Is Bacille Calmette-Guérin (BCG) Vaccine a Known Risk Factor for Latent Tuberculosis Infection?: A Cross-sectional Study on 180 New Immigrants from BCG-vaccinated Countries to Kuwait

Adel AL-Harbie1,*, Kim Picozzi1, Michael Thrusfield2, Ali Sadek3 and Susan C Welburn1

1Division of Pathway Medicine & Centre for Infectious Diseases, School of Biomedical Sciences, College of Medicine & Veterinary Medicine, The University of Edinburgh, Chancellor’s Building, 49 Little France Crescent, Edinburgh, EH16 4SB, UK
2Veterinary Clinical Sciences, Royal (Dick) School of Veterinary Studies. The University of Edinburgh, Easter Bush, EH25 9RG, UK
3Department of Public Health and Community Medicine, Faculty of Medicine, University of Alexandria, Egypt

Abstract

The Bacille Calmette-Guérin (BCG) vaccine has existed for 90 years and is the most widely used of all current national childhood immunization programmes. The impact of BCG vaccination on transmission of Mycobacterium tuberculosis is not clear or whether BCG confers lifelong immunity through sufficient protection against infection and diagnosis of latent tuberculosis infection (LTBI) is recently questionable especially in tuberculosis endemic regions and vaccinated high risk individuals, which was discussed along the paper work following a new evidence-based criteria for LTBI diagnosis.

Keywords: Bacille Calmette Guerin (BCG); Tuberculosis (TB); Latent tuberculosis infection (LTBI); Tuberculin skin test (TST)

Introduction

Since the 20th century the Bacille Calmette-Guérin (BCG) vaccine there is only one defensive tool against Mycobacterium tuberculosis (MTB) introduced as a shot of a 90-year-old vaccine (attenuated virulence of Mycobacterium bovis). BCG limits the growth and protect against MTB and aids in clearance of bacillary loads during chemotherapeutic treatment in the active post-infective stage [1]. BCG vaccination can have obvious public health advantages. It can lessen the burden of TB-related disorders such as LTBI prevalence and active TB incidence in children [2]. TB incidence rose again after abandoning routine BCG in the mid-1980s even with the introduction of other preventive measures [3]. But the efficacy impacts of BCG vaccination whether confers lifelong immunity or adversely causing tuberculosis (TB) through indirectly forming latent tuberculosis infection (LTBI) is still unclear. Worldwide the tuberculin skin test (TST) has been the standard diagnostic test for detection of LTBI for more than 100 years. It was first described by Robert Koch in 1890, and involves injecting intradermally a purified protein derivative (PPD) of 0.1 ml tuberculin (which is a glycerol extract of the bacillus) followed by measurement of the localized delayed-type hypersensitivity (DTH-IV) reaction [4]. The TST has several limitations and cannot distinguish between previous BCG vaccination and current TB infection due to poor specificity in BCG negative people and poor sensitivity in BCG-vaccinated individuals. Meta-analysis has shown that previous BCG administration increases the likelihood of TST false-positive results up to 15 years after-vaccination [5].

BCG vaccination might alter the interpretation of a positive PPD, and therefore TST interpretation is extremely limited as a diagnostic tool to prove whether a positive result is caused by MTB infection or cross-reactivity resulting from BCG vaccination (containing live attenuated M. bovis antigens), and has to be interpreted taking into consideration both the pre-test risk of TB infection and BCG vaccination status.

Objective

To assess the interference effect of BCG vaccination on the results of the tuberculin skin test.
Table 1 presents a new laboratory classification of diagnostic testing, which can be considered as new criteria for LTBI case definition.

**Statistical analysis**

Lowest accepted sample size calculated by the Confidence Interval Analysis Software was 167 out of average annual 80,000 expatriates (theoretically are healthy or free-of-disease) entering to Kuwait [7]. Data were analyzed using the Statistical Package for Social Sciences software, version 17.0 [8]. Categorical variables were analyzed using a Likelihood ratio (LLR χ²(d.f.)) test when Pearson's chi square was inappropriate for small sample (measured by SPSS as expected values were more than 20% or more 5 categories of total table categories). Similar to LLR χ²use, non-normally distributed continuous variables (age of BCG vaccination in years) were analyzed using a Kruskal-Wallis (KW χ²(d.f.)) test and a p-value of less than 0.05 was considered as the level of statistical significance. Data limits using the mid-spread (middle 50%) represented by the interquartile range (IQR) and semi-interquartile range (SIQR) as half the interquartile range [9]. Questionnaire quantitative variables were drawn as box-and-whiskers plots diagrams using SPSS version 17.0 [8].

