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Interferon-gamma release assays for tuberculosis screening of healthcare workers: a systematic review

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► Additional material and appendix tables A1 and A2 are published online only. To view these files please visit the journal online (<http://thorax.bmj.com>).

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ABSTRACT

Healthcare workers (HCWs) are at increased risk of exposure to tuberculosis (TB). Traditionally, screening for latent TB infection (LTBI) is done using the tuberculin skin test (TST). Interferon-gamma release assays (IGRAs) are now increasingly being used for diagnosis of LTBI, but their role in HCW screening is unclear. A systematic review was conducted of all IGRA studies in HCWs to summarise their performance in cross-sectional and serial testing settings. By searching four electronic databases and other sources, all available studies using any one of the commercial IGRA assays in HCWs were retrieved and screened. 50 unique studies were identified which met the inclusion criteria including five from high TB incidence settings. Among 24 cross-sectional studies in low TB incidence settings, the pooled prevalence of positive IGRA using either test was significantly lower than for a positive TST. However, in high-incidence settings (n=2) there were no consistent differences in the prevalence of positive tests. IGRAs showed good correlation with occupational risk factors for TB exposure in low-incidence settings. Only 10 studies assessed use of IGRA for serial testing and all showed large variation in the rates of conversions and reversions, with no data suggesting that IGRAs are better at identifying the incidence of new TB infection than the TST. The use of IGRAs instead of TST for one-time screening may result in a lower prevalence of positive tests and fewer HCWs who require LTBI treatment, particularly in low TB incidence settings. However, the use of IGRAs for serial testing is complicated by lack of data on optimum cut-offs for serial testing and unclear interpretation and prognosis of conversions and reversions. Further longitudinal research will be required to inform guidelines on serial testing using IGRAs.

INTRODUCTION

Tuberculosis (TB) continues to have a significant health impact worldwide with an estimated one-third of the world's population infected with latent TB (LTBI). TB poses a significant occupational health problem and healthcare workers (HCWs) are at increased risk of exposure to TB.^{1,2} A systematic review of 51 studies showed that the prevalence and incidence of TB infection and disease were high among HCWs in low and middle income countries.¹ The emergence of extensively drug-resistant TB has resulted in a renewed interest and emphasis on the protection of the healthcare workforce. The World Health Organization (WHO) recently issued a policy on TB infection control (TBIC) in resource-limited settings and is now actively promoting TBIC programmes.³

In many high-income countries, periodic screening of HCWs for LTBI is an important component of TBIC programmes.⁴ Traditionally, the prevalence of LTBI and incidence of new TB infection (ie, conversion) among HCWs has been estimated using the tuberculin skin test (TST), a test with known limitations.⁵⁻⁷ Recently, interferon-gamma release assays (IGRAs) have emerged as alternatives for the diagnosis of LTBI.⁸⁻¹⁰ Two IGRAs are commercially available—the QuantiFERON-TB Gold In-Tube (QFT) assay (Cellestis Ltd, Carnegie, Australia) and the T-SPOT.TB assay (Oxford Immunotec, Abingdon, UK). With the development of new national guidelines incorporating IGRAs, their use is steadily increasing.¹¹

IGRAs have features that make them attractive for repeated screening: they are ex vivo blood-based tests that, in contrast to the TST, can be repeated any number of times without sensitisation or boosting, they require only one visit and do not need a baseline two-step protocol.

There is strong evidence from systematic reviews that IGRAs, especially QFT, have excellent specificity that is unaffected by BCG vaccination, while the T-SPOT.TB shows improved sensitivity for active TB over both the TST and QFT.^{7,12,13} However, reviews have suggested that IGRA performance differs in high versus low TB incidence settings, with relatively lower sensitivity in high-incidence countries.^{8,14}

Despite the substantial body of literature on IGRAs, almost all the available studies have limitations—namely, lack of a gold standard for LTBI, cross-sectional design, use of sensitivity and specificity as surrogates for patient-important outcomes, and lack of adequate data on predictive/prognostic value of IGRAs. In particular, data are lacking on how to interpret repeated (serial) IGRA testing results.^{15,16} Currently, no guidelines exist on the use of IGRAs in countries with a high incidence of TB. Some guidelines from high-income low-incidence countries have not recommended IGRAs for serial testing of HCWs¹⁷ while others state that IGRAs may be used for serial testing of HCWs in place of the TST.^{4,9}

METHODS

Our objective was systematically to review all studies using IGRA to test HCWs, including cross-sectional, longitudinal and serial testing studies, to summarise their performance characteristics. Secondary objectives included (1) to compare IGRA performance in HCWs in high or low TB incidence settings; (2) to determine if IGRAs are better

correlated with occupational exposure to TB than the TST in cross-sectional studies; (3) to estimate the rate of IGRA conversions and reversions and assess whether IGRA conversions are more closely associated with recent occupational exposure than TST conversions; and (4) to summarise the evidence produced by cost-effectiveness analyses and programmatic studies.

Data sources and searches

We have previously published systematic and narrative reviews on IGRA accuracy and performance in various subgroups.^{10 12 13 18} We updated the database searches used in previous systematic reviews and searched the literature for relevant IGRA studies. PubMed, Embase and Biosis and Web of Science were searched and citations of all original articles published in all languages up to 30 April 2010 that reported data on IGRA performance in HCWs (updated again on 1 October 2010 to include the most recent serial testing studies) were reviewed.

In addition to database searches we reviewed bibliographies of previous reviews and guidelines on IGRAs, conference proceedings, abstracts and also screened the citations of relevant original articles. Experts in the field and commercial test manufacturers were contacted to obtain relevant citations. Authors of primary studies were contacted to obtain additional information where necessary. The detailed search string and a list of conferences reviewed for relevant citations are available in the online supplement.

Outcomes evaluated

A major challenge for studies evaluating the performance of IGRAs is the lack of a gold standard for LTBI. We therefore developed a priori a hierarchy of reference standards for the performance of IGRAs in diagnosing LTBI in HCWs (figure 1). At the time of this review, there was no evidence at the highest two levels of the hierarchy nor were there studies looking at sensitivity and specificity for active TB in HCWs. We therefore evaluated studies that reported one or more of the following outcomes: prevalence and incidence of LTBI; correlation between IGRA results and an exposure gradient; and/or agreement between IGRA and TST results.

Details of study selection, inclusion and exclusion criteria, data extraction procedure and quality assessment are given in the online supplement.

Data synthesis and analysis

Study characteristics and results are presented as tables and plots. In order to compare prevalence estimates for the tests (IGRAs

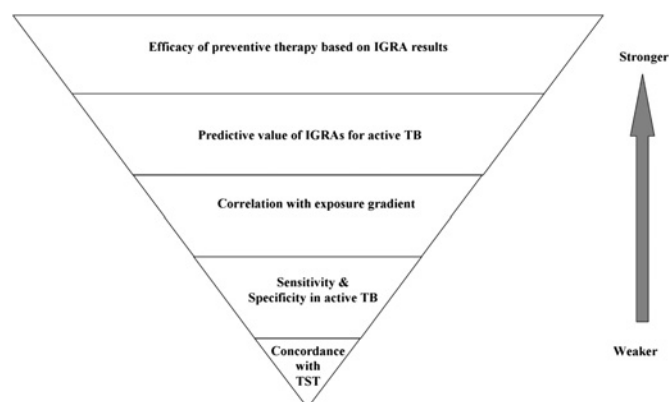


Figure 1 Hierarchy of reference standards for performance of interferon-gamma release assays (IGRAs) for diagnosing latent tuberculosis infection (LTBI) in healthcare workers. TST, tuberculin skin test.

vs TST) we calculated Fisher exact 95% CIs for the prevalence estimates extracted from the original reports. If the 95% CIs did not overlap, differences between proportions were considered as statistically significant, a conservative approach. For the association between occupational risk factors and IGRA we extracted ORs reported by the original authors. When available we extracted both crude and adjusted ORs. Because the included studies varied greatly in their design, execution and outcomes, and because meta-analysis methods are not well defined for such heterogeneous diagnostic studies with no gold standard, we did not perform a meta-analysis. IGRA performance varies across populations⁸ so all the results were stratified by TB incidence in the country where the study was done (high vs intermediate and low incidence). High-incidence countries were defined as countries with more than 100 estimated incident TB cases per year per 100 000 population as determined by the WHO.¹⁹

RESULTS

Description of included studies

Figure 2 shows a flow chart on study selection. The final tables included 42 IGRA studies in HCWs that reported one of our main outcomes of interest. In addition we identified three studies on cost-effectiveness of IGRAs in HCWs and three studies on feasibility and test implementation. Finally, two new serial testing studies were identified by 1 October 2010, giving a total of 50 studies of IGRAs in HCWs.^{20 21} The characteristics of the cross-sectional and serial testing studies identified in this review are shown in tables 1 and 2, respectively. Complete details on study methodology and test performance are given in appendix tables A1 and A2, respectively, in the online supplement. Data are presented stratified by high versus low and moderate TB incidence settings but, even within these strata, the study populations included HCWs with varying risk of TB exposure.

