria for the base trial									
	Healthy infants aged 4–6 months									
	Received BCG vaccination within 7 days of birth									
	 Received all age-appropriate routine immunizations, and 2 doses of pneumococcal conjugate vaccin at least 28 days before study vaccination (amended to 14 days during enrolment) 									
	HIV ELISA-negative									
	QuantiFERON-negative									
	 No substantial exposure to a person with known TB 									
	 Written informed consent obtained from parents/guardian 									
	 Weight: by chart > 3rd percentile on study day 0 or, if < 3rd percentile, infant has shown a stable growth pattern 									
	Ability to complete follow-up period as required by the protocol									
	Completed simultaneous enrolment in the Aeras Vaccine Development Registry protocol									
	Exclusion criteria for the base trial									
	Acute illness on study day 0									
	 Fever ≥ 37.5 °C on study day 0 									
	 Evidence of significant active infection on study day 0 									
	 Received a EPI immunization within 14 days prior to study day 0 									
	 Historical or virological evidence of individual or maternal HIV-1 infection 									
	 History of allergic disease or reactions likely to be exacerbated by any component of the study vaccine 									
	 Previous medical history, or evidence, of an intercurrent illness that may compromise the safety of the infant in the study 									
	Evidence of chronic hepatitis from any cause									
	 History or evidence of any systemic disease on physical examination or any acute, chronic or intercurrent illness that, in the opinion of the investigator, may interfere with the evaluation of the safety or immunogenicity of the vaccine. 									
	evaluation of the safety of infinitunogenicity of the valuence									
	• FISCOLY OF OF KHOWIETE OF LEAGHIEFTE OF THE									
	 Shared residence since birth with a person with active 1B of on ATT for < 2 months 									

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)



Bunyasi 2017 (Continued)	HIV status: negative									
	Other comorbidities: n	one reported								
	Preterms in the initial sample size of the base trial									
	 Intervention group: 412 (29.4%) Control group: 368 (26.4%) 									
Interventions	Intervention group									
	 Vaccine: MVA85A/AE Dosage: 1 × 10⁸ pfu Route: intradermal Schedule: at day 1, 1 Timing after BCG: in 	ERAS-485 in 0.06 mL 1 dose I dose criteria request BCG given during the first 7 days of life.								
	Control group									
	 Vaccine: Candida sk Dosage: 0.06 mL Route: intradermal Schedule: at day 1, Timing after BCG: in 	in test antigen (Candin, AllerMed, USA) 1 dose Iclusion criteria request BCG given during the first 7 days of life.								
Outcomes Outcomes included in this review										
	 Active TB. Definition used was the endpoint 3 described in Tameris 2013: participants ment for TB by a health professional. Latent TB, defined by a positive quantiFERON or a positive TST 									
	Outcomes not included in this review									
	 Subgroup analysis of active TB and latent TB in children who received and did no prophylaxis. 									
Notes	Country: South Africa									
	Setting: rural, near Cap	be Town								
	Background prevalenc estimated to be almost site.	e of TB: extremely high. The overall incidence of TB in South Africa in 2011 was t 1%, and the incidence of TB in children aged < 2 years was about 3% at the trial								
	Study dates: enrolment from 15 July 2009 to 4 May 2011 and follow-up to 2014									
	Study sponsor: Aeras									
	Other funders: Wellcome trust and Oxford Emergent TB Consortium (OETC). No additional f from the Tameris 2013 trial was obtained for the analysis of these data.									
Risk of bias										
Bias	Authors' judgement	Support for judgement								
Random sequence genera- tion (selection bias)	Low risk	Quote from Tameris 2013: "We randomly allocated infants in a 1.1 ratio with a block size of 4 using interactive voice /online response system"								
		"An independent statistician prepared the randomisation schedule."								

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

Bunyasi 2017 (Continued)

Allocation concealment (selection bias)	Low risk	Comment: voice response system adequately concealed allocation of intervention as reported in Tameris 2013.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Comment: same as in Tameris 2013. It did not affect long term follow-up. Quote from Tameris 2013: "Parents or legal guardians of study participants, study staff administering vaccine or undertaking follow-up clinical assess- ments and laboratory staff were masked to intervention group assignment." "Doses were prepared and labelled in masked syringes by an unmasked study pharmacists."
Blinding of outcome as- sessment (detection bias): laboratory assessors	Unclear risk	Not applicable.
Blinding of outcome as- sessment (detection bias): clinical assessors All outcomes	Unclear risk	Quote from report: "We also obtained post-trial data from a regional electronic TB register (ETR) (2012–2014). Comment: no information on how data were collected from this register. Clini- cal diagnosis of TB was also a subjective outcome.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote from report: "199 participants discontinued FU [follow-up] early." Comment: 119 participants were excluded from Tameris 2013 for analysis in the current study. No details on how many participants there were from each group of the study.
Selective reporting (re- porting bias)	Unclear risk	Comment: raw data not reported. Only reported incidence rate ratios. There were no disaggregated data on missing data for each group. Number of participants with TB were not reported per group. Only incidence per year.
Other bias	Unclear risk	Quote from report: "The authors received no specific funding for this work," "Conflicts of interest: none declared." "Study is a follow up to Tameris 2013 where the trial sponsor contributed to study design, data interpretation, and writing of the manuscript."

Ndiaye 2015

Methods Study objective: to assess the safety, immunogenicity, and efficacy of MVA85A vaccine in adults HIVpositive. Study design: multicentre randomized double-blind placebo-controlled trial, Phase 2 Study duration: 46 months (from August 2011 to May 2014) Length of follow-up: ≥ 6 months after enrolment Follow-up method • Diary card to report adverse events during the 7 days following vaccination. Direct questionnaire to enquire about adverse events on days 7 and 28 after vaccination. • • Blood tests for routine haematological and biochemical analysis, and for peripheral CD4 cell count and HIV-1 viral load at screening, before booster vaccination, and on days 7 and 28 after vaccination. Blood test for peripheral CD4 cell count and HIV-1 viral load every 3 months until 6 months after booster vaccination. Active follow-up every 3 months until the last participant enrolled had completed 6 months of follow-up after the booster vaccination.

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Ndiaye 2015 (Continued)	Losses to follow-up: 14 participants. 5/324 (1.5%) in the intervention group and 9/326 (2.7%) in the con- trol group. Additionally, 3 participants in the intervention group and 2 in the control group withdrew consent; and 2 participants in the intervention group and 4 in the control group died before the end of the study. In total, 325/649 participants completed the study.			
	Power calculation: the sample size calculation was planned to detect active TB. However, after the Tameris 2013 efficacy data were revised, the authors changed the trial design with safety as the primary objective. A smaller sample size was considered and follow-up was shortened. Therefore, the present trial was underpowered to detect an effect on active TB.			
Participants	Number: 649 (292 from Cape Town, 358 from Dakar).			
	 Intervention group (324 participants): median age: 38.0 years (range: 21 to 49 years); 18.2% men Control group (325 participants): median age: 39.0 years (range: 22 to 41 years); 22% men 			
	Target group: adults			
	Inclusion criteria			
	 Completed written informed consent process prior to undergoing any screening evaluations. Men or women aged ≥ 18 and ≤ 50 years on study day 0 In general good health, confirmed by medical history and physical examination Had ability to complete follow-up period as required by the protocol Had laboratory evidence of HIV infection, defined as a positive HIV-1 ELISA test plus a positive confirmatory test (e.g. a second HIV-1ELISA, PCR, or rapid ELISA) diagnosed prior to randomization. Was willing to allow the investigators to discuss the participant's medical history with the participant's HIV physician. If not receiving ART at the time of randomization, must have 2 CD4+ lymphocyte count test results > 350 cells/mm³, performed ≥ 4 weeks apart, 1 performed within 6 months prior to randomization and 1 within 45 days prior to randomization. If receiving ART at the time of randomization, must have 2 CD4+ lymphocyte count test results > 300 cells/mm³, performed ≥ 4 weeks apart, 1 performed within 6 months prior to randomization and 1 within 45 days prior to randomization. Participants on ART must have been receiving ART for ≥ 6 months prior to randomization and must have an undetectable HIV viral load within 45 days prior to randomization. Had: a negative QFT test result and tuberculin PPD skin test ≤ 5 mm induration within 45 days prior to randomization. 			
	 * a positive QFT test result or tuberculin PPD skin test > 5 mm (or both) and had completed ≥ 5 months of isoniazid preventive therapy within 3 years prior to randomization or * a positive QFT test result or tuberculin PPD skin test > 5 mm (or both) and had completed treatment for TB disease within 3 years prior to randomization. • Women: ability to avoid pregnancy during the trial. Women physically capable of pregnancy (not sterilized and still menstruating or within 1 year of the last menses if menopausal) in sexual relationships 			
	with men must have avoided pregnancy by using an acceptable method of avoiding pregnancy from 28 days prior to administration of the study vaccine to 6 months after the last study vaccination. Acceptable methods of avoiding pregnancy included a sterile sexual partner, sexual abstinence (not engaging in sexual intercourse), and any contraceptive method deemed clinically suitable by the trial clinician taking into account ART status.			
	 Had completed the written informed consent process for simultaneous enrolment in Aeras Vaccine Development Registry protocol. 			
	Exclusion criteria			
	Acute illness			
	 Fever (temperature > 37.5 °C) 			
	Significant symptomatic infection (including laboratory evidence of HIV-2)			

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Ndiaye 2015 (Continued)