**Results**

**Socio-demographic characteristics**

The overall median age of the immigrants was 31.5 years, with an interquartile range (IQR) of 11 years and the semi-interquartile range (SIQR) of 5.5 year (KW χ²(4)=25.741, p<0.001). Statistically significant differences were detected between nationality and LTBI case categories. Most of the immigrants in this study 77.22% (139/180) came from countries in which TB was endemic: Indians 35%, Filipinos 19.44%, Nepalis 15%, Ethiopians 7.78% and Sri Lankans 3.3%. Egyptians represented the majority of immigrants coming from non-endemic countries: 14.44% (26/180) (LLR χ²(52)=72.522, p=0.032).

**Tuberculin skin test**

Mantoux (induration) reaction was measured in 98.33% (177/180) who fulfilled the inclusion criteria for TST results. The TST results of three excluded immigrants were as follows: two participants came on day 7 (a male Indian electrician and a female Nepali housemaid) and one participant did not return for a TST reading (a female Ethiopian housemaid). The frequency of LTBI reactions in the new immigrants, including the three missed follow-up participants, is shown in Table 2.

According to the induration reaction, the prevalence of LTBI in the sample of new immigrants was zero (0/177) using the TST method. Applying the new classification criteria of LTBI, all participants belonged to the normal results within score I (zero=negative reaction) entering to Kuwait [7].

**BCG vaccination status**

Analysis of the influence of BCG vaccination status on test results
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Table 3: Distribution of tuberculin skin test results according to latent tuberculosis infection categories of 180 new immigrants to Kuwait, February-May, 2010.

| Tuberculin Skin Test | Negligible | Low | Average | High | Extremely High | Can not be judged | Total | Likelihood | p-value |
|----------------------|------------|-----|---------|------|----------------|------------------|-------|------------|---------|
|                      | (n=90)     | (n=36) | (n=0)   | (n=4) | (n=46)         | (n=180)         |
| Score I (<5)         | n %        | n %   | n %     | n %   | n %            | n %             |
| TST                   | 90 100.00  | 36 100.00 | 0 0.00 | 3 75.00 | 46 100.00 | 1 25.00 | 176 | 97.78 |
| Score II (5-9)       | 0 0.00     | 0 0.00 | 0 0.00 | 1 25.00 | 0 0.00 | 0 0.00 | 1 0.56 |<0.001 |
| Missed               | 0 0.00     | 0 0.00 | 0 0.00 | 0 0.00 | 0 0.00 | 0 0.00 | 3 75.00 | 3.667 |
| No reaction          | 85 94.44   | 36 100.00 | 0 0.00 | 3 75.00 | 45 97.83 | 1 25.00 | 194.44 |
| TST <5               | 5 5.56     | 0 0.00 | 0 0.00 | 0 0.00 | 1 21.70 | 0 0.00 | 6 3.33 |
| Presence of Positive cutaneous in Normal | 37.947 |<0.001 |
| duration size (mm)   | 5-9        | Reactive- | Borderline | 0 0.00 | 0 0.00 | 0 0.00 | 0 0.00 | 3 75.00 | 3.667 |
| TST adverse reaction | 90 100.00 | 36 100.00 | 0 0.00 | 4 100.00 | 46 100.00 | 1 25.00 | 177 98.33 |<0.001 |

Average LTBI category is zero (means no detected cases). Missed=excluded case due to missed follow-up or came after 5 days for reading of TST reaction, cannot be judged means excluded cases (3 missed follow-up for TST results and one pregnant female not preformed chest X-ray due to contraindication).

