EDITORIAL

Interferon-gamma release assays for screening of health care workers in low tuberculosis incidence settings: Dynamic patterns and interpretational challenges

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There is considerable evidence that both tuberculin skin tests (TSTs) and interferon-gamma release assays (IGRAs) are valid but imperfect for latent tuberculosis (TB) infection (LTBI). Neither test can distinguish LTBI from TB disease and, therefore, have no value for active TB detection (1). Both tests have suboptimal sensitivity in active TB, especially in HIV-infected persons and children (2,3). Both tests appear to correlate well with gradient of exposure (3). While neither IGRAs nor the TST have high accuracy for predicting active TB, the use of IGRAs in some populations might reduce the number of people considered for preventive treatment (4).

However, there are key differences between the two tests (5). While the TST has high specificity in Bacille-Calmette-Guérin (BCG) unvaccinated persons, its specificity is lower and variable in those who have received BCG vaccination after infancy or have received multiple BCG vaccinations (6). In contrast, IGRA specificity remains high in BCG-vaccinated and unvaccinated populations (7). From a logistical perspective, IGRAs are more convenient for patients who do not have to return for the reading, and the laboratory readout is more objective than the subjective reading of TST induration. Finally, there is an important difference in terms of cost for the health care system – in general, IGRAs are more expensive to implement than the TST.

While many countries have published guidelines on IGRAs (8), the use of IGRAs for routine screening of health care workers (HCWs) remains an area of controversy. The 2005 United States Centers for Disease Control (CDC) guidelines on the QuantiFERON-TB Gold (QFT) assay (Cellestis, Australia) allowed for replacement of the TST with QFT for annual testing of HCWs in the United States (US) (9), and this was expanded to cover both commercial IGRAs in the 2010 update (10). In contrast, the Canadian guidelines on IGRAs have not recommended the use of IGRAs for serial testing of HCWs (11,12).

While the performance of IGRAs in serial testing of HCWs was first reported six years ago (13), this topic has received significantly more attention in the past few years, culminating in a recent systematic review (14). In the current issue of the Canadian Respiratory Journal, Joshi et al (15) (pages 84-88) provide useful data on routine implementation of QFT for HCW screening in Arkansas (USA), where QFT replaced the TST in 2008. Joshi et al describe the challenges they faced in implementing the test and raise concerns about high rates of QFT positivity in a setting with very low historical TST conversion rates. They observed high reversion rates on repeat testing of positives and poor short-term reproducibility of positive QFT results. Based on their results, Joshi et al argue for the need for cautious interpretation of QFT results, especially those in the borderline zone around the cut-off.

The report by Joshi et al (15) is highly consistent with data that have emerged from many countries (16-25). Table 1 summarizes the major serial testing studies of IGRAs in HCWs in low and intermediate TB incidence settings (16-25), including the data from Arkansas (15), building on the systematic review by Zwerling et al (14). Overall, these data suggest that IGRA results vary greatly during serial testing. When simple negative/positive changes are used as cut-offs, IGRAs had high rates of conversions (2% to 15%), which were frequently higher than the rates of TST conversions, and higher than the annual risk of TB infection expected in these low incidence settings. What is striking are the high rates of IGRA reversions – which range from approximately 20% to 40% in most studies.

Thus, while both TST and IGRAs are impacted by nonspecific variations, conversions and reversions during serial testing, IGRAs appear to have a very high frequency of conversions and reversions, and the consequences of such changes are unclear, posing challenges for implementation in routine occupational health programs. Indeed, a series of recent studies have described the challenges faced by US hospitals that switched to IGRA for HCW screening after the 2005 CDC guideline (15,17,25-28).

Despite the growing evidence base on IGRAs, there are several unresolved questions, especially in the context of serial testing (29). What cut-off should be used for IGRA conversion? We now know that a simple negative to positive definition is probably inappropriate and will result in a much higher conversion rate than what epidemiologically might be expected. This would then expose more HCWs to potentially unnecessary preventive therapy. Several studies have examined the within-person reproducibility of IGRA results over time, and many have proposed more stringent cut-offs based on reproducibility data, and proposals for a borderline zone of uncertainty have also been made (13,30-35). However, there is no consensus on the optimal cut-offs for conversions and reversions, and the need for a borderline zone for both IGRAs.