- Any evidence of active TB disease, as determined by any clinical, radiological, or microbiology measurements.
- Any AIDS defining illness by WHO criteria
- Use of any investigational or non-registered drug, vaccine, or medical device other than the study vaccine within 182 days preceding dosing of study vaccine, or planned use during the study period
- Previous receipt of a recombinant MVA or FP vector at any time.
- Enrolled in any other clinical product trial
- Administration of methotrexate, azathioprine, cyclophosphamide, oral corticosteroids (for corticosteroids, this will mean prednisolone, or equivalent, ≥ 0.5 mg/kg/day; inhaled and topical steroids are allowed), and other immunosuppressive therapies, or blood products or blood derivatives within the 6 months prior to randomization
- History of allergic disease or reactions likely to be exacerbated by any component of the vaccine, e.g. egg products
- History of cancer (except basal cell carcinoma of the skin and cervical carcinoma in situ), or renal failure
- Severe depression, schizophrenia, or mania
- Pregnant, breast-feeding, or both
- History of anaphylaxis in reaction to vaccination
- Principal investigator assessment of lack of willingness to participate and comply with all requirements of the protocol, or identification of any factor felt to significantly increase the participant's risk of experiencing an adverse outcome

HIV status: positive

Other comorbidities: none reported

	Preterms: not mentioned		
Interventions	Intervention group		
	Vaccine: MVA85A/AERAS-85		
	 Dosage: 1 × 10⁸ pfu 		
	Route: intradermal.		
	Schedule: at day 1, and 2nd (booster) dose given 6 months after the 1st injection		
	Timing after BCG: not mentioned		
	Control group		
	Vaccine: Candida skin test antigen		
	Dosage: not mentioned		
	Route: intradermal		
	Schedule: at day 1, and 2nd (booster) dose given 6 months after the 1st injection		
	Timing after BCG: not mentioned		
Outcomes	Outcomes included in this review		
	Active TB		
	* Endpoint 1: culture or Xpert [®] MTB/RIF positivity		
	* Endpoint 2: endpoint 1 and a composite clinical endpoint; see detailed criteria in Table 4		
	* Endpoint 3: participants placed on treatment for TB by a health professional		
	Latent TB		
	Adverse effects of any severity		
	Serious adverse effects		
	Adverse events of any severity		
	Outcomes not included in this review		

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Ndiaye 2015 (Continued)

(continued)	Immunogenicity tests
Notes	Countries: South Africa and Senegal
	Setting: Cape Town (South Africa) and Dakar (Senegal), urban
	Background prevalence of TB
	 In Cape Town: TB case notification rate was at least 1500 per 100,000 population per year In Dakar: TB incidence rate of 0.14% in 2013
	Study dates: 4 August 2011 to 24 April 2013 for enrolment, with follow-up until 19 May 2014
	Study sponsor: Aeras. Collaborators: University of Oxford and European and Developing Countries Clin- ical Trials Partnership (EDCTP) (IP.2007.32080.002)

Funders: Bill & Melinda Gates Foundation, Wellcome Trust, and Oxford-Emergent Tuberculosis Consortium

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote from the report: "Participants were randomly assigned (1:1) in blocks of four by a randomly generated sequence of participant identification numbers via an interactive voice response system to receive two intradermal injections of either 1 × 10 ⁸ pfu MVA85A or placebo."
Allocation concealment (selection bias)	Low risk	Quote from the report: "A statistician uninvolved with study analyses prepared the interactive voice response system randomisation schedule."
		Comment: the interactive automated voice response system would make it impossible to predict the allocation sequence.
Blinding of participants and personnel (perfor- mance bias)	Low risk	Quote from the report: "Participants, nurses (who were involved in assessment and follow-up) investigators, and laboratory staff were masked to group allo- cation."
Alloutcomes		"Doses of vaccines were prepared and labelled in masked syringes."
		Quote from the protocol: "The MVA85A/AERAS-485 and the placebo will be packaged and labelled to appear indistinguishable from each other at the time of injection. Identical syringes and needles will be used for preparation and administration of injections of vaccine/placebo, and labels accompanying the syringes of prepared vaccine/placebo doses will not indicate which is in the sy- ringe."
Blinding of outcome as- sessment (detection bias): laboratory assessors	Low risk	Comment: as quoted above and outcome objective
Blinding of outcome as- sessment (detection bias): clinical assessors All outcomes	Low risk	Quote from the protocol in supplement: "The study vaccine manager and the study monitor will be the only persons unblinded at the site during the study and must not reveal individual subject treatment assignments to any other member of the study team. The study vaccine manager must be a designat- ed study team member who is not an employee of Aeras and who will have no other clinical or regulatory responsibilities associated with the conduct of the study during the entire study period. Unblinded study personnel must not par- ticipate in the evaluation of adverse events."

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Ndiaye 2015 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote from the report: "650 were randomly assigned; 649 were included in the safety analysis and 645 in the per-protocol analysis." Median follow-up for the 320 recipients of MVA85A was 655 days and for the 325 recipients of placebo was 654 days. "Other than 4 participants, all participants were included in the analysis."
		Comment: when authors refer to "per-protocol analysis," this is actually re- garding the analysis for the efficacy outcome. Results for per-protocol analy- ses were noted to be not different from the intention-to-treat results that were not reported.
Selective reporting (re- porting bias)	Unclear risk	Quote from the report: "Routine haematological and biochemical test results did not differ between study groups (data not shown)."
		Adverse effects solicited by the vaccine were not disaggregated by type of event.
Other bias	Unclear risk	Quote from the report: "The secondary outcome was the efficacy of MVA85A for the prevention of active tuberculosis in the per-protocol population which was determined by the incidence of active tuberculosis meeting the definition of endpoint 1, calculated as the number of new cases of active tuberculosis with a date of diagnosis from 28 days after the first vaccination until the end of the study follow-up (May 19, 2014)."
		Comment: the start of the intervention did not coincide with the start of fol- low-up; therefore a period of follow-up was excluded, and participants who experienced the outcome soon after intervention were missing from analyses. As such, the way in which outcomes were measured may bias effect estimates.
		This study was stopped early owing to data from the Tameris 2013 trial. As such, it was underpowered to measure efficacy outcomes.
		Quote: "Aeras was the trial sponsor and contributed to study design and data analysis."
		Comment: impact of sponsor involvement in analysis of results unclear

Nemes 2018

 Methods
 Study objective: to assess safety and immunogenicity of MVA85A vaccination in newborns of HIV-positive mothers, followed by selective deferred BCG vaccination at 8 weeks for HIV-negative infants.

 Study design: double-blind, randomized controlled trial
 Study duration: not mentioned

 Length of follow-up: 52 weeks
 Follow-up method

 •
 For safety endpoints: infants were monitored at weeks 1, 4, 6, and 8 after MVA85A/control vaccination and thereafter, at weeks 9, 12, and 16 (corresponding to weeks 1, 4, and 8 following delayed BCG vaccination at 8 weeks of age), and at week 52. Method of follow-up not detailed.

 •
 For immunogenicity analyses: blood was collected at weeks 4, 8, 16, and 52

 Losses to follow-up: 9 participants (3 in the intervention group, 6 in the control group)
 Power calculation: the sample size had 90% probability of detecting a serious adverse event with a true occurrence rate of 1.5% in infants receiving MVA85A vaccine and 80% power to detect a 15% differ

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Nemes 2018 (Continued)

	0.05).			
Participants	Number: 248			
	• Intervention group (123 participants): mean age: day of birth; 49% boys			
	Control group (125 participants): mean age: day of birth; 49% boys			
	Target group: infants of HIV-positive mothers			
	Inclusion criteria			
	 HIV-positive mother receiving either cART, or started on PMTCT prophylaxis Maternal antenatal and postnatal written informed consent Maternal age ≥ 18 years at the time of informed consent Infant age < 96 hours; any sex Infant birth and residence in the study area Mother contactable and able to attend follow-up visits 			
	Exclusion criteria			
	 Neonatal Apgar score < 7 at 5 minutes Infant birth weight < 2000 g or > 4500 g Estimated infant gestational age < 32 weeks Neonatal respiratory distress History or evidence of infant congenital abnormality, or immunosuppressive condition, other than hilly infaction 			
	 Any maternal or infant condition or systemic illness that in the opinion of the investigator was likely to affect safety or immunogenicity of study vaccine 			
	Infant BCG vaccination prior to enrolment			
	 Residence in a household, or frequent close contact, with an adult diagnosed with active TB who has not yet completed TB treatment 			
	Mother with active TB who has not yet completed TB treatment			
	Unknown or negative maternal HIV status			
	 Intention to leave the study area or unable to attend follow-up visits, or both 			
	HIV status: infants of HIV-positive mothers			
	 Intervention group * Mother receiving ARTs: 80% 			
	* Median maternal CD4 count: 442 cells/mm ³ (IQR 306 to 607)			
	Control group: * Mother receiving ARTs: 81%			
	* Median maternal CD4 count: 400 cells/mm ³ (IQR 262 to 554.5)			
	Other comorbidities: none reported			
	Preterms: median gestational age			
	Intervention group: 39 weeks (IQR 39 to 40)			
	Control group: 40 weeks (IQR 39 to 40)			
Interventions	Intervention group			
	 Vaccine: MVA85A Dosage: 1 × 10⁸ pfu Route: intradermal Schedule: 1 dose within 96 hours of birth 			

ence in the rate of non-serious adverse events (20% compared to 35%) between the 2 study groups (P <

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

Nemes 2018 (Continued)	
	• Timing after BCG: BCG 1–4 × 10 ³ cfu was selectively given at 8 weeks of age only to HIV-negative infants
	Control group
	 Vaccine: Candida skin test antigen (Candin[®])
	 Dosage: 1 × 10⁸ pfu
	Route: intradermal
	Schedule: 1 dose within 96 hours of birth
	• Timing after BCG: BCG 1–4×10 ⁵ cfu was selectively given at 8 weeks of age only to HIV-negative infants.
Outcomes	Outcomes included in this review:
	Active TB: culture-positive or on clinical/radiological grounds and TB contact history
	Latent TB: quantiFERON conversion at 1 year
	Adverse effects of any severity
	Serious adverse effects
	Adverse events of any severity
	Abnormal laboratory tests
	Outcomes not included in this review
	Immunogenicity tests
Notes	Country: South Africa
	Setting: urban (Cape Winelands east district and Khayelitsha)
	Background prevalence of TB: not mentioned
	Study dates: not reported. According to Clinicaltrial.gov, the study started in October 2012, and was completed in October 2015.
	Study sponsor: Aeras. Other funders: UK Medical Research Council, Department for International Devel- opment, and Wellcome Trust Joint Global Health Trials programme and AERAS.
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote from supplementary: "Assignment to study arm was double-blinded and based on a random number sequence prepared by an independent statis- tician."
Allocation concealment (selection bias)	Low risk	Quote from supplementary: "The study pharmacist, the only unblinded mem- ber of the study team, controlled the numbered sealed envelopes containing randomization arm and sequential 3-digit enrolment number."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Comment: from the statement that the pharmacist was the only unblinded member in the team we assumed everyone else was blinded and it was effec- tive.
Blinding of outcome as- sessment (detection bias): laboratory assessors	Low risk	Comment: from the statement that the pharmacist was the only unblinded member in the team we assumed everyone else was blinded and it was effec- tive.
Blinding of outcome as- sessment (detection bias): clinical assessors	Low risk	Comment: from the statement that the pharmacist was the only unblinded member in the team we assumed everyone else was blinded and it was effec- tive.