Table 4: Distribution of Bacillus Calmette-Guérin (BCG) vaccination status against tuberculosis according to latent tuberculosis infection categories of 180 new immigrants to Kuwait, February-May, 2010.

| BCG vaccination against | Negligible | Low | Average | High | Extremely High | Can not be judged | Total | Likelihood | p-value |
|-------------------------|------------|-----|---------|------|----------------|------------------|-------|------------|---------|
| History of vaccination  | TB (n=90)  | (n=36) | (n=0)   | (n=4) | (n=46)         | (n=180)         |
| Yes                     | 78 87      | 28 87 | 77.80   | 0 0.00 | 100.00 | 41 89 | 3 75.00 | 154 85.56 |
| No                      | 10 11      | 7 19.44 | 0 0.00 | 0.00 | 5 11 | 1 25.00 | 23 | 12.78 |
| Unknown                 | 2 2        | 1 2.78 | 0 0.00 | 0.00 | 0 0 | 0 0.00 | 3 | 1.67 |
| Age of BCG vaccination  | Median (IQR) | 5.00 | (2) | 6.00 | (3) | - | 7.50 | (6) | 5.00 | (2) | 4.00 | (-) | 5 | (2) | 3.330 |
| Presence of BCG scar   | Positive  | 79 87.78 | 27 75.00 | 0 0.00 | 4 100.00 | 41 89.13 | 4 | 100.00 | 155 | 86.11 |
| Negative               | 11 12.22 | 9 25.00 | 0 0.00 | 0.00 | 5 10.87 | 0 | 0.00 | 25 | 13.89 |
| Animal exposure in county | Yes  | 51 56.67 | 16 50 | 0 | 0 | 1 | 25.00 | 24 | 52.17 | 0 | 0.00 | 94 | 52.22 |<0.001 |
| Having cow?            | No         | 39 43.33 | 18 50 | 0 | 0 | 0 | 75.00 | 22 | 47.83 | 4 | 100.00 | 86 | 47.78 |

is shown in Table 3. BCG vaccination status was ascertained from the participants through availability of reliable past history of vaccination and/or presence of a characteristic BCG scar. Analysis of the effects of BCG vaccination status (either vaccinated or non-vaccinated) represented in relation to the defined cases of latent tuberculosis infection categories is shown in Table 4.

The overall median age of BCG vaccination for the 180 participants was 5 years (IQR=2 years). The 'high' LTBI group had been vaccinated at older ages (median=7.5 years, IQR=6 years). The minimum (youngest) age of vaccination was one year versus the oldest maximum age of 13 years was answered (Table 4). However these findings did not show a statistically significant difference between normal vaccination ages for LTBI development (KW χ²(4)=3.330, p=0.504) (Figure 1).

Presence of BCG vaccination scar was not a statistically significant risk factor for LTBI development or to cause positive a TST reaction, even in those expatriates classified as 'high' and 'extremely high' LTBI cases who were having forearm scar.

A positive scar was detected in 86.11% (155/180) of participants (LLR χ²(4)=6.104, p=0.192). The overall median age of BCG vaccination for the participants was 5 years (IQR=2 years) (Table 4).

Immigrants having a history of exposure to poultry animals e.g.
negative results of all immigrants were not significantly associated with the risk of being diagnosed as active or latent TB was also revealed by Li et al. (2010) [13] detected LTBI prevalence of positive TST in 24.4% populations can be effective and might add suspicion for LTBI carriers. at immigrant entry centres in Kuwait that serve large foreign-born to predict LTBI suspicious carriers [12]. Therefore TB skins testing finding was emphasized on skin testing in BCG-vaccinated populations can be assessed by the presence or absence of a BCG scar [11]. Similar vaccine administration and age at vaccination. Successful vaccination confounder, which affects TST reactivity, is prior vaccination with BCG such as interferon gamma release assays (IGRAs) [10]. A common (reducing TB reservoirs) by using other higher specific diagnostics in infection and highlights the need for LTBI early and accurate detection from TB high-incidence regions represents a potential pool for new TB

Discussion

The influx of BCG-vaccinated immigrants in non-endemic countries from TB high-incidence regions represents a potential pool for new TB infection and highlights the need for LTBI early and accurate detection
(reducing TB reservoirs) by using other higher specific diagnostics such as interferon gamma release assays (IGRAs) [10]. A common confounder, which affects TST reactivity, is prior vaccination with BCG vaccine, and/or the strain and dose of BCG used, and/or method of vaccine administration and age at vaccination. Successful vaccination can be assessed by the presence or absence of a BCG scar [11]. Similar finding was emphasized on skin testing in BCG-vaccinated populations to predict LTBI suspicious carriers [12]. Therefore TB skins testing at immigrant entry centres in Kuwait that serve large foreign-born populations can be effective and might add suspicion for LTBI carriers. Li et al. (2010) [13] detected LTBI prevalence of positive TST in 24.4% (higher among foreigners) in BCG-vaccinated which help to target TST testing before starting immigrant’s chemoprophylaxis.