Of the 44 included studies reporting a main outcome, 35 (79%) evaluated QFT only and 3 (7%) used T-SPOT.TB only, while the remaining 6 (14%) evaluated both IGRAs. While most studies performed both an IGRA and the TST, 14% of studies only performed IGRA testing and therefore could not compare IGRA results with TST results. Overall, only 5 (11%) were done in high-incidence settings. Only 10 (23%) used a longitudinal or serial testing design. Study sizes ranged from 12 to 1313 HCWs, for a total of 11 963 HCWs across the 44 studies. Most included BCG-vaccinated HCWs; however, the proportion of BCG-vaccinated HCWs varied considerably (7–100%).

Results of cross-sectional IGRA studies

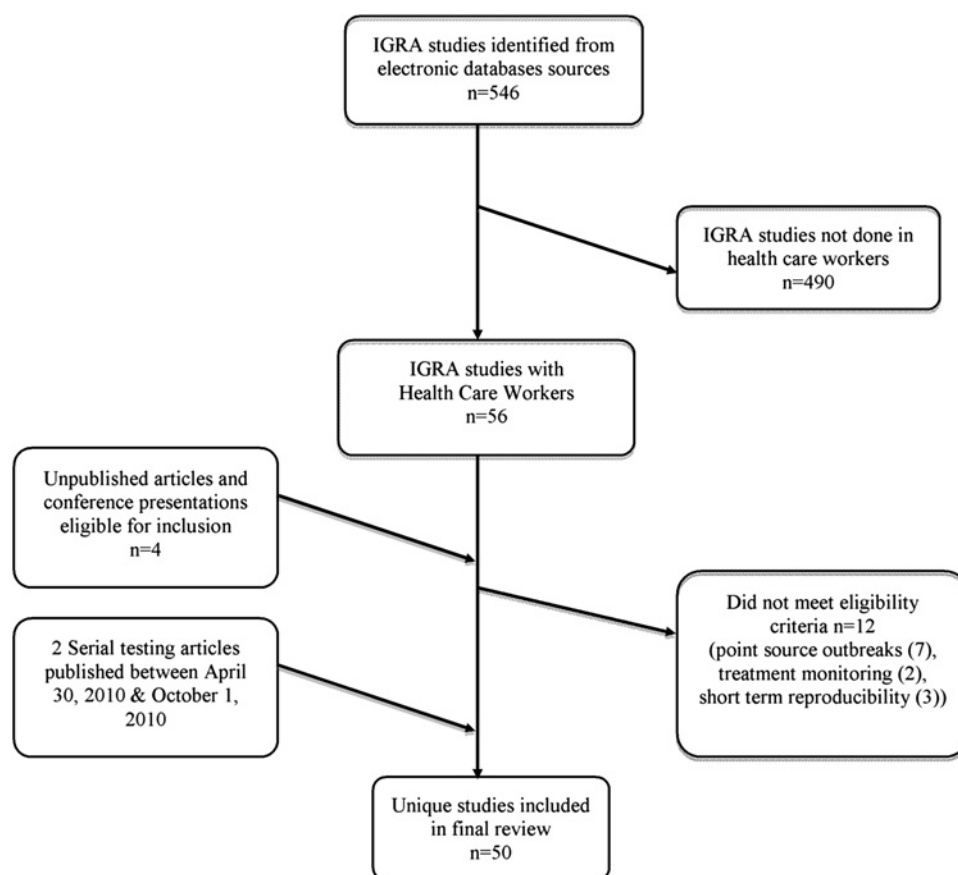
IGRA versus TST positivity rates in HCW populations in high-incidence countries

Three cross-sectional studies evaluated IGRA performance in HCWs in India, Russia and Vietnam, although TST was not performed in the Russian study.^{22–24} As shown in figure 3, TST and IGRA positivity rates were high in HCWs (40–66%); IGRA positivity was slightly lower than TST positivity in the studies in India and Vietnam comparing TST and IGRAs, but the difference in estimated prevalence between the two tests was significant only in Vietnam.²⁴ The Vietnamese study also reported the lowest rate of BCG vaccination among its participants at 37.3% compared with 71% in the Indian study.

IGRA versus TST positivity rates in HCW populations in low- and moderate-incidence countries

We identified 31 cross-sectional studies from low or intermediate TB incidence countries.^{25–52 54–56} Twenty-five studies used only

Figure 2 Flow chart of studies included and excluded at each stage of the review. IGRA, interferon-gamma release assay.



the QFT-Gold test, 2 studies only the T-SPOT.TB and 4 performed head-to-head comparisons of the QFT and T-SPOT.TB tests.^{41 44 50 55} The prevalence of positive QFT ranged from 1% to 66.8% and of positive T-SPOT.TB tests from 1% to 60%. Among 25 studies that compared IGRA with TST, all but one⁴⁴ reported a lower prevalence of positive QFT or T-SPOT.TB than positive TST (figures 3 and 4), with statistically significant differences found in 17 of these 24 studies. The studies in figure 4 are shown in order of increasing proportion of participants BCG vaccinated; studies in which a higher proportion of study participants were BCG vaccinated did not necessarily have a higher prevalence of positive TST or a larger difference between the prevalence of positive TST and positive IGRA.

Concordance was weak between TST and IGRAs in these studies with κ values ranging widely from 0.05 in Denmark²⁹ (using a 12 mm TST cut-off) to 0.56 in a study from Spain⁴⁹ (using a 15 mm TST cut-off for BCG-vaccinated individuals). Three studies evaluated a range of TST cut-offs in the analysis and all showed that agreement could be improved by employing a more stringent TST cut-off (ie, 15 mm vs 10 mm).^{33 46 47} In all 19 studies that reported on discordance, TST+/IGRA- subjects was the predominant type of discordance.^{25 27 29 32–35 37 39 46–52 54–56}

In summary, among these 34 cross-sectional studies the prevalence of positive IGRA was lower than positive TST. The difference in prevalence was significant in low and moderate TB incidence settings (figure 4) but not in high-incidence settings.

Association between occupational risk factors and test results in HCW populations

The majority of studies reporting such analyses were done in low-incidence settings. Of three cross-sectional studies

conducted in high-incidence settings, only Pai *et al* evaluated associations between occupational risk factors and both TST and IGRA. They found a stronger but non-significant association between occupational risk factors and IGRA positivity than the TST.²²

Of the 31 cross-sectional studies conducted in low-incidence countries, 22 evaluated potential risk factors for IGRA and/or TST positivity rates. Fourteen studies^{26 30 32 35 36 40 42 43 46 49 50 52 55 57} reported a positive association between IGRA positivity and occupational risk factors, including higher risk for clinical staff working in a high-risk ward, TB clinic or geriatric care and increased duration of healthcare employment. Two studies^{38 39} reported no association between test positivity and risk factors; the remaining studies either did not perform the TST or did not calculate ORs for risk factors associated with test positivity. The age-adjusted ORs for TST and IGRAs and occupational risk factors for relevant studies are compared in figure 5; all studies are from low- and moderate-incidence countries with the exception of the India study. All three tests correlated well with established indicators of occupational risk of TB exposure, although no test was consistently more often associated with these indicators of exposure. Being of foreign birth or having lived in a high TB incidence country was correlated with QFT positivity in four studies,^{35 40 46 48} and three of the four studies also showed correlation with TST positivity while the fifth study to do this using the T-SPOT.TB test—found no association with place of birth although there was an association for TST.³⁹ From these cross-sectional studies, IGRAs appear to be well correlated with TB infection risk factors (including occupational risk factors) in HCWs in low- and intermediate-incidence countries.