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Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: minimal attrition and balance between groups; 16 in MVA85A group and 19 in control group as set out in figure 1b.
Selective reporting (re- porting bias)	Low risk	Comment: reported everything they set out in protocol. Additionally reported QFT conversion and incident TB disease; outcomes were not specified in pro- tocol but of importance to mention.
Other bias	Unclear risk	Comment: authors declared no conflict of interest. 1 author declared that they were patent holders for MVA85A and were responsible for its development in Scriba 2011.

Scriba 2011				
Methods	Study objective: to assess the safety of and to characterize the T-cell response induced by 3 doses of the candidate vaccine, MVA85A, in BCG-vaccinated infants from a setting where TB was endemic.			
	Study design: open-label, Phase 2a safety, immunogenicity, and dose-finding study			
	Study duration: 23 months			
	Length of follow-up: 168 days (24 weeks)			
	Follow-up method			
	 Diary cards the first 7 days for registration of local and systemic adverse effects Onsite safety data at 60 minutes and on days 2, 7, 28, 84, and 168 Blood sample for haematology and biochemistry on days 7 and 84 Blood sample for immunogenicity on days 0, 7, 28, 84, and 168 			
	Losses to follow-up: none			
	Power calculation: not mentioned			
Participants	Number: 144			
	 Intervention group Vaccine group 1 (36 participants): median age: 270.5 days; 42% male Vaccine group 2 (36 participants): median age: 278.5 days; 47% male Vaccine group 3 (36 participants): median age: 188 days; 39% male Control group (36 participants): median age: 252 days; 62% male 			
	Target group: infants aged 5–12 months			
	Inclusion criteria			
	 Children or infants aged 6 months to 11 years Participant's parent/guardian willing and able to give written informed consent for participation in the study Participant is BCG vaccinated within the first 4 weeks of life Informed assent from all children aged ≥ 7 years unless judged incapable of understanding the basic concepts covered in the informed assent form, and from children aged < 7 years if judged capable of understanding the basic concepts covered in the informed assent form and from children aged < 7 years if judged capable of understanding the basic concepts covered in the informed assent form Healthy Clinically acceptable laboratory results from screening visit 			
	 Chest x-ray normal with no evidence of active or past TB 			

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Scriba 2011 (Continued)

Trusted evidence. Informed decisions. Better health.

	 Participant's parent/legal guardian willing to allow child to undergo an HIV test Parent/guardian and participant able (in the Investigators opinion) and willing to comply with all study requirements
	Exclusion criteria
	 Participant Mantoux (> 10 mm) or ELISPOT (> 50 spots/million PBMC) positive for Mycobacterium tu- berculosis (PPD, ESAT-6 or CFP-10, or both)
	 HIV-positive Any other significant disease or disorder which, in the opinion of the investigator, may either put the person at risk because of participation in the study, or may influence the result of the study, or the person's ability to participate in the study
	Have participated in another research study involving an investigational product in the past 12 weeks
	 Previously enrolled into this study Received a live vaccine (e.g. measles) in the previous 4 weeks or due to receive a live vaccine in the 4 weeks following enrolment
	HIV status: negative
	Other comorbidities: none reported
	Preterms: not mentioned
Interventions	Intervention group
	 Vaccine: MVA85A (manufactured at Impfstoffwerk Dessau-Tornau; Biologika) Dosage:
	• * Vaccine group 1: 2.5×10^7 pfu in 35μ L
	* Vaccine group 2: 5×10^7 pfu in 70 µL * Vaccine group 2: 1×10^8 s fu in 125 vL
	^{**} Vaccine group 3: $1 \times 10^{\circ}$ ptu in 135 µL
	 Roule: intrademation decod ann Schedule: at day 1, 1 dose
	 Timing after BCG: inclusion criteria request BCG given during the first 4 weeks of life.
	Control group
	Vaccine: pneumococcal 7 valent conjugate (Prevenar, Wyeth)
	Dosage: not specified
	Route: intramuscular, site of injection not mentioned
	 Schedule: at day 1, 1 dose Timing after BCG: inclusion criteria request BCG given during the first 4 weeks of life.
Outcomes	Outcomes included in this review
	 Latent TB (reported under the safety profile) Adverse effects of any severity Serious adverse effects Adverse events of any severity Abnormal biochemical tests
	Outcomes not included in this review
	Immunogenicity tests
Notes	Country: South Africa
	Setting: Cape Town, urban
	Background prevalence of TB: extremely high (incidence of 1%)

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Scriba 2011 (Continued)

Study dates: February 2008 to December 2009 (according to data published in clinicaltrial.gov, not mentioned in the paper)

Study sponsor: University of Oxford. Funders: EuropeAID European commission, Wellcome trust

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	High risk	Quote from the report: "The aim was to enroll 144 infants into 3 consecutive vaccine dose groups of 48, who would be systematically allocated at a 3:1 ratio to receive either MVA85A (groups 1–3) or placebo." Comment: randomization method predictable.
Allocation concealment (selection bias)	High risk	Quote from the report: "systematically allocated at a 3:1 ratio"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote from report: "MVA85A, contract manufactured at Impfstoffwerk Dessau- Tornau (Biologika), was administered intradermally over the deltoid region of the arm contralateral to where BCG was administered." Prevenar was adminis- tered intramuscularly.
		Comment: open label with 2 different routes of administration. Subjective out- comes, so could influence participants when reporting the symptoms.
Blinding of outcome as- sessment (detection bias): laboratory assessors	Low risk	Comment: open label, but with no repercussion on objective laboratory out- comes.
Blinding of outcome as- sessment (detection bias): clinical assessors All outcomes	High risk	Comment: open label, with high repercussion on subjective clinical outcomes.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No attrition
Selective reporting (re- porting bias)	Low risk	Comment: all the outcomes mentioned in the methods section were reported in the results.
Other bias	Unclear risk	Quote from report: "are named inventors on a composition of matter patent for MVA85A filed by the University of Oxford and are shareholders in a joint venture formed for the further development of this vaccine."
		Comment: unknown role of funders in the elaboration of the study and 2 au- thors with potential conflict of interest.

Tameris 2013

Methods

Study objective: to assess safety, immunogenicity, and efficacy of MVA85A against TB and *Mycobacterium tuberculosis* infection in infants.

Study design: parallel-group, randomized, placebo-controlled double-blind Phase 2b trial

Study duration: 39 months

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Tameris 2013 (Continued)

Length of follow-up: ≥ 15 months after enrolment, and up to 39 months

Follow-up method

- Follow-up at study day 7, study day 28, study day 84, and every 84 days (i.e. every 3 months) thereafter until the end of the study.
- Safety diary cards for first 7 days, direct questioning at study days 7 and 28 and serious adverse events throughout the study.
- Peripheral blood for routine haematological and biochemical tests at screening and on days 7 and 28 after vaccination in an initial safety cohort.
- QFT testing at screening, day 336, at end of study visit, and for infants admitted to a dedicated study ward for investigation for TB.
- Active follow-up every 3 months to identify signs, symptoms, or exposure that merited further investigation.

Losses to follow-up

- Intervention group: 61/1399 (4.4%) participants; 37 (2.6%) withdrew consent
- Control group: 65/1398 (4.6%) participants; 25 (1.8%) withdrew consent

Power calculation: given a TB cumulative incidence of 3% over 18 months in the control group, 1392 participants per treatment group (2784 participants total) would be required to demonstrate positive efficacy when the true efficacy of MVA85A/AERAS-485 was approximately 60%. An estimate of 7.5% of participants lost to follow-up in each treatment group was assumed over 18 months.