Exposure to previous BCG vaccination did not significantly reduce the risk of being diagnosed as active or latent TB was also revealed by Caley et al. (2010) [14], in common with our findings.

This study proved that the presence of a BCG scar and related TST negative results of all immigrants were not significantly associated with a higher prevalence of LTBI as also noted by Demkow et al. (2008) [15], Other similar significant findings was detected the absence of risk difference in TB patient contacts having positive BCG scar compared to other normal control contacts without scars [16]. Also Kik et al. (2009) [17] showed no association between the presence of BCG scar with recent exposure to TB in immigrants having positive IGRAs and TST results.

Since BCG-TST positive reactivity wanes with time (if more than five years have elapsed since administration of BCG vaccine) - a positive TST reaction is most likely a result of exposure to MTB infection [18]. Gomes and colleagues (2011) [19] have recently concluded that waning of the BCG-induced protection was associated with raised risks of TB morbidity rates in children aged between three and five years. Soysal et al. concluded that absence of an immunization scar can determine the likelihood for TB infection and is significantly associated with TB disease severity, which can be related to the level of exposure to MTB [2].

TST reactions can be interpreted regardless of BCG vaccination history [6]. Absence of positive TST reactions in our study strengthens the inference that BCG is not interfering with TST interpretation even though 86.11% (155/180) of participants had already been vaccinated against TB with BCG around pre-school ages. Similar related finding we recompiled by Minodier et al. (2010) [20], namely that a positive TST is more likely to be related to an increased duration of TB exposure in the TB-endemic country of birth rather than to previous BCG vaccination. TST cannot be positive in certain biological factors due to suppression of DTH-IV reaction and T-lymphocytes such as malignancies and viral infections (e.g. HIV) [21].

Bradshaw et al. (2011) [22] suggested that advancing age increases the likelihood of exposure to unpasteurized dairy products and cross-reactions of environmental mycobacterial antigens using IGRA tests. A similar significance in our research results of those participants having past history of exposures to various animals and positive BCG scar without LTBI diagnosis. On the contrary Grafein et al. (2011) [23] concluded that there was an absence of a significant association between the presence of LTBI and a BCG scar, but presence of a significant association with consumption of unpasteurized (cows) milk in the past 6 months, which can be related to Mycobacterium bovis.

Similar to the majority of BCG-related publications, the size of BCG scar was not measured in our research because did not correlate with protection against TB, and also is not an indication for diagnosis (or presence) of LTBI. A similar result was also achieved by Crampin et al. (2009) [24].

TST reactivity caused by BCG vaccine generally wanes with the passage of time, but periodic skin testing may prolong reactivity in vaccinated persons, the phenomenon of ‘boosting reactivity’ [6] (CDC, 2010). On the contrary, TST reaction size and TST results were not affected by the time since the last dose of BCG vaccination, the number of BCG scars or BCG vaccination schedule in children, and significantly affected the decision for LTBI prophylaxis [25].

Study Limitation and Strength

The absence of a gold standard test for latent TB infection and TB can be considered as a limiting factor. Another limitation is the small sample size which might constrain generalization of the results to a larger population and wider community. On the other hand presenting and comparatively testing the new evidence-based diagnostic criteria for LTBI in addition to absence of any previous data on LTBI prevalence in immigrants or Kuwait residents add strength for future comparison.
Conclusions

Tuberculin skin test is extremely limited as a diagnostic tool for latent tuberculosis infection to prove whether a positive result is caused by cross-reactivity resulting from BCG vaccination. Still a history of BCG and/or presence of vaccination scar (those answered ‘No’ or ‘Unknown’ vaccination status) is not a contraindication for tuberculin skin testing or LTBI chemoprophylaxis in suspected individuals.

Recommendations

The relationship between immunogenicity against tuberculin skin test in the response to BCG vaccination needs to be elucidated. Considering the confounding effect of Mycobacterium tuberculosis risk factors, BCG efficacy and protection need further evaluation using interferon gamma release assays to compare between LTBI detected in both vaccinated and non-vaccinated populations. Emphasis on skin testing in BCG-vaccinated populations should be considered in the appropriate clinical setting to predict LTBI suspicious carriers.

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