Review

Table 1 Characteristics of 34 cross-sectional IGRA studies in HCWs

Study	Year	Country	N	TST (PPD dose)	IGRA (QFT, TSPOT.TB or both)	BCG vaccinated (%)
High TB incidence countries						
Pai <i>et al</i> ²²	2005	India	726	1 TU	QFT-G-IT	71
Drobniewski <i>et al</i> ²³	2007	Russia	500	—	QFT-G-IT	84
Lien <i>et al</i> ²⁴	2009	Vietnam	300	5 TU	QFT-G-IT	37.3
Intermediate and low TB incidence countries						
Kang <i>et al</i> ²⁵	2005	Republic of Korea	171	2 TU	QFT	92.3
Harada <i>et al</i> ²⁶	2006	Japan	332	2.5 TU	QFT-G	91.3
Ozekinci <i>et al</i> ²⁷	2007	Turkey	66	5 IU	T-SPOT.TB	67
Veeser <i>et al</i> ²⁸	2007	USA	55	—	QFT-G	45
Soborg <i>et al</i> ²⁹	2007	Denmark	139	2 TU	QFT-G	76
Nienhaus <i>et al</i> ³⁰	2007	Germany	454	—	QFT-G-IT	42
Nienhaus <i>et al</i> ³¹	2007	Germany	161	2 TU	QFT-G-IT	36
Mirtskhulava <i>et al</i> ³²	2008	Georgia	265	5 TU	QFT-G-IT	77.7
Hotta <i>et al</i> ³³	2007	Japan	207	3 TU	QFT-TB-2G	92% (48% with >1 BCG)
Nienhaus <i>et al</i> ³⁴	2008	Germany	261	2 TU	QFT-G-IT	37.5
Ciaschetti <i>et al</i> ³⁵	2007	Italy	590	—	QFT-G	56
Eum <i>et al</i> ³⁶	2008	Republic of Korea	73	—	QFT-GIT	100
Choi <i>et al</i> ³⁷	2008	Republic of Korea	84	2 TU	QFT-G	100
Carvalho <i>et al</i> ³⁸	2008	Italy	65	5 IU	QFT-G	85
Barsegian <i>et al</i> ³⁹	2008	Germany	95	—	T-SPOT.TB	36
Stebler <i>et al</i> ⁴⁰	2008	Switzerland	777	—	QFT-G-IT	87.4
Thijssen <i>et al</i> ⁴¹	2008	The Netherlands	19*	—	QFT-G-IT and T-SPOT.TB	16
Demkow <i>et al</i> ⁴²	2008	Poland	155	—	QFT-G	100
Schablon <i>et al</i> ⁴³	2009	Germany	270	—	QFT-G-IT	52.8
Dorman <i>et al</i> ⁴⁴	2009	USA	1313	—	QFT-G-IT and T-SPOT.TB	—
Mehta <i>et al</i> ⁴⁵	2009	USA	12†	—	QFT-G	—
Vinton <i>et al</i> ⁴⁶	2009	Australia	481	10 IU	QFT-G-IT	78
Zrinski Topić <i>et al</i> ⁴⁷	2009	Croatia	54	2 TU	QFT-G-IT	100
Khanna <i>et al</i> ⁴⁸	2009	UK	171	2 TU	QFT-G-IT	82.5
Álvarez-León <i>et al</i> ⁴⁹	2009	Spain	134	2 TU	QFT-G-IT	35
Casas <i>et al</i> ⁵⁰	2009	Spain	147	2 TU	T-SPOT.TB & QFT-G-IT	16
Fox <i>et al</i> ⁵¹	2009	Israel	100	5 PPD	QFT-G-IT	37
Costa <i>et al</i> ^{52, 53 *}	2009	Portugal	1218	2 TU	QFT-G-IT	100
Zhao <i>et al</i> ⁵⁴	2009	USA	40	—	QFT-G-IT	—
Girardi <i>et al</i> ⁵⁵	2009	Italy	115	5 IU	T-SPOT.TB and QFT-G-IT	37.4
Cummings <i>et al</i> ⁵⁶	2009	USA	182	—	QFT-G-IT	7

*Two studies based on same cohort, ERJ data displayed in table.

†All subjects were TST converters at recruitment.

BCG, Bacille Calmette-Guerin vaccine; HCW, healthcare worker; IGRA, interferon-gamma release assay; IU, international unit; PPD, purified protein derivative; QFT, QuantiFERON test; QFT-G, QFT Gold test; QFT-G-IT; QFT Gold In-Tube test; TB, tuberculosis; TST, tuberculin skin test; TU, tuberculin unit.

Longitudinal serial testing IGRA studies and their results**IGRA conversion and reversion rates in HCW populations in high-incidence countries**

We identified only two serial testing studies from high-incidence settings. Studies conducted repeat testing at 0, 6 and

12 months⁵⁸ and 0 and 18 months (table 2).¹⁵ The rates of IGRA conversions from these two studies ranged from 11.6% to 21%. The study by Pai *et al* was the only study to calculate the TST conversion rate and found a 4% rate after 18 months.¹⁵ Pai *et al* also found conversions rates varied for both the TST and the

Table 2 Characteristics of eight longitudinal and serial testing IGRA studies in HCWs

Study	Year	Country	N	TST (PPD dose)	IGRA (QFT, TSPOT.TB or both)	Timing between repeat testing
High TB incidence countries						
Pai <i>et al</i> ¹⁵	2006	India	216	1 TU	QFT-G-IT	18 months
Joshi <i>et al</i> ⁵⁸	2009	India	79	—	QFT-G-IT	0, 6 and 12 months
Intermediate and low TB incidence countries						
Pollock <i>et al</i> ⁵⁷	2009	USA	143 ²⁵	5 TU	QFT-G-IT and T-SPOT.TB	1–7 months
Zwerling <i>et al</i> ⁵⁹	2009	Canada	117	5 TU	QFT-G-IT	1 year
Yoshiyama <i>et al</i> ⁶⁰	2009	Japan	311	—	QFT-G	2 and 4 years
Chee <i>et al</i> ⁶¹	2009	Singapore	207	2 TU	T-SPOT.TB	1 year
Lee <i>et al</i> , ICHE ⁶²	2009	Republic of Korea	196	2 TU	QFT-G	1 year
Belknap <i>et al</i> ⁶³	2010	USA	1281	5 TU	QFT-G-IT and T-SPOT.TB	6 months
Costa <i>et al</i> ²⁰	2010	Portugal	670	2 TU	QFT-G-IT	1–2.5 years
Ringshausen <i>et al</i> ²¹	2010	Germany	182	2 TU	QFT-G-IT	18 weeks

HCW, healthcare worker; IGRA, interferon-gamma release assay; IU, international unit; PPD, purified protein derivative; QFT, QuantiFERON test; QFT-G, QFT Gold test; QFT-G-IT; QFT Gold In-Tube test; TB, tuberculosis; TST, tuberculin skin test; TU, tuberculin unit.

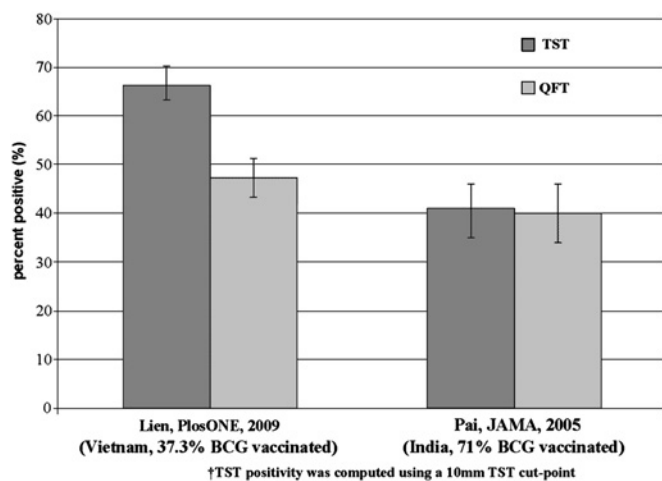


Figure 3 Cross-sectional studies in countries with a high incidence of tuberculosis (N=2). QFT, QuantiFERON-TB Gold In-Tube; TST, tuberculin skin test.

IGRA when different cut-offs were used. Neither study reported data to suggest that IGRA conversions were better associated with TB exposure than TST conversions.