Participants Number: 2797

- Intervention group (1399 participants): mean age: 146.6 days; 50.6% boys
- Control group (1395 participants; 1398 randomized): mean age: 145.7; 51.2% boys

Target group: infants aged 4-6 months

Inclusion criteria

- Healthy infants aged 4–6 months
- Received BCG vaccination within 7 days of birth
- Received all age-appropriate routine immunizations, and 2 doses of pneumococcal conjugate vaccine ≥ 28 days before study vaccination (amended to 14 days during enrolment)
- HIV ELISA-negative
- QuantiFERON-negative
- No substantial exposure to a person with known TB
- Written informed consent obtained from parents/guardian
- Weight: by chart > 3rd percentile on study day 0 or, if < 3rd percentile, infant showed a stable growth pattern
- Ability to complete follow-up period as required by the protocol
- Completed simultaneous enrolment in the Aeras Vaccine Development Registry protocol

Exclusion criteria

- Acute illness on study day 0
- Fever ≥ 37.5 °C on study day 0
- Evidence of significant active infection on study day 0
- Received a EPI immunization within 14 days prior to study day 0
- Historical or virological evidence of individual or maternal HIV-1 infection
- History of allergic disease or reactions likely to be exacerbated by any component of the study vaccine
- Previous medical history, or evidence, of an intercurrent illness that may compromise the safety of the infant in the study
- · Evidence of chronic hepatitis from any cause

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Tameris 2013 (Continued)	 History or evidence of any systemic disease on physical examination or any acute, chronic, or intercurrent illness that, in the opinion of the investigator, may have interfered with the evaluation of the safety or immunogenicity of the vaccine History of or known TB or treatment for TB Shared residence since birth with a person with active TB or on ATT for < 2 months
	HIV status: negative
	Other comorbidities: none reported
	Preterms
	 Intervention group: 412 (29.4%) participants Control group: 368 (26.4%) participants
Interventions	Intervention group
	 Vaccine: MVA85A/AERAS-485 Dosage: 1 × 10⁸ pfu in 0.06 mL Route: intradermal Schedule: at day 1, 1 dose Timing after BCG: inclusion criteria request BCG given during the first 7 days of life.
	Control group
	 Vaccine: Candida skin test antigen (Candin, AllerMed, USA) Dosage: 0.06 mL Route: intradermal Schedule: at day 1, 1 dose Timing after BCG: inclusion criteria request BCG given during the first 7 days of life.
Outcomes	Outcomes included in this review
	 Active TB Endpoint 1: see detailed criteria in Table 2 Endpoint 2: participants diagnosed with TB based on the presence of specific clinical, radiological, and microbiological findings. Endpoint 3: participants placed on treatment for TB by a health professional Latent TB Adverse effects of any severity Serious adverse effects Adverse events of any severity
	Outcomes not included in this review
	Immunogenicity tests
Notes	Country: South Africa
	Setting: rural, near Cape Town
	Background prevalence of TB: extremely high. The overall incidence of TB in South Africa in 2011 was estimated to be almost 1%, and the incidence of TB in children aged < 2 years was about 3% at the trial site.
	Study dates: enrolment 15 July 2009 to 4 May 2011 and follow-up to 25 October 2012
	Study sponsor: Aeras. Collaborators: University of Oxford and University of Cape Town. Funders: Aeras, Wellcome trust and Oxford Emergent tb consortium (OETC)

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Tameris 2013 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote from the report: "We randomly allocated infants in a 1.1 ratio with a block size of 4 using interactive voice /online response system."
		"An independent statistician prepared the randomisation schedule."
Allocation concealment (selection bias)	Low risk	Comment: voice response system adequately concealed allocation of inter- vention.
Blinding of participants and personnel (perfor- mance bias)	Low risk	Quote from the report: "Parents or legal guardians of study participants, study staff administering vaccine or undertaking follow up clinical assessments and laboratory staff were masked to intervention group assignment."
All outcomes		"Doses were prepared and labelled in masked syringes by an unmasked study pharmacist."
		Comment: syringes had equal amount of placebo and control.
		Quote from the protocol: "packaged and labelled to appear indistinguishable to each other."
Blinding of outcome as- sessment (detection bias): laboratory assessors	Low risk	Comment: laboratory staff were masked to intervention group assignment.
Blinding of outcome as- sessment (detection bias): clinical assessors All outcomes	Low risk	Comment: staff undertaking clinical assessments were masked to intervention group assignment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote from the report: "The number of participants discontinuing the study did not differ between the two treatment groups." 1126 infants (5%) were lost to follow-up, 11 died (< 1%), and 62 (2%) had consent withdrawn.
		Comment: reasons for missing outcome data balanced between the 2 groups and proportion of missing data not enough to have a clinically relevant impact on the intervention effect estimate. Per-protocol analysis was done and only 1 person was excluded from analysis from the placebo group due to dose devia- tion.
Selective reporting (re- porting bias)	Unclear risk	Quote from study description from clinical trials.gov: "Adverse events and clin- ically relevant laboratory results for the safety cohort will be summarized to examine the relationship between treatment group and key safety endpoints including number (percentage) of solicited and spontaneous adverse events, rates of reactogenicity, and number (percentage) of subjects with newly ab- normal post-vaccination laboratory values based on predefined neonatal toxi- city criteria."
		Comment: data were collected; however, no summary provided on biochemi- cal or haematological adverse effects.
		Unclear if endpoints were specified a priori as endpoint definition was only published alongside the trial and approach outlined a priori on clinical trial registry was amended.
		The differences between endpoint point 1 and 2 were 5 mm on TST; 2 posi- tive smears compared to 1 positive smear and residence in household with positive AFB member. These endpoints were significantly different from the

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Tameris 2013 (Continued)		endpoints used in 2 other trials that included efficacy measures (Ndiaye 2015; Nemes 2018).
Other bias	Unclear risk	Quote from the report: "Aeras was the trial sponsor. Aeras and the Ox- ford-Emergent Tuberculosis Consortium (OETC) contributed to study design, data interpretation, and writing of the manuscript."
		Comment: impact of sponsor involvement on study findings unclear.

AFB: acid-fast bacilli; ART: antiretroviral therapy; ATT: antituberculosis therapy; BCG: Bacillus Calmette-Guérin; cART: combination antiretroviral therapy; CFP-10: culture filtrate protein-10; cfu: colony-forming unit; ELISA: enzyme-linked immunosorbent assay; ELISPOT: enzyme-linked immune absorbent spot; EPI: Expanded Programme on Immunization; ESAT-6: early secretory antigenic-6; FP: floating point; IQR: interquartile range; MVA: modified Vaccinia Ankara; PBMC: peripheral blood mononuclear cell; PCR: polymerase chain reaction; pfu: plaque-forming unit; PMTCT: prevention of mother-to-child transmission; PPD: purified protein derivative; QFT: QuantiFERON-TB Gold In-Tube; SD: standard deviation; TB: tuberculosis; TST: tuberculin skin test; WHO: World Health Organization.

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Brookes 2008	Different study design
Bunyasi 2015	Different outcomes measured
Dieye 2013	Different study design
Harris 2011	Different study design
Harris 2014a	Different study design
Hawkridge 2008	Different study design
Matsumiya 2014a	Different outcomes measured
Matsumiya 2014b	Different outcomes measured
Matsumiya 2014c	Different outcomes measured
McShane 2004	Different study design
Meyer 2013	Different study design
Minassian 2011	Different study design
Minhinnick 2016	Different study design
Mulenga 2015	Different intervention
Odutola 2012	Different study design
Ota 2011	Different study design
Pathan 2007	Different study design
Pathan 2012	Different study design

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Study	Reason for exclusion
Rowland 2012	Different study design
Rowland 2013	Different study design
Sander 2009	Different study design
Satti 2014	Different study design
Scriba 2010	Different study design
Scriba 2012	Different study design
Sheehan 2015	Different study design
Tameris 2014	Measured different outcomes
Tanner 2014	Different study design
Whelan 2009	Different study design

DATA AND ANALYSES

Comparison 1. MVA85A versus placebo

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Active tuberculosis (TB): con- firmed by culture or Xpert® MTB/RIF longest reported fol- low-up	2	3439	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.58, 1.62]
2 Active TB: started on TB treatment	3	3687	Risk Ratio (M-H, Fixed, 95% CI)	1.10 [0.92, 1.33]
3 Latent TB	4	3831	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.85, 1.21]
4 Adverse effects of any sever- ity	3		Risk Ratio (M-H, Fixed, 95% CI)	Totals not selected
4.1 Local: skin	3		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
4.2 Malaise	1		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
4.3 Lethargy	1		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
4.4 Any fever	1		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
4.5 Vomiting	1		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]

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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5 Adverse effects of any severi- ty: aggregated	4		Risk Ratio (M-H, Fixed, 95% CI)	Totals not selected
6 Serious adverse effects	3	3692	Risk Difference (M-H, Fixed, 95% CI)	0.00 [-0.00, 0.00]
7 Adverse events of any sever- ity	4	3836	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [1.02, 1.08]
8 Abnormal biochemical tests	2	392	Risk Ratio (M-H, Fixed, 95% CI)	1.09 [0.60, 1.97]

Analysis 1.1. Comparison 1 MVA85A versus placebo, Outcome 1 Active tuberculosis (TB): confirmed by culture or Xpert[®] MTB/RIF longest reported follow-up.

Study or subgroup	MVA85A	Placebo		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-H	, Fixed, 95%	CI			M-H, Fixed, 95% CI
Ndiaye 2015	6/320	9/325		-				30.84%	0.68[0.24,1.88]
Tameris 2013	22/1399	20/1395						69.16%	1.1[0.6,2]
Total (95% CI)	1719	1720			•			100%	0.97[0.58,1.62]
Total events: 28 (MVA85A), 29 (Placebo	o)								
Heterogeneity: Tau ² =0; Chi ² =0.64, df=1	L(P=0.42); I ² =0%								
Test for overall effect: Z=0.13(P=0.9)									
		Favours MVA85A	0.01	0.1	1	10	100	Favours placebo	

Favours MVA85A 0.01 0.1

¹⁰ ¹⁰⁰ Favours placebo

Analysis 1.2. Comparison 1 MVA85A versus placebo, Outcome 2 Active TB: started on TB treatment.

Study or subgroup	MVA85A	Placebo		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-	H, Fixed, 95%	6 CI			M-H, Fixed, 95% Cl
Ndiaye 2015	8/320	9/325			+			4.72%	0.9[0.35,2.31]
Nemes 2018	5/123	3/125						1.57%	1.69[0.41,6.93]
Tameris 2013	196/1399	177/1395			+			93.71%	1.1[0.91,1.33]
Total (95% CI)	1842	1845			•			100%	1.1[0.92,1.33]
Total events: 209 (MVA85A), 189 (Placebo)									
Heterogeneity: Tau ² =0; Chi ² =0.53, d	f=2(P=0.77); I ² =0%								
Test for overall effect: Z=1.05(P=0.2	9)								
		Favours MVA85A	0.01	0.1	1	10	100	Favours BCG alone	

Analysis 1.3. Comparison 1 MVA85A versus placebo, Outcome 3 Latent TB.

