Tuberculosis infection in foreign-born children: a screening survey based on skin and blood testing

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This study, carried out in a low tuberculosis (TB) prevalence country with high immigration rates from high TB prevalence countries, deals with the interferon-gamma release assay, QuantiFERON®-TB Gold In-Tube, for the diagnosis of latent TB infection (LTBI) in foreign-born children. The results of our study highlight the potential advantages and concerns of using a blood test for diagnosing LTBI in a ‘two-step’ strategy in foreign-born children.

KEY WORDS: latent tuberculosis infection; foreign-born children; active tuberculosis; interferon-gamma assays

OVER THE LAST 30 YEARS, industrialised countries have witnessed a progressive increase in tuberculosis (TB) incidence, partly related to the growing number of immigrants from high TB incidence areas. Accurate diagnosis of latent TB infection (LTBI) cases is a desirable goal in achieving effective TB control in low-incidence countries, particularly in young children, who are more prone to severe disease as a consequence of recent infection and in whom preventive chemotherapy is effective and relatively safe.1,2

Until recently, LTBI detection relied only on the tuberculin skin test (TST), known to be associated with false-positive results in children infected with nontuberculous mycobacteria (NTM) or bacille Calmette-Guerin (BCG) vaccinated, as is the case with the majority of the children born in high TB incidence countries.3,4 Furthermore, TST sensitivity may be impaired in young children, and in several other conditions such as immunosuppression.3,5 The positive cut-off of the TST induration therefore depends on the likelihood of infection and on the individual risk of developing active TB.4,6 Interferon-gamma release assays (IGRAs) represent an attractive tool for TB screening due to their high diagnostic accuracy, although their performance in paediatric subjects needs corroborating evidence.

Policies for TB screening in immigrant children vary widely across Europe.8 For immigrants from TB-endemic countries, the current Italian national guidelines set the cut-off for a positive TST at 10 mm, regardless of age and BCG vaccination status.

RESULTS

We performed 621 TSTs in children and adolescent immigrants, mostly from Africa (40.3%) and Eastern Europe (26.9%): 104 (16.7%) were TST-positive, 128 (20.6%) were TST-borderline and 389 (62.6%) TST-negative. Children in the first two groups were tested with QFT at 31 ± 29 days (mean ± standard deviation) after TST. Three children had an invalid QFT result and were excluded from further analyses. QFT was positive in 80 (34.5%) children: 30 had borderline TST (30/125, 24%) and 50 were TST-positive (50/104, 48.1%; Figure 1). By logistic regression analysis, a positive correlation was found between a positive QFT and the size of TST induration (odds ratio [OR] 1.23, 95% confidence interval [CI] 1.12–1.36, P < 0.001).
The distribution of TST and QFT results according to age is shown in Figure 2. TST-positive, QFT-negative results diminish with increasing age (OR 0.88, 95%CI 0.82–0.95, \( P = 0.001 \)), while TST-borderline, QFT-positive results are not related to age (OR 1.04, 95%CI 0.94–1.15, \( P = 0.458 \)). All QFT-negative, TST-positive tests had interferon-gamma (IFN-\( \gamma \)) levels <0.20 international units (IU)/ml, while the majority \( (n = 26) \) of the 30 QFT-positive children with borderline TST had IFN-\( \gamma \) levels > 0.50 IU/ml.

All children were healthy on clinical examination and none complained of respiratory symptoms. Nevertheless, four TST-positive, QFT-positive children had radiological findings consistent with active TB, confirmed by microbiological culture (acid-fast bacilli smear-negative). These TB cases were aged >10 years and BCG-vaccinated: TST size and IFN-\( \gamma \) levels were respectively 12 mm/17.0 IU, 13 mm/15.9 IU, 11 mm/16.2 IU and 10 mm/9.4 IU. Isoniazid (INH) was offered to all QFT-positive children \( (n = 76) \), with the exception of two children with coexisting chronic liver disease. Based on clinical judgment, children aged <8 years with TST > 15 mm \( (n = 3) \) were also treated with INH, as were those subjects for whom reliable follow-up could not be guaranteed \( (n = 4) \). For TST-positive, QFT-negative children, paediatricians were advised to provide strict follow-up; at the time of writing, no additional cases of active TB have been notified in this group.

**DISCUSSION**

Using blood tests to diagnose LTBI in immigrant children permits the diagnosis of infection in a sizeable fraction of children with TST results between 5 and 9 mm: we report here that about one third of these children were QFT-positive, usually with high IFN-\( \gamma \) levels. Although longitudinal data in children are not yet available, it is currently assumed that individuals with TST induration between 5 and 9 mm and a positive QFT are likely to be infected with *Mycobacterium tuberculosis*.\(^{10}\) Based on this assumption, we offered treatment to all of these children. Assuming that children with a positive QFT are truly infected and that those with a negative TST are non-infected, LTBI...
prevalence in this group could be estimated to be around 13%. A limitation of this study is that TST-negative children did not undergo QFT testing, and we cannot exclude that some of them might also be infected.

As previously reported, only half of the children with a positive TST were QFT-positive. Both BCG vaccination and exposure to NTM might help in interpreting some of these TST results as falsely positive; recent longitudinal data in household contacts would support this concept.11 Active pulmonary TB was diagnosed in four BCG-vaccinated, asymptomatic adolescents: all were positive on both QFT (at high IFN-γ levels) and TST (>10 mm). These findings confirm recent data obtained in the context of contact screening showing that quantitative results of both TST and IGRA may assist in the identification of subjects with active disease.12,13 A combination of a TST > 10 mm and high QFT levels might also help to identify children with active disease in the context of screening programmes.

BCG vaccination is usually given in the first days of life, and its effect on TST is presumed to be strongest in the first 10–15 years.4 On the other hand, younger children are more prone to developing severe disease. As the sensitivity of IGRA in young children has not yet been proven, the decision to disregard a positive TST because of a negative QFT is debatable, and caution should be used in this regard. It is therefore crucial to emphasise the importance of parental education regarding symptoms of TB. Consistent clinical follow-up to monitor for disease progression in young children should be in place, particularly considering the poor (TST) or still unproven (IGRA) sensitivity of diagnostic tests in this population group. Last, our data indicate that clinicians must always consider age and risk of disease progression when deciding whether to disregard a positive TST in young immigrant children.

References


Notre étude, menée dans un pays à faible prévalence de
la tuberculose (TB) avec des taux d’immigration élevés
provenant de pays à haute prévalence de la TB, traite du
test de libération de l’interféron-gamma, QuantiFERON®-
TB Gold In-Tube pour le diagnostic de l’infection tuber-
culeuse latente (LTBI) chez les enfants nés à l’étranger.

Les résultats de notre étude illustrent les avantages po-
tentiels et les préoccupations concernant l’utilisation
d’un test sanguin pour le diagnostic d’une LTBI dans
une stratégie en deux étapes concernant les enfants nés à
l’étranger.

RÉSUMÉ

El presente estudio se llevó a cabo en un país de baja
prevalencia de tuberculosis con un alto nivel de inmi-
gración procedente de países con tasas altas de preva-
lencia de esta enfermedad; se analizó el rendimiento de
la prueba de liberación de interferón gama, Quanti-
FERON®-TB Gold En Tubo, en el diagnóstico de la in-
fección tuberculosa latente (LTBI) en los niños nacidos
en el extranjero. Los resultados destacan las posibles
ventajas y las dificultades del uso de una prueba san-
guínea para establecer el diagnóstico de LTBI en los
niños nacidos en el extranjero, mediante una estrategia
en dos etapas.

RESUMEN