Reversion rates in the study by Joshi *et al* ranged from 27% in the first 6-month period to 40% in the second 6-month period.⁵⁸ Pai *et al* reported 18-month IGRA reversion rates around 7% among baseline concordant positives, but up to 70% among those with discordant baseline results (ie, TST-/IGRA+).¹⁵

IGRA and TST conversion and reversion rates in HCW populations in low-incidence countries

Four studies have been recently published in this area, while two others were presented at conferences in 2009 and 2010 (table 2).^{59–64} As summarised in table 3, conversion rates ranged from 1.8% (5/277) in Japan⁶⁰ (testing every 2 years) to 14.4% (21/146) in Korea⁶² (testing every year). A study by Lee *et al*⁶² was the only one to report a higher TST conversion rate than IGRA conversion rate (21% vs 14%). Yoshiyama *et al* in Japan⁶⁰

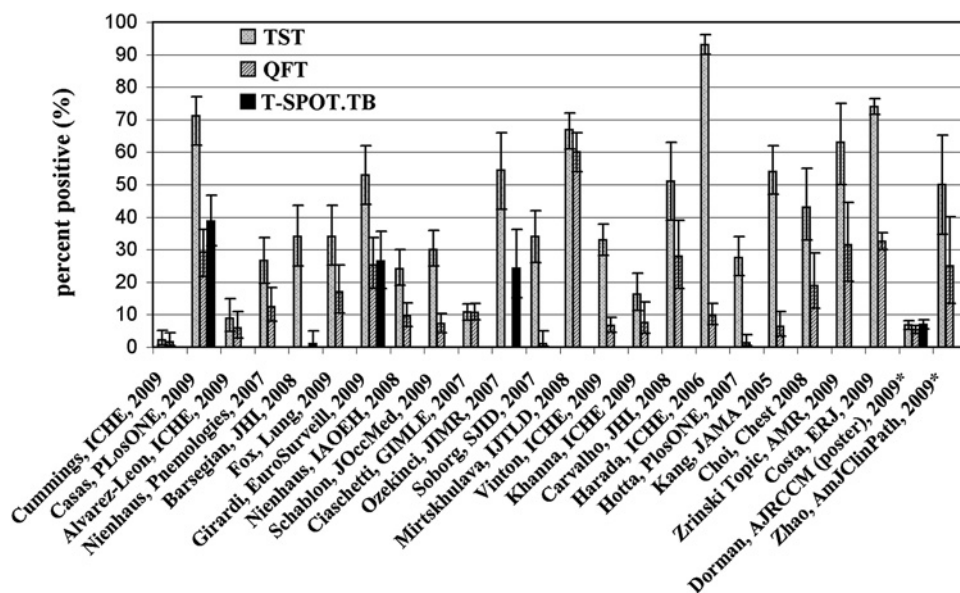
and Pollock *et al* in the USA⁶⁴ found IGRA conversions were associated with TB exposure; however, neither performed repeat TST. Belknap *et al* found IGRA conversion rates were associated with older age and male gender but not occupational exposure to patients with active TB.⁶³ In the only published study to examine the association of IGRA and TST conversions with exposure to TB,⁶² occupational TB exposure was not associated with TST or IGRA conversions.

Three studies from low- and moderate-incidence settings including Japan,⁶⁰ Canada⁵⁹ and the USA reported IGRA reversion rates of 40–52.9%.^{63–65} In the Japanese study, where participants were tested at baseline, at 2 years and at 4 years, all reversions had initial interferon-gamma values close to the cut-off.⁶⁰ This study also showed that QFT conversion was associated with working in a TB ward. A study from Canada found half of the subjects with discordant baseline results (QFT +/TST-) reverted to negative QFT at 1 year. TST was not repeated if the participant was TST+ so these studies were not able to estimate TST reversions.

The two most recent serial testing publications in HCWs confirm these early findings. Ringshausen *et al* in Germany reported a large reversion rate (6/18, 33%) while only 3/162 (1.9%) reported IGRA conversions and demonstrated that, when continuous IGRA results were employed, higher baseline results were more stable than those closer to the cut-off.²¹ Costa *et al* found conversion rates of 3.6–11% and reversion rates of 5.2–22% depending on the cut-off used. Using simple negative to positive cut-offs whether for reversions or conversions always gave the largest estimates (11% and 22%, respectively).

Overall, serial testing data from low-incidence countries suggest that IGRA results vary greatly during serial testing, and rates of conversions and reversions vary depending on the test used and the cut-off definition used. When simple negative/positive changes are used as cut-offs, IGRAs had higher rates of reversions and conversions which were frequently higher than the TST. Owing to the limited number of studies evaluating conversions and a relationship between exposure, there are no data to show that IGRAs are better at identifying the incidence of new TB infection than the TST.

Figure 4 Cross-sectional studies comparing tuberculin skin test (TST) and interferon-gamma release assays (IGRAs) in countries with a low and intermediate incidence of tuberculosis (N=24) in order of increasing proportion of BCG vaccinated. QFT, QuantiFERON-TB Gold In-Tube.

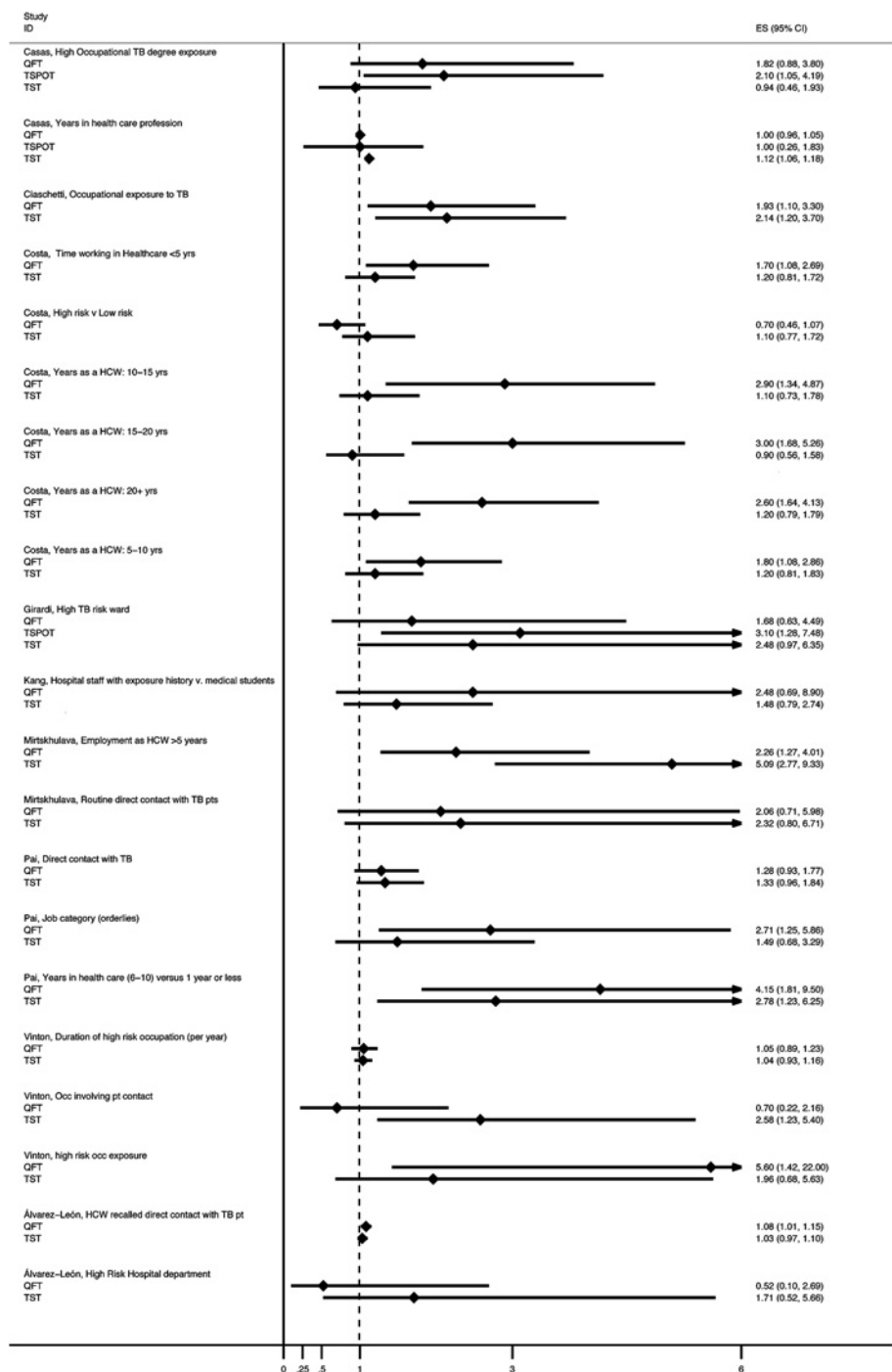


*proportion BCG vaccinated was not reported for these studies

TST positivity was computed using the TST cut-point used by authors of the primary studies.