Study or subgroup	MVA85A	Placebo		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-H, Fixed, 95% CI				M-H, Fixed, 95% CI	
Ndiaye 2015	38/320	40/325		1	+	1		18.4%	0.96[0.64,1.46]
		Favours MVA85A	0.01	0.1	1	10	100	Favours placebo	

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Study or subgroup	MVA85A	Placebo			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H	l, Fixed, 95%	6 CI			M-H, Fixed, 95% CI
Nemes 2018	1/123	4/125	_					1.84%	0.25[0.03,2.24]
Scriba 2011	3/108	0/36						0.35%	2.38[0.13,44.93]
Tameris 2013	178/1399	171/1395			+			79.41%	1.04[0.85,1.26]
Total (95% CI)	1950	1881			•			100%	1.01[0.85,1.21]
Total events: 220 (MVA85A), 215 (Pla	icebo)								
Heterogeneity: Tau ² =0; Chi ² =1.98, df	f=3(P=0.58); I ² =0%								
Test for overall effect: Z=0.16(P=0.87	7)								
		Favours MVA85A	0.01	0.1	1	10	100	Favours placebo	

Analysis 1.4. Comparison 1 MVA85A versus placebo, Outcome 4 Adverse effects of any severity.

Study or subgroup	MVA85A	Placebo	Risk Ratio	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% Cl	M-H, Fixed, 95% CI
1.4.1 Local: skin				
Nemes 2018	121/123	118/125	•	1.04[0.99,1.09]
Scriba 2011	106/108	6/36		5.89[2.84,12.23]
Tameris 2013	1251/1399	628/1396	+	1.99[1.87,2.11]
1.4.2 Malaise				
Scriba 2011	6/108	1/36		2[0.25,16.06]
1.4.3 Lethargy				
Scriba 2011	6/108	2/36		1[0.21,4.74]
1.4.4 Any fever				
Scriba 2011	18/108	2/36	+	3[0.73,12.3]
1.4.5 Vomiting				
Scriba 2011	6/108	2/36		1[0.21,4.74]
		Favours MVA85A	0.01 0.1 1 10	¹⁰⁰ Favours placebo

Analysis 1.5. Comparison 1 MVA85A versus placebo, Outcome 5 Adverse effects of any severity: aggregated.

Study or subgroup	MVA85A	Placebo		Risk Ratio				Risk Ratio	
	n/N	n/N		M-H, F	ixed, 959	% CI		M-H, Fixed, 95% CI	
Ndiaye 2015	318/324	307/325			+			1.04[1.01,1.07]	
Nemes 2018	105/123	30/125						3.56[2.58,4.9]	
Scriba 2011	106/108	6/36					\rightarrow	5.89[2.84,12.23]	
Tameris 2013	1251/1399	628/1396				+	1	1.99[1.87,2.11]	
		Favours MVA85A	0.2	0.5	1	2	5	Favours placebo	

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Analysis 1.6. Comparison 1 MVA85A versus placebo, Outcome 6 Serious adverse effects.

Study or subgroup	MVA85A	Placebo		Ris	sk Differend	e	Weight Risk Di		Risk Difference
	n/N	n/N		M-H	, Fixed, 95%	6 CI			M-H, Fixed, 95% Cl
Ndiaye 2015	1/324	0/325			•			17.58%	0[-0.01,0.01]
Nemes 2018	0/123	0/125			+			6.72%	0[-0.02,0.02]
Tameris 2013	0/1399	1/1396						75.7%	-0[-0,0]
Total (95% CI)	1846	1846						100%	0[-0,0]
Total events: 1 (MVA85A), 1 (Placebo)								
Heterogeneity: Tau ² =0; Chi ² =1.01, df	=2(P=0.6); I ² =0%								
Test for overall effect: Z=0(P=1)									
		Favours MVA85A	-1	-0.5	0	0.5	1	Favours placebo	

Analysis 1.7. Comparison 1 MVA85A versus placebo, Outcome 7 Adverse events of any severity.

Study or subgroup	MVA85A	Placebo	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% Cl		M-H, Fixed, 95% Cl
Ndiaye 2015	321/324	312/325	-	20.84%	1.03[1.01,1.06]
Nemes 2018	122/123	121/125	+	8.03%	1.02[0.99,1.06]
Scriba 2011	1/36	1/12		0.1%	0.33[0.02,4.93]
Scriba 2011	6/36	0/12		0.05%	4.57[0.28,75.58]
Scriba 2011	3/36	0/12		0.05%	2.46[0.14,44.48]
Tameris 2013	1120/1399	1059/1396	-	70.93%	1.06[1.01,1.1]
Total (95% CI)	1954	1882	•	100%	1.05[1.02,1.08]
Total events: 1573 (MVA85A), 149	3 (Placebo)				
Heterogeneity: Tau ² =0; Chi ² =5.76	5, df=5(P=0.33); I ² =13.17%				
Test for overall effect: Z=3.29(P=0))				
		1	Favours placebo		

Analysis 1.8. Comparison 1 MVA85A versus placebo, Outcome 8 Abnormal biochemical tests.

Study or subgroup	MVA85A	Placebo			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H	, Fixed, 95% (1			M-H, Fixed, 95% Cl
Nemes 2018	14/123	13/125			-			68.25%	1.09[0.54,2.23]
Scriba 2011	13/108	4/36						31.75%	1.08[0.38,3.11]
Total (95% CI)	231	161			•			100%	1.09[0.6,1.97]
Total events: 27 (MVA85A), 17 (Placebo)								
Heterogeneity: Tau ² =0; Chi ² =0, df=1(P=	:0.99); l ² =0%								
Test for overall effect: Z=0.29(P=0.77)									
		Favours MVA85A	0.01	0.1	1	10	100	Favours BCG alone	

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Comparison 2. Comparison of endpoints

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Tameris 2013: incidence of tuberculosis (TB) according to post-hoc endpoints	1		Risk Ratio (M-H, Fixed, 95% CI)	Totals not select- ed
2 Ndiaye 2015: incidence of TB according to post hoc defined endpoints	1		Risk Ratio (M-H, Fixed, 95% CI)	Totals not select- ed

Analysis 2.1. Comparison 2 Comparison of endpoints, Outcome 1 Tameris 2013: incidence of tuberculosis (TB) according to post-hoc endpoints.

Study or subgroup	or subgroup MVA85A		Risk Ratio	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl	
Tameris 2013	32/1399	39/1395		0.82[0.52,1.3]	
Tameris 2013	55/1399	52/1395		1.05[0.73,1.53]	
Tameris 2013	196/1399	177/1395		1.1[0.91,1.33]	
Tameris 2013	22/1399	20/1395		1.1[0.6,2]	
		Favours MVA85A	0.5 0.7 1 1.5 2	Favours placebo	

Analysis 2.2. Comparison 2 Comparison of endpoints, Outcome 2 Ndiaye 2015: incidence of TB according to post hoc defined endpoints.

Study or subgroup	p MVA85A		Placebo					Risk Ratio	
	n/N	n/N		М-Н,	Fixed, 959	% CI		M-H, Fixed, 95% Cl	
Ndiaye 2015	6/320	9/325		_	-+			0.68[0.24,1.88]	
Ndiaye 2015	6/320	9/325		-	-+			0.68[0.24,1.88]	
Ndiaye 2015	8/320	9/325						0.9[0.35,2.31]	
		Favours MVA85A	0.01	0.1	1	10	100	Favours placebo	

ADDITIONAL TABLES

Table 1. Summary of Phase 1 studies

NCT trial number	Route	Dates	Intervention and schedule de- tails	Country	Partici- pants (age)	HIV	Adverse events	Reference
NCT00423566	ID	2002–2003	MVA85A; 1 dose	UK	14 adults (18–45 years)	-ve	7 trials (112 partic- ipants); combined in 1 report: no seri- ous AE attributable to the vaccine	McShane 2004; Rowland 2012
NCT00423839	ID	2003-2005	MVA85A; 1 dose, 2 doses (5 × 10 ⁷ pfu)	Gambia	21 adults	NR	No serious AE attrib- utable to the vaccine	Brookes 2008; Ibanga 2006; Owiafe 2012
NCT00427830	ID	2003–2005	MVA85A; 1 dose (5 × 10 ⁷ pfu)	UK	21 adults	-ve	No serious AE attrib- utable to the vaccine	McShane 2004; Pathan 2012; Rowland 2012; Tanner 2014; Whelan 2009
NCT00427453	ID	2003-2005	MVA85A; 1 dose (5 × 10 ⁷ pfu)	UK	10 adults	-ve	No serious AE attrib- utable to the vaccine	Pathan 2012; Rowland 2012
NCT00456183	ID	2005–2007	MVA85A, (5 × 10 ⁷ pfu)	UK	12 adults with latent tuberculosis	-ve	No vaccine-related serious AEs 7 trials (112 partic- ipants; data com- bined in 1 report)	Rowland 2012; Sander 2009; Tanner 2014
NCT00465465	ID	2005–2007	MVA85A; 1 dose (1 × 10 ⁸ pfu for 12 participants, and 1 × 10 ⁷ pfu for 12 participants)	UK	24 adults	-ve	No serious AE attrib- utable to the vaccine	Griffiths 2011; Matsumiya 2013; Pathan 2007; Rowland 2012
NCT00460590	ID	2005–2008	MVA85A (5 × 10 ⁷ pfu)	South Africa	36 adults and adoles- cents	-ve	No vaccine-related serious AEs	Hawkridge 2008; Scriba 2010; Tameris 2014; Tanner 2014