Should all those who are IGRA positive at baseline get retested routinely? Should IGRA positive results be repeated before recommending treatment? If yes, then what is the best approach for managing the anticipated reversions? Should those with IGRA reversions be carefully monitored but not given preventive therapy? What is the prognostic significance of IGRA reversions? Studies that have performed multiple time-point testing have shown diverse patterns of changes over time, and the prognosis of these phenotypes remains unclear (36), although there is evidence that both TST and IGRAs have suboptimal predictive value (4).

Regardless of the test used for serial testing, there is a need to reconsider the general strategy of annual testing of HCWs in North America, where TB incidence has declined to a very low level. Millions of HCWs undergo screening for LTBI every year, and a substantial proportion are low-risk employees who probably should not be screened at all. Arkansas is an excellent example, where the TB case rate was only 2.7 per 100,000 population in 2010 (37), which is reflected in the historical annual TST conversion rate of less than...
TABLE 1
Serial testing studies of interferon-gamma release assays in health care workers (HCWs) in low and intermediate incidence countries

| Author (reference), year, country | Duration between testing | Tuberculin skin test | IGRA* | IGRA reversions*, n/N (%) |
|----------------------------------|--------------------------|----------------------|-------|--------------------------|
| Joshi et al (15), 2012, USA      | 2 to 30 days             | N/A                  | N/A   | 18/45 (40)               |
| Rafiza et al (16), 2012, Malaysia | 1 year                   | N/A                  | 69/703 (9.8) | 14/59 (23.7) |
| Fong et al (17), 2012, USA       | 1 year or 1 to 6 months for positive IGRA | N/A                  | 52/1857 (2.8) | 8/10 (80)† |
| Torres Costa et al (18), 2011, Portugal | 1 year | N/A                  | 61/199 (30.7) | 51/462 (11) |
| Schablon et al (19), 2010, Germany | High-risk HCWs tested annually, all others evaluated every other year | Reversion rates: 4/188 (2.1) | 51/462 (11) | 46/208 (22.1) |
| Ringshausen et al (20), 2010, Germany | 18 weeks | N/A                  | 3/162 (1.9) | 13/42 (32.6) |
| Park et al (21), 2010, South Korea | 1 year | N/A                  | 14/244 (5.7) | 6/18 (33.3) |
| Lee et al (22), 2009, South Korea | 1 year        | N/A                  | 21/146 (14.4) | N/A |
| Chee et al (23), 2009, Singapore | 1 year | 16/75 (21.3) | 5/182 (4.9) | N/A |
| Yoshiyama et al (24), 2009, Japan | 2 and 4 years | N/A                  | 5/277 (1.8) | N/A |
| Pollock et al (25), 2008, USA    | 1 to 7 months           | 2/43 (4.6)           | N/A   | 13/32 (41)               |

*All conversions/reversions using simple negative/positive; †Testing was performed among individuals with positive QuantiFERON-TB (Cellestis Ltd, Australia) results close to the cut-off point. IGRA Interferon-gamma release assay; N/A Not available

0.1% among HCWs (15). Yet, more than 3000 employees at the Central Arkansas Veterans Healthcare System alone are expected to undergo LTBI screening every year. Not only does the current testing strategy pose obvious dilemmas in terms of who to treat, there are significant economic costs associated with occupational TB testing programs in North America.

In summary, routine implementation of IGRAs in serial testing programs poses challenges, especially when a test with poorly understood dynamic characteristics is deployed in a predominantly low-risk population. Current guidelines on the use of IGRAs do not adequately address these challenges, nor do they provide specific guidance on how to handle conversions, reversions and borderline responses, and not rely on simplistic cut-offs. As is always the case, it is particularly important to consider clinical context, TST results, if available, and history of exposure in making decisions about preventive therapy. Using an IGRA result alone will not suffice.

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