When results for multiple cut-points were presented, data for TST cut-point 10mm was displayed

Figure 5 Occupational exposure to TB as a risk factor for tuberculin skin test (TST) and interferon-gamma release assay (IGRA) positivity: comparison of multivariate ORs. QFT, QuantiFERON-TB Gold In-Tube.



Information on secondary outcomes including cost-effectiveness, feasibility and implementation are available in the online supplement.

DISCUSSION

While the TST has been successfully used in TBIC programmes, the availability and growing use of IGRAs raises the issue of whether IGRAs could replace the TST for screening of HCWs. Given the repeated nature of routine screening, there are particular issues which may not be relevant in routine practice or contact investigations but become very important when contemplating serial screening. These include test reproducibility, performance of IGRA when repeated frequently,

interpretation of discordant TST and IGRA results and the interferon-gamma thresholds (cut-off values) which most accurately distinguish new TB infection (ie, conversion) from random variation. Despite the paucity of data on the above critical issues, some countries such as the USA have recommended the use of IGRAs for HCW screening.⁹ In contrast, other guidelines (eg, Canada, Australia, UK)^{17 66-68} have been cautious. In June 2010 the US Centers for Disease Control (CDC) released an updated guideline on IGRAs.⁶⁹ This guideline is more cautious about serial testing with IGRAs and emphasises that using a 'lenient criterion to define IGRA conversion might produce more conversions than are observed with the more stringent criteria applied to TSTs. Furthermore, an association between an IGRA conversion and subsequent disease risk has

Table 3 Summary of rates of conversions and reversions in serial testing studies (N=8)

Study	Duration between testing	TST converters, n/N (%)	IGRA converters, n/N (%)	IGRA reverters, n/N (%)
High TB incidence countries				
Pai, 2006 ¹⁵	18 months	6/147 (4.1%)	17/147 (11.6%)	7/38 (18.4%)
Joshi, 2009 ⁵⁸	6 months	—	11/57 (19%)	6/22 (27%)
	6 months (6–12 months)	—	11/52* (21%)	11/27† (40%)
Moderate and low TB incidence countries				
Pollock, 2009 ⁶⁴	1–7 months	—	2/43‡ (4.6%)	—
Zwerling, 2009 ⁵⁹	1 year	0/57 (0%)	4/56 (7.14%)	4/5 (80%)
Yoshiyama, 2009 ⁶⁰	2 and 4 years	—	5/277 (1.8%)	13/32 (41%)
Chee, 2009 ⁶¹	1 year	0/18‡	9/182 (4.9%)	—
Lee, 2009 ⁶²	1 year	16/75 (21.3%)	21/146 (14.4%)	—
Belknap, 2010 ⁶³	6 months	4/1202 (0.3%)	TSPOT 44/1117 (3.9%) QFT-GIT 44/1169 (3.8%)	TSPOT 36/68 (52.9%) QFT-GIT 20/50 (40%)
Costa, 2010 ²⁰	1–2.5 years	98/199 (49.2%)	51/462 (11%)	46/208 (22.1%)
Ringshausen, 2010 ²¹	18 weeks	(baseline only)	3/162 (1.9%)	6/18 (33.3%)

All conversions/reversions using simple negative/positive definition.

*Denominator includes only participants negative at 6 months.

†Denominator includes only participants positive at 6 months.

‡Denominator includes only baseline concordant negatives.

IGRA, interferon-gamma release assay; PPD, purified protein derivative; TB, tuberculosis; TST, tuberculin skin test.

not been demonstrated. The criteria for interpreting changes in an IGRA that identify new infections remain uncertain.⁶⁹ Likewise, the most recent 2010 Canadian guidelines also called for caution in the use of IGRAs in serial testing.⁷⁰ In December 2010, the WHO released a STAG report including recommendations which discourage the use of IGRAs, “for the detection of latent TB infection (LTBI) in adults, children, health-care workers, contacts and those involved in outbreak investigations in low-income and middle-income countries.”⁷¹ This move towards more cautious guidelines in the domain of serial testing is supported by findings in our review, and reinforces the need for an addendum to existing guidelines on IGRAs that specifically address issues related to interpretation of conversions and reversions.

Our systematic review provides several useful insights into the performance of IGRAs in HCWs. The observed prevalence of LTBI in HCWs depends on the test used and the particular TB incidence setting. In low-incidence countries IGRAs estimate a significantly lower prevalence of LTBI than the TST. Some attribute such discordance to the higher specificity of IGRAs compared with the TST and, indeed, agreement and κ values are improved in non-BCG-vaccinated individuals compared with BCG-vaccinated, although there is not a strong trend (figure 4).

Studies have suggested that the lower prevalence of LTBI using IGRAs will result in fewer numbers of HCWs who require preventive therapy. However, the higher rate of subsequent conversions found by IGRA in these studies suggests that, while fewer individuals may be identified as LTBI at baseline, more individuals could be diagnosed with conversions by IGRA leading to more HCWs requiring preventive therapy upon repeated screening. This finding has major implications for TBIC policies and relevant cost-effectiveness analyses.

Along with high conversion rates, studies reported high rates of subsequent reversions. This poses concerns of both a scientific and clinical nature. What biological phenomenon is at work here? Are these individuals clearing the infection naturally? Is reinfection an issue in high TB incidence settings? Given a high expected reversion rate, should individuals with positive results be treated, or tested again at a later date? Treating individuals who might have reverted in the absence of treatment could create an environment where more individuals take potentially harmful preventive therapy unnecessarily. This has raised much

interest in what is an appropriate definition for an IGRA conversion or reversion.

As summarised in a recent systematic review,¹⁸ within-subject reproducibility of IGRAs is moderate and a previous TST can potentially boost subsequent IGRA results. IGRAs are dynamic assays and interferon-gamma values tend to fluctuate around the cut-off and cause apparent conversions and reversions.^{72–74} The exact cause of the conversions and reversions remains unclear, and might indicate spontaneous clearance of TB infection, reinfection or dynamic changes within the spectrum of latent TB infection.^{75 76} Conversion rates are highest when a simple negative to positive change is used to define a conversion. This is also true for reversion rates where a simple positive to negative change is used as the definition. Our review suggests that this is true in both high- and low-incidence settings and has implications for deciding on criteria (cut-offs) for conversions and reversions. Alternatives to a simple negative to positive definition for conversion have been proposed by others, including definitions involving an absolute increase over baseline (similar to the TST), a proportionate increase over baseline or a proposed ‘grey zone’.^{73 74} These more stringent definitions may lead to smaller rates of conversions and reversions; however, it remains to be seen which conversion definition will be most strongly associated with TB exposure or subsequent disease development.

Studies found good correlation between occupational risk factors and positivity rates, but very few studies have looked at an association between IGRA test conversion and known occupational exposure or progression to disease. Without these, the interpretation and prognosis of conversions and reversions remains unclear.

Strengths and limitations of the review

Our systematic review had several strengths. We employed a comprehensive search strategy using multiple sources and databases to retrieve relevant studies, including unpublished studies and conference proceedings. Two review authors independently assessed eligible articles for inclusion. Owing to heterogeneity in study designs and outcomes assessed in each study, it was not appropriate to pool the data, and instead we analysed study results in subgroups by study design and by background TB incidence.

Despite the comprehensive search, we may have missed relevant studies and publication bias is always a concern. Furthermore, very few studies reported prevalence of HIV among their study populations. Lastly, there is lack of evidence at the highest level of the hierarchy of reference standards; a majority of the included studies were cross-sectional and predominantly from low TB incidence settings. Serial testing data, evidence on predictive value in HCWs and reproducibility data are still quite limited.