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NCT00480454	ID	2006–2009	MVA85A;	The Gambia	214 infants	NR	No serious AE judged	Odutola 2012;
			1 dose MVA85A (2.5 × 10 ⁷ pfu, 5 × 10 ⁷ pfu)		(4 montins)		vaccine	018 2011
			Groups					
			 EPI vaccines: MVA85A + EPI: MVA85A + EPI 1 week later 					
NCT00395720	ID	2006-2010	MVA85A; 1 dose (5 × 10 ⁷ pfu for 10 participants, and 1 × 10 ⁸ pfu for 10 participants)	UK	20 adults	+ve	No serious AE attrib- utable to the vaccine	Minassian 2011
NCT00480558	ID	2007–2011	MVA85A; 1 dose (5 × 10 ⁷ pfu)	South Africa	48 adults	+ve	No vaccine-related	Scriba 2012;
			4 groups with background of		(18–50 years)		Serious AES	Tameris 2014;
			• MTB					
			• HIV					
			HIV on ART					
NCT00653770	ID	2007–2010	FP85A, MVA85A (5 × 10 ⁷ pfu)	UK	31 adults	-ve	No serious AE attrib- utable to the vaccine	Rowland 2013
NCT00548444	ID	2007–2010	MVA85A; 1 dose	UK	12 adults	-ve	7 trials (112 partic-	Porter (unpub-
			(1 × 10 ⁸ pfu), administered as 2 injections (5 × 10 ⁷ pfu each injec- tion)				bined in 1 report: no serious AE attribut- able to the vaccine	source Rowland 2012)
NCT00731471	ID	2008–2011	MVA85A; 2 doses (spaced by 6–12 months) (1 × 10 ⁸ pfu)	Senegal	24 adults	+ve	No serious AE attrib- utable to the vaccine	Dieye 2013
NCT01181856	ID	2010-2011	MVA85A; 1 dose (1 × 10 ⁸ pfu)	UK	24 adults	-ve	No serious AE attrib-	Matsumiya
	IM						utable to the vaccine	2013; Meyer 2013
NCT01194180	ID	2010-2012	MVA85A, BCG;	UK	49 adults re-	-ve	No serious AE attrib-	; Harris 2014b;
			1 dose (1 × 10 ⁸ pfu)		cruited; 48 completed study		utable to the vaccine	Matsumiya 2013

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	Table 1. Sum	mary of Phas	e 1 studies (Co	ntinued)					
				Group A: BCG naive, no MVA85A Group B: BCG naive, MVA85A					
•				Group C: BCG vaccinated, no MVA85A					
				Group D: BCG vaccinated, MVA85A.					
	NCT01497769	Aerosol ID	2011-2013	MVA85A; 1 dose: 1 × 10 ⁸ , 1 × 10 ⁷ pfu	UK	24 adults	-ve	No vaccine related serious adverse ef- fects.	Satti 2014
	NCT01683773	ID	2012-2014	AERAS-402 MVA85A;	UK	40 adults	-ve	No vaccine related	Sheehan 2015
				Group A: 2 doses AERAS-402 then MVA85A				Serious AES	
				Group B: 1 dose AERAS-402 then MVA85A					
	NCT01879163	ID	2013-2014	MVA85A IMX313;	UK	30 BCG vac-	-ve	No vaccine-related	Minhinnick
				Group A: low-dose MVA85A- IMX313 (1 × 10 ⁷ pfu)		adults		SELIOUS AE	2010
				Group B: dose MVA85A-IMX313 (5 × 10 ⁷ pfu)					
				Group C: MVA85A (5 × 10 ⁷ pfu)					
	NCT01829490	IM	2013–2016	MVA85A, ChAdOx1 85A;	UK	42 adults	-ve	No data reported yet	No publication
				Group A: 1 dose ChAdOx1 85A					NCT01829490
				Group B: 1 dose ChAdOx1 85A then MVA85A					
				Group C: 2 doses ChAdOx1 85A then MVA85A (1 × 10 ⁸ pfu)					
	NCT01954563	Aerosol	2013-2016	MVA85A;	UK	37 adults	-ve	No data reported yet	Manjaly 2016
		ID		Group 1: aerosol then ID					(conference ab-
				Group 2: ID then aerosol					SUDCI

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Table 1. Sum	mary of Phas	e 1 studies (Co	ontinued) Group 3: ID then ID (5 × 10 ⁷ pfu)						
NCT02532036	Aerosol ID	2015-2018	MVA85A; 1 × 10 ⁷ pfu aerosol in- haled,	UK	15 adults	-ve	No data reported yet	NCT02532036	
			5 × 10 ⁷ aerosol and ID						

-ve: negative; +ve: positive; AE: adverse event; ART: antiretroviral therapy; BCG: bacillus Calmette-Guérin; EPI: Expanded Programme on Immunization; ID: intradermal; IM: intramuscular; MTB: Mycobacterium tuberculosis; NR: not reported; pfu: plaque-forming unit.

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Table 2. Adverse events risk of bias assessment methods

Criterion	Assessment	Explanation
Participant-reported sy	mptoms	
Was monitoring active or passive?	Active Passive Unclear	We classified monitoring as 'active' when authors reviewed participants at set time points and enquired about symptoms.
Was blinding for partic- ipants and outcome as- sessors adequate?	Adequate Inadequate Unclear	We classified blinding as 'adequate' when both participants and outcome as- sessors were blinded to the intervention group, and the methods of blinding (including use of a placebo) were described.
Was outcome data re- porting complete or in- complete?	Complete Incomplete	We classified outcome data reporting as 'complete' when data were presented for all the time points where it was collected.
Were all participants in- cluded in reporting?	Yes No	We reported the percentage of randomized participants included in adverse event reporting.
Was the analysis inde- pendent of study spon- sor?	Yes No Unclear	We classified the analysis of trials sponsored by pharmaceutical companies as independent of the sponsor when it was clearly stated that the sponsor had no input to the trial analysis
Laboratory tests		
Number of tests under- taken	_	We extracted the type and number of laboratory tests were taken.
Timing of tests: was number and timing of tests adequate?	Adequate Inadequate	We classified the number and timing of tests as 'adequate,' when tests were taken at baseline, plus 2 other time points within the first week after treat- ment, plus the last day of the study. We classified the number of test taken as 'inadequate,' if either the laboratory controls in the first week or controls at 4 weeks were not performed.
Reporting of test re- sults: was reporting of test results complete?	Complete Incomplete	We classified reporting as 'complete' when test results of all time points were reported. For the trials with inadequate number of tests taken, we considered completeness of reporting as inconsequential, and therefore did not record a judgement.
Independence of da- ta analysis: was data analysis independent?	Yes No Unclear	We classified the analysis of trials sponsored by pharmaceutical companies as independent of the sponsor when it is clearly stated that the sponsor had no input to the trial analysis.

Adapted from Bukirwa 2014.

Table 3.	Differences in tuberculosis endpoint assessment							
Study	Endpoint 1	Endpoint 2	Endpoint 3					

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

All participants

placed on treat-

ment for TB by a

health profession-

al with the intent of

treating TB regard-

less of whether they

have met the other

efficacy endpoints.

Table 3. Differences in tuberculosis endpoint assessment (Continued)

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Tameris 2013

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Any of the following criteria. • Isolation of *M tuberculosis* from any • Isolation define TB infection ar

- Identification *of M tuberculosis* by an approved molecular diagnostic technique from any site.
- Histopathology diagnostic for TB disease (e.g. caseating granulomas).
- Choroidal tubercle diagnosed by an ophthalmologist.
- Miliary pattern on chest x-ray in an HIV-negative infant.
- Clinical diagnosis of TB meningitis (CSF protein concentrations > 0.6 g/L and pleocytosis of > 50 cells/ µL with > 50% mononuclear cells) with features of basal meningeal enhancement and hydrocephalus on head CT.
- Vertebral spondylosis.
- 1 smear or histology specimen positive for auramine-positive bacilli from a normally sterile body site.
- 1 of each of the following:
 - * evidence of mycobacterial infection defined as 2 acid-fast positive smears (each from a separate collection) that were morphologically consistent with mycobacteria from either sputum or gastric aspirate that were not found to be non-tuberculous mycobacteria bacteria on culture; QuantiF-ERON-TB Gold In-tube test conversion from negative to positive; or tuberculin skin test ≥15 mm and
 - * radiographic findings compatible with TB defined as ≥ 1 of the following factors identified independently by ≥ 2 of 3 paediatric radiologists serving on a masked review panel: calcified Ghon focus, pulmonary cavity, hilar or mediastinal adenopathy, pleural effusion, or airspace opacification and
 - * clinical manifestations compatible with TB defined as cough without improvement for > 2 weeks; weight loss > 10% of bodyweight for > 2 months; or failure to thrive, defined as crossing > 1 complete major centile band (< 97th-90th, < 90th-75th, <75th-50th, < 50th-25th, < 25th-10th, and < 10th-3rd weight-for-</p>

"Included all infants who met endpoint 1 criteria; had marginally less stringent criteria to define TB infection and household exposure."

Any of the following numerical categories.