Research directions and implications

Until further evidence is available, TBIC programmes that include IGRA testing must use caution, as emphasised by recent US and Canadian guidelines.^{69 70} In particular, TBIC programmes may observe higher conversion numbers with IGRAs, and health professionals should be cautious about using a simplistic negative-to-positive definition of conversion, and instead consider the amount of change in absolute interferon-gamma responses as well as relevant clinical information (eg, likelihood of exposure or contact and concurrent TST results, if available) to detect and treat conversions. This is particularly relevant for individuals with borderline results because these results are most likely to change upon retesting.

Current guidelines and evidence available on the use of IGRAs do not adequately address questions raised by serial testing, nor do they provide the guidance or understanding needed to properly interpret IGRA results in serial testing. With rapidly accumulating evidence from serial testing studies, existing IGRA guidelines will need to be updated with specific recommendations on interpretation of serial testing results.

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WEB-ONLY SUPPLEMENT FOR:

Interferon-gamma release assays for tuberculosis screening of healthcare workers: a systematic review

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METHODS

Study Selection

We included studies that used a commercial IGRA assay: QuantiFERON-TB Gold® or In-Tube version and the T-SPOT.TB, for TB screening in HCWs in any setting (high or low incidence). We included cross-sectional, longitudinal and serial testing designs.

The following studies were excluded: 1) case reports and case series, 2) studies with 10 or fewer participants, 3) reviews and commentaries; 4) letters that did not report original data; 5) studies evaluating the use of IGRAs for treatment monitoring in HCWs (ie: not diagnostic purposes); 6) short term serial testing studies (serial testing within one month) and reproducibility studies (which have been systematically reviewed recently), [1] 7) non-commercial/in-house assays, and finally 8) IGRA testing in the context of a known nosocomial outbreak or point source exposure. We would expect these studies to report higher rates of conversions and reversions by both tests, and they may not reflect the typical level of LTBI prevalence or conversions in an occupational environment.

Data extraction and Quality assessment

Two review authors (AZ and MP) independently assessed eligible articles for inclusion, and disagreements were resolved by consensus. All included articles were evaluated by a reviewer (AZ), who extracted data that included study design, participants, country, period of recruitment, proportion BCG vaccinated, IGRA methods, TST methods and outcome data, including: baseline TST and IGRA positivity rates, indeterminate rates, concordance between TST and IGRA (agreement and kappa [agreement adjusted for chance]), predominant type of discordance and correlations found between risk factors and test results. A second reviewer (MP) verified a subset of the extracted data (approximately one-third of all studies).

Because IGRA studies in HCWs do not use the conventional diagnostic study design for sensitivity and specificity estimation, we chose a few design features as quality indicators. These included study design (cross-sectional vs longitudinal), use of standardized, commercial assays, use of standardized tuberculin material (PPD-S in North America, RT23 in Europe), proportion of indeterminate results, and duration of follow-up in longitudinal studies.

Database Search String:

The search terms used in database searching included: ((interferon-gamma release assay*) OR (T-cell-based assay*) OR (antigen-specific T cell*) OR (T cell response*) OR (T-cell response*) OR (interferon*) OR (interferon-gamma) OR (gamma-interferon) OR (IFN) OR (elispot) OR (ESAT-6) OR (CFP-10) OR (culture filtrate protein) OR (Enzyme Linked Immunosorbent Spot) OR (Quantiferon* OR Quantiferon-TB)) AND ((tuberculosis OR mycobacterium tuberculosis)).

Detailed outcomes evaluated

In cross-sectional studies, the potential outcomes of interest were: a) Prevalence of positive TST (i.e. LTBI prevalence) versus positive IGRA in HCWs, and risk factors associated with prevalence of positive TST or IGRA, and b) Concordance (i.e. agreement) between TST and IGRA in HCWs; and factors associated with concordance and discordance (e.g. BCG status).

In longitudinal studies, the potential outcomes of interest were: a) Incidence of TST and IGRA conversions and risk factors for conversions, b) Incidence of TST and IGRA reversions and risk factors for reversions.

Conferences reviewed for relevant publications:

- 40th Union World Conference on Lung Health, December 2009
- American Thoracic Society International Conference, 2009
- 2nd Global Symposium on IGRAs, 2009

RESULTS

Table A1. Methodology and results of cross-sectional IGRA studies among HCWs, stratified by TB incidence [N = 34 studies]

Study, Journal, Year & Country	Date of Subject Recruitment	N	TST (PPD dose)	IGRA (QFT or TSPOT or both)	Performance and outcome data						
					BCG vaccinated (%)	TST positivity n/N (%) [TST cut-point]	IGRA Positivity rate n/N (%)	Indeterminate rate n/N (%)	TST & IGRA Concordance (% agreement and kappa)	Predominant type of discordance	Correlation between risk factors (exposure) and test results
High Incidence Countries											
Pai et al, JAMA 2005, India [2]	Jan – May 2004	726	1 TU	QFT-G-IT	71%	298/720 (41%) [≥10mm]	291/725 (40%)	-	81.4% κ= 0.61 (95% CI:0.56-0.67)	+TST/-QFT (But very close)	-Both TST and QFT was associated with more years working in health care and increasing age (different age categories were significant for TST and IGRA). Being an orderly was associated with QFT only.
Drobniewski et al, Plos Med 2007, Russia[3]	Oct 2004 – Oct 2005	500	-	QFT-G-IT	84%	-	131/500 (26.2%)	-	-	-	- Pos QFT higher in doctors and nurses than medical students, especially those in TB service. - TB service HCWs has much higher risk than primary health HCWs. -No difference between risk for nonmedical and medical students.
Lien et al, PlosONE 2009, Vietnam[4]	Nov 2007	300	5 TU	QFT-G-IT	37.3%	191/288 (66.3%) [≥10mm]	142/300 (47.3%)	35/300 (11.6%)	72.5% κ=0.44	+TST/-QFT	-Working in a TB hospital, less than university education, and high body mass were all associated with QFT positivity.

Intermediate & Low Incidence Countries

Kang et al, 2005 JAMA, Republic of Korea[5]	Feb 2004-2005	171	2 TU	QFT	92.3%	93/171 (54%) [≥10mm]	11/171 (6.4%)	-	κ=0.16	+TST/-QFT	-
Harada et al, ICHE 2006, Japan[6]	Mar – Apr 2003	332	2.5 TU	QFT-G	91.3%	283/304 (93.1%) [≥10mm]	33/332 (9.9%)	-	-	-	- Pos QFT results were associated with increased age and with a history of working in a TB ward or an outpatient department of a tuberculosis clinic. - Pos TST results were not correlated with any of the risk factors evaluated.
Ozekinci et al, JIntMedRes, 2007, Turkey[7]	Jan-June 2006	66	5IU	T-SPOT.TB	67%	36/66 (54.5%) [≥10mm in BCG vaccinated otherwise ≥15mm]	16/66 (24.2%)	-	63.6% κ=0.305	+TST/-TSPOT	-
Veesser et al, JACH 2007, USA[8]	June 2005-Aug 2006	55	-	QFT-G	45%	-	2/55 (3.6%)	5/55 (9%)	-	-	-
Soborg et al, Scan JID 2007, Denmark[9]	2005	139	2 TU	QFT-G	76%	47/139 (34%) [≥12mm]	2/139 (1%)	-	κ=0.05 (95%CI :0.0009 – 0.11)	+TST/-QFT	-BCG vaccination only factor significantly associated with TST positivity.
Nienhaus et al, (German) Pneumologie 2007, Germany[10]	-	454	-	QFT-G-IT	42%	-	53/454 (11.7%)	-	-	-	-Strong positive association between Pos QFT and older age. -Weak assoc with work in Geriatric care

Nienhaus et al, (German) Pneumologie 2007, Germany[11]	Dec 2005	161	2TU	QFT-G-IT	36%	43/161 (26.7%) [≥5mm]	20/161 (12.4%)	-	-	+TST/-QFT	-Positive association between increasing age and pos QFT, but not TST. -Pos TST associated with foreign birth.
Mirtskhulava et al, IJTLD 2008, Georgia[12]	June – Aug 2006	265	5 TU	QFT-G-IT	77.7%	177 /265 (66.8%) [≥10mm]	159/265 (60%)	-	50.2 % κ =0.43 (95%CI:0.33-0.55)	+TST/-QFT	-Length of employment as a HCW for ≥5 years was associated with both QFT and TST positivity, while age >30 years, was associated with QFT only.
Hotta et al, Plos ONE 2007, Japan[13]	May – June 2006	207	3TU	QFT-TB-2G	92% (48% with >1 BCG)	57/207 (27.5%) [≥15mm]	3/207 (1.4%)	5/207 (2.4%)	5 mm: 18.8% 10mm: 41.1% 15mm: 72.5% κ = 0.007 (95%CI:0.02 – 0.077)	+TST/-QFT	-
Nienhaus et al, IAOccEH 2008, Germany[14]	Dec 2005-Aug 2006	261	2 TU	QFT-G-IT	37.5%	63/261 (24.1%) [≥5mm]	25/261 (9.6%)	-	77.7% κ = 0.24 (p=0.001)	+TST/-QFT	-Older age associated with IGRA positivity.
Ciaschetti et al, (Italian) GI tal Med Lav Erg 2007, Italy[15]	2006-2007	590	-	QFT-G	56%	64/590 (10.8%) [≥10mm]	63/590 (10.7%)	-	-	+TST/-QFT (Very Close)	-Both tests were significantly associated with non-Italian nationality age ≥45 years old, history of household contacts, and occupational exposure to TB patients
Eum et al, DiagMicroID 2008, Rep. of Korea[16]	-	73	-	QFT-GIT	100%	Using whole blood assay stimulated by PPD, 100% nurses were positive	5/48 med students (10.4%) 13/25 nurses (52%)	-	-	-	-Significant positive correlation was found with duration worked among nurses and IGRA positivity but not with TST positivity.
Choi et al, CHEST 2008,	Aug 2006-	84	2 TU	QFT-G	100%	36/82 (43.9%)	16/82 (19.5%)	2/82 (2.4%)	67.5% κ=0.031	+TST/-QFT	-Main objective of study was boosting of TST

Rep. of Korea[17]	Dec 2006					[≥10mm]			(95%CI:0.22-0.4)		
Carvalho et al, JHospInf 2008, Italy[18]	Feb 2007	65	5 IU	QFT-G	85%	33/65 (51%) 26/33 were previous TST+ [≥10mm + increase ≥10mm]	18/65 (28%)	-	52% κ=0.05	*Nurses had more discordant results than doctors	-No significant association between QFT and gender, professional category or BCG status.
Barsegian et al, JHospInf 2008, Germany[19]	Oct-Dec 2006	95	-	T-SPOT.TB	(36%)	32/95 (34%) [≥5 mm]	1/95 (1%)	-	-	+TST/-TSPOT	-Significant association between birthplace and Pos TST. -T-SPOT.TB results were independent of foreign birth, prior TST, and prior BCG vaccination
Stebler et al, ICHE 2008 Switzerland[20]	June 2005-May 2006	777	-	QFT-G-IT	87.4%	-	59/ 777 (7.6%)	-	-	-	-Country of origin and working in a high risk location for TB exposure was associated with QFT positivity
Thijssen et al, JOEM 2008, Netherlands[21]	2005	19*	-	QFT-G-IT & T-SPOT.TB	16%	All TST >5mm at recruitment	3/10 QFT (30%) 5/18 TSPOT (28%)	-	-	-	*All TST at recruitment
Demkow et al, JPhysPharm 2008, Poland[22]	-	155	-	QFT-G	100%	On average: 42/155 (27.1%) [≥10mm]		-	Correlation between diameter of skin test and IFN-g was highly significant (p<0.001)	-	-QFT pos associated with TB labs and clinics and nurses. -QFT pos was associated with age and duration of employment
Schablon et	Dec	270	-	QFT-G-IT	52.8%	-	19/270	-	-	-	- QFTpos were associated with older

al, JOccMed Tox 2009, Germany[23]	2005-Jan 2008					(30% Previous positive)	(7.2%)				age, being a physician or nurse, and having no previous TST on record (compared with TST – on record).
Dorman et al. AJRCCM 2009 (conference poster), USA[24]	-	1313	-	QFT-G-IT & T-SPOT.TB	-	89/1313 (6.8%)	5.3% QFT 6.9% TSPOT	2.7% QFT 0.8% TSPOT (Failed: 1.1% TSPOT & 3.8% TSPOT borderline)	-	-	-
Mehta et al. CVI 2009, USA[25]	2006	12**		QFT-G	-	100% [≥10mm increase over previous year]	1/12 (8%)	1/12 (8%)	-		**All TST converters at recruitment
Vinton et al, ICHE 2009, Australia[26]	-	481	10IU	QFT-G-IT	78%	10mm: 120/364 (33%) 15mm 73/364 (20%) 20mm: 39/364 (10.7%)	32/481 (6.7%)	8/481 (1.6%)	10mm:71% $\kappa = 0.16$ 15mm:82% $\kappa = 0.23$ 20mm:89% $\kappa = 0.25$	+TST/-QFT	-Birth in high prevalence country, the number of years living in TB endemic country, and high risk occupation were all associated with higher odds of having a positive QFT. -Number of years in a TB endemic country, having received BCG vaccinations and an occupation involving patient contact were all associated with TST positivity (10mm cut-point).
Zrinski Topić et al, ArchMedRes 2009,	June - Sept 2007	54	2 TU	QFT-G-IT	100%	5 mm: 45/54 (83%) 10mm:	17/54 (31.5%)	-	15mm: 74% $\kappa = 0.418$ (95% CI: 0.155-0.68)	+TST/-QFT	-HCWs with positive QFT were 5 years older (p=0.05)

Croatia[27]						34/54 (63%) 15mm: 19/54 (35%)					
Khanna et al, ICHE 2009, United Kingdom[28]	-	171	2 TU	QFT-G-IT	82.5%	24/148 (16.2%) [≥15mm]	13/171 (7.6%)	-	κ=0.44 (95% CI:0.29-0.589)	+TST/-QFT	-Birth in high prevalence country was associated with QFT and TST positivity, BCG vaccination and age were not associated with TST or QFT.
Álvarez-León et al, ICHE 2009, Spain[29]	May - Sept 2007	134	2 TU	QFT-G-IT	35%	12/134 (8.9%) [≥5mm in BCG non-vaccinated, otherwise ≥15mm]	8/134 (5.9%)	3/134 (2.2%)	94% κ=0.56 (95% CI:0.27-0.85)	+TST/-QFT	-Only significant risk factor for +TST was working as an orderly, while older age and direct contact with patients were risk factors for QFT positivity.
Casas et al, PLoSONE 2009, Spain[30]	Nov 2004 - Jul 2005	147	2 TU	T-SPOT.TB & QFT-G-IT	16%	103/147 (71.1%) [≥5mm in BCG non-vaccinated, otherwise ≥15mm]	TSPOT 57/147 (38.7%) QFT 42/147 (29.3%)	TSPOT 2/147 (1.4%) QFT 2/147 (1.4%)	TSPOT 62.9% κ=0.32 QFT 58.7% κ=0.288	+TST/-IGRA	-In non BCG vaccinated subjects, the only risk factor to be significantly associated with TSPOT.TB in multivariate analyses was high occupational exposure to TB.
Fox et al., Lung 2009, Israel[31]	-	100	5 PPD	QFT-G-IT	37%	34/100 (34%) [≥10mm]	17/100 (17%)	9/100 (9%)	κ=0.19	+TST/-QFT	-Only significant factor associated with TST positivity in multivariate analysis is BCG vaccination (OR=4.69) Nothing associated significantly with QFT positivity.
Costa et al, ERJ 2009, Portugal[32, 33]*	May 2005- Sept 2008	1218	2 TU	QFT-G-IT	100%	903/1218 (74%) [≥10mm]	397/1219 (32.6%)	1/1219 (0.08%)	54% κ=0.22	+TST/-QFT	-Probability of positive IGRA increased with age, TST induration and years as a HCW, only ≥3 BCG was associated significantly with TST positivity.