- Isolation of *M tuberculosis* from any site.
- Identification *of M tuberculosis* by an approved molecular diagnostic technique from any site.
- Histopathology diagnostic for TB disease (such as caseating granulomas).
- Choroidal tubercle diagnosed by an ophthalmologist.
- Miliary pattern on chest x-ray in a HIV-negative infant.
- Clinical diagnosis of TB meningitis (CSF protein > 0.6 g/L and pleocytosis > 50/mm³ with mononuclear cell > 50%) or^a features of basal meningeal enhancement and hydrocephalus on head CT.
- Vertebral spondylosis
- A single smear/histology specimen positive for auramine-positive bacilli from a normally sterile body site.
- 1 of each of the following:
 - evidence of mycobacterial infection defined as:
 - 2 acid fast-positive smears each from a separate collection morphologically consistent with mycobacteria from either sputum or gastric aspirate that are not found to be non-tuberculous mycobacteria bacteria on culture, or
 - QFT conversion from negative to positive, or
 - \Box Tuberculin skin test \geq **10 mm**,^{*a*} or
 - ☐ household contact with AFB smear positive person^a and
 - * radiographic findings compatible with TB defined as ≥ 1 of the following identified independently by at least 2 out of 3 paediatric radiologists serving on a blinded review panel: calcified Ghon focus, pulmonary cavity, hilar/mediastinal adenopathy, pleural effusion, or airspace opacification and
 - clinical manifestations compatible with TB defined as either
 - cough without improvement for > 2 weeks, or
 - \Box weight loss \geq 10% of bodyweight for \geq 2 months, or
 - ☐ failure to thrive (crossing ≥ 1 entire major centile band downward) for ≥

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	age centiles) downward for > 2 months.	2 months, where the major centile bands are defined as < 97th–90th, < 90th–75th, < 75th–50th, < 50th– 25th, < 25th–10th, and < 10th–3rd weight-for-age centiles.		
Andrews 2017	Revised endpoint 1 from Tameris 2013 that removed QFT conversion from the diagnostic criteria to avoid bias to- wards association with QFT status.	Not used	Not used	
Bunyasi 2017	Not used	Not used	Same definition as for Tameris 2013.	
Ndiaye 2015	 Any of the following numerical categories. Isolation of <i>M tuberculosis</i> from any site. Identification of <i>M tuberculosis</i> by an approved molecular diagnostic technique from any site. Histopathology diagnostic for TB disease (such as caseating granulomas). Choroidal tubercle diagnosed by ophthalmologist. 	 Any of the following numerical categories: Isolation of <i>M tuberculosis</i> from any site. Identification of <i>M tuberculosis</i> by an approved molecular diagnostic technique from any site. Histopathology diagnostic for TB disease (such as caseating granulomas). Choroidal tubercle diagnosed by ophthalmologist. A single smear/histology specimen positive for AFB from a normally sterile body site. 2 acid-fast smears positive each from a separate collection morphologically consistent with mycobacteria from either pulmonary or gastric sampling that are not found to be non-tuberculous mycobacteria bacteria on culture, and ≥ 1 of the following: * a compatible radiographic feature: airspace opacification, cavity, hilar or mediastinal adenopathy, or pleural effusion; * a compatible clinical feature, i.e. > 2 weeks of fever, night sweats, anorexia, cough, or weight loss (≥ 5 kg by history or noticeable change in clothing fit); or ≥ 1 episodes of haemoptysis. 	Same definition as for Tameris 2013.	
Scriba 2011	Not applicable	Not applicable	Not applicable	

Nemes 2018	Outcomes not specified in the meth- ods section.	Not used	Not used
	In results, authors specified that 8 par- ticipants were diagnosed as TB:		
	"of whom one was M.tb [<i>Mycobacteri- um tuberculosis</i>] culture positive and 7 were diagnosed on clinical/ radi- ographic grounds and TB contact his- tory. Two of the TB cases were QFT positive."		

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

AFB: acid-fast bacilli; CSF: cerebrospinal fluid; CT: computerized tomography; QFT: quantiFERON; TB: tuberculosis. ^aIn Tameris 2013, endpoint 2: criteria in bold indicate where different from endpoint 1.

Study	Protocol		Published findings	Differences between — protocol and pub- lished findings	
	Stated out-Measurementcomes pub-of outcomelished pri-stated a priceor to com-mencementof trial thatdiffer to pub-lished out-comes		Measurement of out- come as stated in published findings		
Andrews 2017	No protocol pu	blished.			
Bunyasi 2017	No protocol pu	blished (extended p	ost-trial follow-up of Tame	eris 2013).	
Ndiaye 2015	Adverse events: blood tests ^a	"Percentage of participants with adverse events" AEs measured up to day 28 SAEs measured up to 6 months.	"Phlebotomy for rou- tine haematologi- cal and biochemical analysis was done at screening, before booster vaccination, and on days 7 and 28 after each vaccina- tion."	"Routine haematologi- cal and biochemical test results did not differ be- tween study groups (data not shown)."	Haematological and biochemical blood tests not outlined as a measure of safety in the study protocol. Blood test findings re- ported unclearly.
Nemes 2018	Safety	Clinicaltrial- s.gov – local, regional, and systemic AEs and SAEs which would be re- ported as cu- mulative 12- month inci- dences.	"Infants followed for safety end points at weeks 1, 4, 6, and 8 af- ter MVA85A/control vaccination and there- after, at weeks 9, 12, and 16 (corresponding to weeks 1, 4, and 8 following delayed BCG vaccination at 8 weeks of age), and at week 52."	Reported total events for AEs per group after MVA85A and before BCG and for whole follow-up period. Data including for laboratory AEs were not disaggregated as prespec- ified.	Data including for lab- oratory AEs were not disaggregated as pre- specified.
Scriba 2011	Safety ^a	Local and sys- temic AEs for the first week.	Diary cards	Local and systemic AEs re- ported on ≥ 1 day of the first 7 days after MVA85A vaccination.	None
		Blood tests (days 7, 28)	Biochemical and haematological tests (days 7, 28)	Reported number and per- centages of participants with abnormal results and reported that, "all except one patient that had el- evated liver enzymes re- mained unresolved by day 28."	-
		Immunology	ESAT-6/CFP-10	Infants converted – sug- gestive of TB infection but	-

Table 4. Differences between details of studies published prior to commencement and reported outcomes

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

Tameris 2013	Safety profile – AEs ^a	AEs measured up to day 28 SAEs measured throughout fol- low-up.	Collected data on so- licited and unsolicit- ed local and systemic AEs. Active surveillance for SAEs.	AEs broken down by type of event and reported in supplementary material. Only local events at the in- jection site were consid- ered to be related to the vaccine.	Causal relationship with AEs other than lo- cal injection site reac- tions was not report- ed.
	Safety profile – blood tests ^a	Testing up to 28 days postvacci- nation.	"Peripheral blood for routine haematolog- ical and biochemi- cal tests was taken at screening and on day 7 and day 28 after vaccination in an ini- tial safety cohort of at least 330 infants."	Not reported	Primary outcome not reported
	Efficacy of MVA85A ^b	Using an end- point derived from epidemi- ological cohort surveys in BCG vaccinated in- fants.	Not reported – sim- ply stated clinical end- points 'developed.'	Composite clinical end- points 1, 2, 3 (see Table 3) Microbiologically con- firmed cases reported in appendix.	The "primary efficacy endpoint" was mea- sured using an end- point not derived from cohort studies. The endpoint defi- nition differed from all other implied or reported ways of measuring efficacy in the other studies. The point estimate showed clinically sig- nificant benefit for endpoint 1 (no benefit seen at the 95% confi- dence level). This end- point was reported as the main efficacy find- ing. All other point es- timates show no clini- cally significant bene- fit or harm.

Table 4. Differences between details of studies published prior to commencement and reported outcomes (Continued)

seemed to be reported as safety data not efficacy.

AE: adverse events; BCG: bacillus Calmette-Guérin; ESAT-6/CFP-10: early secretory antigenic-6/culture filtrate protein-10; SAE: severe adverse events; TB: tuberculosis.

^{*a*}Primary outcomes as outlined in study protocols.

^bSecondary outcomes as outlined in study protocols.

Study	Participar	Participant reported adverse events			Outcome data reporting				Laboratory tests			
	Monitor- ing ac- tive or passive	Blinding of partic- ipants or outcome assessors	Times data collected	Times data re- ported	Com- plete/not com- plete	Percent- age of partic- ipants report- ed on	Analy- sis inde- pendent of study sponsor	Number of tests taken	Timing of tests and ade- quacy	Complete reporting of test re- sults	Inde- pen- dence of data analysis	
Scriba 2011	Active	Inadequate	60 min, D 2, 7, 28, 84, and 168	D 7, 28	Incom- plete	100%	Unclear	Biochemistry and haematol- ogy	Inade- quate	Inconse- quential	Unclear	
Ndiaye 2015	Active	Inadequate	D 7, 28, and 84 after boost 3 monthly until end of study	NR	Incom- plete	99.8%	No	Haematology, chemistry, viro- logical markers	Ade- quate	Incom- plete	No	
Tameris 2013	Active	Adequate	Baseline, D 7 and 28, throughout up to D 84	NR	Incom- plete	99.9%	No	Biochemistry and haematol- ogy	Inade- quate	Incom- plete	No	
Nemes 2018	Active	Adequate	Week 1, 4, 6, 8, 16, and 52	NR	Incom- plete	85.9%	Unclear	Not specified	Ade- quate	Incom- plete	Unclear	

D: day; min: minute; NR: not reported.

Table 6. Results of the different endpoints of active tuberculosis

Active TB	Tameris 2013		Andrews	Andrews 2017		Bunyasi 2017		Ndiaye 2015		Scriba 2011		Nemes 2018	
10	MVA85A	Placebo	MVA85A	Placebo	MVA85A	Placebo	MVA85A	Placebo	MVA85A	Placebo	MVA85A	Placebo	
End- point 1 ^a	32/1399 (2.3%)	39/1395 (2.8%)	58/2797 (2 NDD	.1%) with	N/A	N/A	6/320 (1.9%)	9/325 (2.8%)	N/A	N/A	5/123 (4.1%)	3/125 (2.4%)	
End- point 2 ^a	55/1399 (3.9%)	52/1395 (3.7%)	N/A	N/A	N/A	N/A	6/320 (1.9%)	9/325 (2.8%)	N/A	N/A	N/A	N/A	
End- point 3 ^a	196/1399 (14.0%)	177/1395 (12.6%)	N/A	N/A	3.3/100 pyo	3.0/100 pyo (95% Cl 2.6 to 3.5)	8/320 (2.5%)	9/325 (2.8%)	N/A	N/A	N/A	N/A	

Table 6. Results of the different endpoints of active tuberculosis (Continued)

(95% CI 2.9 to 3.9)

CI: confidence interval; N/A: not applicable; NDD: no disaggregated data; pyo: person-years of observation; TB: tuberculosis. ^aSee Table 3 for description of endpoints.