Zhao et al, AmJClinPathol 2009, USA[34]	Mar-May 2008	40	-	QFT-G-IT	-	20/40 Historical TST	10/40 (25%)	0	75% $\kappa=0.5$	+TST/-QFT	-
Girardi et al, Euro Surveill, 2009, Italy[35]	2004-2005	115	5 IU	T-SPOT.TB & QFT-G-IT	37.4%	61/115 (53%) [$\geq 10\text{mm}$]	TSPOT 42/115 (26.5%) QFT 29/115 (25.3%)	-	TSPOT 67% $\kappa=0.34$ QFT 65% $\kappa=0.32$	+TST/-IGRA	-TSPOT positivity was significantly associated with working in a high risk ward, all 3 tests were associated with increased age, only TST positivity was associated with BCG vaccination.
Cummings et al, ICHE 2009, USA[36]	June 2007-Feb 2008	182	-	QFT-G-IT	7%	4/182 (2.2%) [$\geq 10\text{mm}$]	3/182 (1.64%)	10/182 (5.5%)	96% But 0% on positive results	+TST/-QFT	-

* two studies based on same cohort, ERJ data displayed in table

TB: Tuberculosis, HCW: health care worker, TU: Tuberculin Unit, IU: International Unit, TST: Tuberculin Skin Test, QFT : QuantiFERON test,, QFT-G: QFT Gold test, QFT-G-IT: QFT Gold In-Tube test IGRA: Interferon gamma release assay, IFN- γ : Interferon-gamma, BCG: Bacille Calmette-Guerin vaccine, κ : kappa, pos: positivity

Table A2. Methodology and results of longitudinal, serial testing IGRA studies in HCWs, stratified by TB incidence
[N = 10 studies]

Study, Journal, Year & Country	Date of Subject Recruitment	N	TST (PPD dose)	IGRA	Repeating testing data							
					Timing between repeat testing	Baseline TST positivity rate n/N (%) [cut-point]	Baseline IGRA positivity rate n/N (%)	TST conversion rate during f-up [cut-off for conversion]	IGRA conversion rate during f-up [cut-off for conversion]	Concordance between TST and IGRA conversions (% agree and k)	IGRA reversion rates	Correlation between risk factors (exposure) and conversions
High incidence countries												
Pai et al, AJRCCM 2006, India[37]	Jan-May 2004	216	1 TU	QFT-G-IT	18 months	48/216 (22%) [≥10mm]	38/216 (18%)	6mm: 14/147 10mm: 6/147 (4.1%) [TST ≥10 mm, with an increase of 6 mm]	17/147 (11.56%) [0.35 IU/ml] 11/147 [0.70 IU/ml]	96% k=0.7	2/28 (7.1%) baseline concordant positive 7/10 (70%) among baseline discordant	-
Joshi et al., 2 nd Global Symposium on IGRAs (conference presentation), India, 2009[38]	2008	79	-	QFT-G-IT	0, 6 and 12 months	-	22/79 27.8%	11/57 (19%) converted at 6 months 8/46 (17%) convert at 12 months.	10 of these initial converters reverted back to a negative test result at 12 months (“unstable conversion”). Of the 22 students QFT positive at baseline, 6 (27%) had reverted to a negative result at 6 months. Three of them converted back to a positive test result at 12 months (“unstable reversion”)			-
Intermediate & Low Incidence Countries												

Pollock et al, ICHE 2009, USA[39]	2006	143 (43)* ***	5 TU	QFT-G-IT & T-SPOT.TB	1 -7 months	143/143 100% [≥10mm]	26/143 (18%) 0/43**** (0%) 2/143 (Indeterminate)	-	2/43*** (4.6%) 5/36 tested positive by TSPOT.TB (TSPOT not done at baseline) **** HCWs at increased risk with positive TST and negative QFT	-	-	-HCWs at increased risk of exposure had an increased positivity rate by QFT (28%). - More than 5 yrs living in a highly TB endemic area was significantly associated with QFT positivity.
Zwerling et al. AJRCCM 2009 (conference presentation), Canada[40]	May 2007-Oct 2008	117	5 TU	QFT-G-IT	1 year	13/117 (11%) [≥10mm]	9/117 (7.7%)	0/57 baseline TST negative	4/56 (7.14%) baseline QFT negatives	-	2/4 (50%) QFT positive reverted	-
Yoshiyama et al, Epi&Inf 2009, Japan[41]	2003, 2005, & 2007	311	-	QFT-G	2 years & 4 years	-	-	-	5/277 (1.8%) converted	-	13/32 (41%) reverted	-Association between QFT conversion and working in a TB ward, all reversions had initial values close to the cut-off.
Chee et al, ICHE 2009, Singapore[42]	2005-2007	207	2 TU	T-SPOT.TB	1 year	177/205 (86.3%)	9/205 (4.3%)	0/18 baseline TST and TSPOT.TB negatives converted	9/182 (4.9%) baseline TSPOT.TB negative converted (all were baseline TST positive)	-	-	-
Lee et al, ICHE 2009, Korea[43]	May-Sept 2007	196	2 TU	QFT-G	1 year	101/196 (51.5%)	28/196 (14.3%)	16/75 (21.3%)	21/146 (14.4%)	84% κ=0.417	-	-Neither hospital department nor exposure to TB

						≥10mm]				(among 75 who were tested at 1 year with both tests)		patients was associated with conversion status
Belknap et al. AJRCCM 2010 (abstract), USA[44]	-	1281	5 TU	QFT-G-IT & T-SPOT.TB	6 months	43/1828 (2.4%)	QFT 70/1828 (3.8%) TSPOT.TB 101/1828 (5.5%)	4/1202 (0.3%) ***9/19 (47.3% TST reversion)	QFT 44/1169 (3.8%) TSPOT 44/1117 (3.9%)	-	QFT 20/50 (40%) TSPOT 36/68 (52.9%)	-Older age was associated with T-SPOT conversion and male gender was associated with QFT-GIT conversion
Costa et al. IAOEH, 2010 Portugal [45]	Feb. 2007-Sept 2009	670	2 TU	QFT-G-IT	1-2.5 years	580/670 (86%) ≥10mm]	208/670 (31%)	≥6mm increase: 98/199 (49.2%) ≥10mm increase: 61/199 (30.7%)	51/462 (11%)	-	IGRA: 46/208 (22.1%) TST: 4/188 (2.1%)	-
Ringshausen et al. BMC ID, 2010 Germany	Dec. 2005-Jan. 2008	182	2 TU	QFT-G-IT	18 weeks	98/178 (55%) Includes prior positives ≥10mm]	18/178 (10.1%)	-	3/160 (1.7%)	-	6/18 (33%)	Persistent positive QFT was associated with advanced age and a positive TST.

TB: Tuberculosis, HCW: health care worker, TU: Tuberculin Unit, IU: International Unit, TST: Tuberculin Skin Test, QFT : QuantiFERON test, IGRA: Interferon gamma release assay, IFN-γ: Interferon-gamma, BCG: Bacille Calmette-Guerin vaccine, κ : kappa, pos: positivity

RESULTS

Secondary Outcomes

Limited data exist on cost effectiveness of IGRAs when used for HCW screening. While over 10 studies have been published on cost effectiveness of IGRAs, only three have focused on screening of HCWs. de Perio et al. reported that both IGRA tests were less costly and more effective than the TST for HCW screening.[46] They found there was no prevalence of LTBI for which TST became the more cost effective option, among BCG vaccinated and unvaccinated HCWs. However, this analysis was restricted to use of IGRAs for screening at entry; it did not consider serial testing. Recently Fox et al, published a cost comparison analysis for IGRA versus the TST to screen HCWs.[31]. Authors found costs could be minimized by using a QFT to confirm a positive TST as compared to QFT only or TST only strategies.[31] Lastly a group from Spain compared costs of TST and QFT in HCWs and found the costs per test quite comparable, cost structures between the two tests differed significantly.[47] None of these cost analyses account for serial testing of HCWs with IGRAs.

Sahni et al. conducted a study to evaluate potential improved uptake of LTBI therapy after implementation of the new QFT test to replace the TST in their health care facility located in the US.[48] The authors reported a 3.3 times increased odds (95%CI: 1.3-8) of accepting LTBI therapy and OR=8.8 (95%CI:3.1-23) for taking therapy post implementation (n=45, pre-implementation, n=62 post-implementation).

A few studies reported operational and programmatic issues in using IGRAs. A report by Miranda et al. from the US investigated an unusually high rate of indeterminate results in HCWs by QFT (11%).[49] Authors found indeterminate rates could be reduced with successful interventions (ie: manual vortexing before incubation and the use of a modified in-tube method). Budnick et al. also reported an unexpectedly high rate of indeterminate results, (20%) in HCWs in the US, but found the rate could be reduced by further standardizing sample processing procedures.[50]

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