Table 7. Adverse effects of the MVA85A vaccine

Study	MVA85A		Placebo		Breakdown		Author conclusions	
	Number of partici- pants with ≥ 1 event caused by the inter- vention	Total par- ticipants	Number of partici- pants with ≥ 1 event caused by the control	Total par- ticipants	Detailed AEs	MVA85A	Placebo	_
Ndiaye 2015	318	324	307	325	Solicited AEs ^a	288	235	"Solicited adverse events were more common in MVA85A group and most were local injection site reactions."
Nemes 2018	105 ^b	123	30p	125	Not detailed	N/A	N/A	"Infants in MVA85A arm were more likely to experience an AE than in con- trol arm. Injection site reactions were more frequent in MVA85A recipients and mild."
Scriba 2011	106 ^c	108 ^c	6	36	Injection site ^d	106	6	"Desquamation significantly increased with greater vaccine dose "
					Malaise	6	1	with greater vacune dose.
					Lethargy	6	2	
					Tactile fever	18	0	
					Documented fever	13	2	_
					Vomiting	6	2	
					Elevated liver en- zyme levels	13	4	

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Table 7. Adverse effects of the MVA85A vaccine (Continued)

					Increased white cell count	0	1	
Tameris 2013	Local 1251 ^e	1399	Local 628 ^e	1396	Not detailed	1251	628	None

AE: adverse event; N/A: not applicable.

^{*a*}Included injection reactions, mild influenza-like symptoms, and regional lymphadenopathy.

^bAuthors of the study reported 105 participants with at least one adverse effect in the vaccine group and 30 participants in the placebo group, where causal relationship was defined as definite.

^cAggregated between three groups receiving different doses.

^dIncluded desquamation (scaling), pain, redness, and swelling.

eAuthors of the study reported local and systemic adverse events. Authors specified in their protocol that, "Solicited adverse events of local injection site reactions will be considered causally related to study vaccine (adverse reaction)." Therefore, we reported such adverse events as adverse effects. Causal relationship with other adverse events was not reported.

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Table 8. Adverse events summary table

Study	Adverse events of an	y severity		
	MVA85A			Placebo
Tameris 2013	1120/1399 (80.1%)			1059/1396 (75.9%)
Andrews 2017	NR			NR
Bunyasi 2017	NR			NR
Ndiaye 2015	321/324			312/325
	(99.1%)			(96%)
Scriba 2011	2.5×10^{7}	5 × 10 ⁷	1×10 ⁸	1/36
	pfu = 35 μL	pfu = 70 μL	pfu = 135 μL	
	1/36	3/36	6/36	-
Nemes 2018	Mild 122/123			121/125
	(99.2%)			(96.8)
	Moderate 62/123			54/125
	(50.4%)		(3.6%)	
	Severe 11/123			14/125
	(8.9%)			(11.2%)

NR: not reported; pfu: plaque-forming unit.

Table 9. Abnormal haematological and biochemical tests

Study	Haematolog tests	gical blood				
	MVA85A	Placebo	MVA85A			Placebo
Tameris 2013	NR	NR	NR			NR
Andrews 2017	NR	NR	NR			NR
Bunyasi 2017	NR	NR	NR	NR		
Ndiaye 2015	NR ^a	NR ^a	NR ^a			NR ^a
Scriba 2011	0/108 ^b	1/36 ^b	2.5 × 10 ⁷ pfu = 35 μL	5 × 10 ⁷ pfu = 70 μL	1 × 10 ⁸ pfu = 135 μL	4/36 (11%)
			1/36	3/36	9/36	-
			(2.8%)	(8.3%)	(25%)	

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Table 9. Abnormal haematological and biochemical tests (Continued)

Nemes 2018	NR	NR	14/123 (11.4%)	13/125
				(10.4%)

NR: not reported; pfu: plaque-forming unit.

^{*a*}Authors stated that routine haematological and biochemical test results did not differ between study groups but did not present data. ^bOne participant had increased white cell count concurrently with an increase in alanine aminotransferase during an episode of gastroenteritis. Authors did not describe any other case of abnormal haematological test in the rest of the participant, although it was not stated explicitly.

APPENDICES

Appendix 1. Search strategies

Cochrane Central Register of Controlled Trials

#1 tuberculosis or TB:ti,ab,kw (Word variations have been searched)

#2 MeSH descriptor: [Tuberculosis] explode all trees

#3 MeSH descriptor: [BCG Vaccine] explode all trees

#4 "BCG vaccin*":ti,ab,kw (Word variations have been searched)

#5 bacill* Calmette-Guerin

#6 #1 or #2 or #3 or #4 or #5

#7 "antigen 85A" or Ag85A or "modified vaccinia ankara" or MVA85A

#8 MVA85*

#9 #7 or #8

#10 #9 and #6

MEDLINE (PubMed)

#12	Search #7 and #11
#11	Search ((#8) OR #9) OR #10
#10	Search "drug therapy" [Subheading]
#9	Search randomized or placebo or randomly or trial or groups Field: Title/Abstract
#8	Search "Randomized Controlled Trial" [Publication Type] OR "Controlled Clinical Trial" [Publica- tion Type]
#7	Search #3 and #6
#6	Search 4 or 5
#5	"antigen 85A" OR Ag85A OR "modified vaccinia ankara" OR MVA85A Field: Title/Abstract
#4	"antigen 85A, Mycobacterium tuberculosis" [Supplementary Concept] or "MVA 85A" [Supplemen- tary Concept])

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(Continued)	
#3	Search 1 or 2
#2	(("BCG Vaccine"[Mesh]) OR ("bcg vaccin*" or "bacille Calmette-Guérin")Field: Title/Abstract
#1	"Tuberculosis"[Mesh] or (tuberculosis or TB) Field: Title/Abstract

Embase

1 (tuberculosis or tuberculous or TB).mp.

2 tuberculosis/

31 or 2

4 BCG vaccine/ or BCG vaccin*.mp. or BCG vaccination/

53 or 4

6 MVA85A.mp.

7 antigen 85A.mp.

8 Ag85A.mp.

9 modified vaccinia virus ankara.mp.

10 modified vaccine ankara.mp.

11 6 or 7 or 8 or 9 or 10

12 5 and 11

13 (randomized or randomised or placebo or double-blind* or single-blind*).mp.

14 randomized controlled trial/ or controlled clinical trial/

15 crossover procedure/

16 13 or 14 or 15

17 12 and 16

CINAHL (EBSCOHost)

#	Search terms
S1	TX (tuberculosis or TB or BCG)
S2	TX ((MVA85A or "antigen 85A" or "modified vaccinia ankara")
S3	TX ((randomized trial or controlled trial or placebo or double-blind * or single-blind *)
S4	S1 AND S2 AND S3

Web of Science

MVA85A vaccine to enhance BCG for preventing tuberculosis (Review)

# 2	TOPIC: (tuberculosis or TB or BCG) <i>AND</i> TOPIC: (MVA85A or "antigen 85A" or "modified vaccinia ankara") <i>AND</i> TOPIC: (randomized trial or controlled trial or placebo or double-blind* or single-blind*)
	Timespan=All years
	Search language=Auto
#1	TOPIC: (tuberculosis or TB or BCG) <i>AND</i> TOPIC: (MVA85A or "antigen 85A" or "modified vaccinia ankara")
	Timespan=All years

CONTRIBUTIONS OF AUTHORS

RK drafted the review, screened abstracts, extracted data, analysed results, and wrote the final review.

SoJ (Sophie Jullien) drafted the review, screened abstracts, extracted data, analysed results, and wrote the final review.

PG contributed to the methods, coherence, and writing of the final review.

SaJ (Samuel Johnson) co-ordinated the review, screened abstracts, extracted data, performed analysis of data, and helped draft the final review.

DECLARATIONS OF INTEREST

RK has no known conflicts of interest.

SoJ worked for the CIDG from September 2015 to April 2016.

PG is the Director of the Research, Evidence and Development Initiative (READ-It) project (project number 300342-104) and CIDG Coordinating Editor.

SaJ worked for the CIDG from January 2017 to July 2018.

None of the review authors receive salary, payment, academic fees, or academic status related to vaccine development.

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Internal sources

• Liverpool School of Tropical Medicine, UK.

External sources

• Department for International Development, UK.

Project number 300342-104

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

Changes to the author team: Taryn Young stepped down from the review author team.

We intended to pilot data extraction forms; however, given the small number of included studies we assessed the appropriateness of the form during the actual data extraction.

In our protocol, we mentioned that the control for the type of intervention would be "BCG alone." However, we did include in our review studies that they used Candin[®] as control intervention, as this is currently used in control groups for randomized controlled trials assessing MVA85A.

We encountered multiple different definitions of active tuberculosis in different trials. We took the approach of defining active tuberculosis as confirmed by culture and participants starting on tuberculosis treatment to allow a consistent approach across the included studies.

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We reported adverse effects of any severity disaggregated by local reactions of the skin and systemic symptoms and we gave justification for this decision in the result section.

The initial risk of bias for adverse event assessment tool had three options to assess completeness of reporting of participant-reported outcomes. The options complete/incomplete/unclear were reduced to complete/incomplete as there was no difference between the options incomplete and unclear reporting.

The detailed subgroup analysis prespecified in the protocol was not done due to too few studies.

INDEX TERMS

Medical Subject Headings (MeSH)

*BCG Vaccine; *Tuberculosis Vaccines; HIV Seropositivity [complications] [immunology]; Primary Prevention; Randomized Controlled Trials as Topic; Tuberculosis [*prevention & control]

MeSH check words

Humans