Kinetics of a Tuberculosis-Specific Gamma Interferon Release Assay in Military Personnel with a Positive Tuberculin Skin Test

Sigrid E. van Brummelen, Anja M. Bauwens, Noël J. Schlösser, and Sandra M. Arend

Central Military Hospital, Department of Pulmonology, Utrecht, Netherlands, and Leiden University Medical Center, Leiden, Netherlands

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Each year, about 3,000 Dutch army personnel are deployed to regions where tuberculosis (TB) is highly endemic. Screening of military personnel for latent Mycobacterium tuberculosis infection (LTBI) has thus far been based on the tuberculin skin test (TST). The Netherlands is a country with a low prevalence of TB, with a yearly incidence of 5.9 cases/100,000 population in 2007, only one-third of which occurred among native Dutch persons (Tuberculosis in The Netherlands 2007 [www.kncvtbcenter.nl]). Personnel are screened by the TST upon initial recruitment into the army, after deployment, or in the presence of other risk factors for TB exposure. Military personnel with TST conversion are prescribed isoniazid for 6 months to prevent TB disease. The risk of progression from untreated LTBI to active TB is generally believed to be about 10%, with half of the cases occurring within 2 years after infection. However, the risks observed in different studies comparing subjects treated with isoniazid or placebo varied widely, depending on the setting and the characteristics of the study population (38). A major disadvantage of the current policy is that a substantial fraction of initially low-positive TST results and to evaluate the kinetics of the Quantiferon TB Gold In-Tube assay (QFT-Git) in military personnel with a positive TST result. QFT-Git was performed at the time of inclusion in the study and was repeated after 2, 6, 12, and 18 or 24 months. Of 192 participants, 17 were recruits and 175 were screened after deployment (n = 169) or because of travel or health care work. Baseline positive QFT-Git results were observed in 7/17 (41.2%) and 12/174 (6.9%) participants, respectively. During follow-up, a negative QFT-Git result remained negative in 163/165 (98.8%) participants. Of 18 subjects with an initial positive QFT-Git result, reversion to a negative result occurred in 1/6 (16%) recruits, whereas it occurred in 8/12 (66%) subjects after deployment or with other risk factors (P = 0.046). The quantitative result was significantly lower in subjects with reversion than in those with consistent positive results (P = 0.017). This study confirmed a low rate of positive QFT-Git results among military personnel with a positive TST result after deployment, supporting the hypothesis of exposure to NTM. Reversion of the majority of initially low-positive QFT-Git results indicates that QFT-Git may be useful for the diagnosis of later reinfections.

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where the military personnel were deployed, in order to justify a change in treatment policy, the previously observed low rate of positive IGRA results in association with a positive TST result needed to be studied in the current setting. In the previous study (13), QFT-Git was performed only once, and the subjects were not assessed for eventual later conversion or reversion. Previous studies of the kinetics of IGRA gave variable and partly conflicting results (9, 12, 14, 15, 17, 20, 29, 30, 35–37, 40), although the main trend was for high-positive results to usually remain positive, and reversion can occur when the results are low or moderately positive (12, 15, 20, 29, 30, 33, 35–37, 42). In a setting of LTBI, the relevance of follow-up testing by IGRA may lie in the possibility of detecting later reactivation if reversion to a negative result has been documented.

The aims of this study were to study the kinetics of QFT-Git during at least 6 months of follow-up in order to evaluate the possibility of detection of later reactivation and to confirm the previously observed very low rate of positive QFT-Git results in military personnel with a positive TST result after deployment.

MATERIALS AND METHODS

Study design. This prospective observational study included BCG-naive military personnel with documented TST conversion during screening. TST screening is performed for all new recruits in order to obtain a baseline value upon entry into the army as well as in all previously TST-negative military personnel 6 to 8 weeks after they return from deployment to a region where TB is endemic. Finally, TST screening is performed on a yearly basis for nondeployed military personnel with risk factors, such as medical work or travel. Exclusion criteria were suspected or proven immune deficiency, BCG vaccination in the past, or exposure to active TB disease. Isoniazid treatment was offered according to protocol of the Dutch Defense Department. The QFT-Git results were not used for clinical decision making. Subjects who declined isoniazid treatment were monitored by the use of chest radiography every 6 months for 2 years. QFT-Git was performed at 0, 2, 6, and 12 months (the last time point of testing for subjects treated with isoniazid) and 18 or 24 months (when isoniazid was not used or was discontinued), whenever possible, considering the duties and deployment of the subjects. The subjects answered the questions on a written questionnaire regarding past employment in the army (median of 7 years and range of 1 to 40 years for those screened after deployment and median of 7.2 years and range of 3 to 12 years for those screened for other reasons), reported TB contact, TST results, and the proportion of military personnel with documented TST conversion during screening. For deployed personnel, the interval between their return to the Netherlands and skin testing was always 6–8 weeks after they return from deployment to a region where TB is endemic. The characteristics of the subjects are listed in Table 1.

TST results. The average TST result was higher among the recruits (Table 1), with 75% having an induration of ≥15 mm, whereas 25% of military personnel in the after deployment/other risk factors group had a positive TST result comprising an induration of ≥15 mm, which could be explained by the preferential referral of recruits with TST inductions of ≥15 mm for isoniazid treatment. TST results were not affected by age, sex, birth outside the Netherlands, travel to tropical countries, or reported contact with a TB patient. QFT-Git results. Samples for testing by QFT-Git were collected between February 2007 and July 2009. QFT-Git result at inclusion. Valid QFT-Git results at inclusion were obtained for 191/192 participants, of which a positive
Almost all (163/165) initially negative QFT-Git results remained consistently negative during follow-up, with only two conversions being observed: one in a recruit who had a negative result at 0, 2, and 6 months, followed by a result of 0.39 IU/ml at 12 months, and another in a subject after deployment with IFN-γ values of 0.32 and 0.28 IU/ml, values just below the cutoff, at 0 and 2 months, respectively, followed by values of 0.59 and 0.56 IU/ml at 6 and 12 months, respectively.

Of 18 subjects with an initial positive QFT-Git result, 7 were consistently positive during follow-up, 2 had one negative value between low-positive values, while 9 reverted to negative; 7 of the last 9 subjects had reverted at the second available time point of 2 months (n = 6) or 6 months (n = 1). Reversion occurred in 1/6 (16%) recruits with an initial positive test result, whereas reversion occurred in 8/12 (66%) subjects in the after deployment/other risk factors group (P = 0.046). Figure 2 shows that for subjects with an initial positive QFT-Git result, the quantitative test result was significantly lower for subjects with reversion than for those who demonstrated a consistent positive pattern (0.72 ± 0.33 and 3.76 ± 3.3 IU/ml IFN-γ, respectively, P = 0.017).

Among 18 subjects with a baseline positive QFT-Git result, consistently positive results were more frequent in subjects with a larger TST induration size, which was not a statistically significant difference, possibly due to the overall low number of positive test results. There was a trend toward higher quantitative test results in subjects with a TST induration of ≥15 mm than in those with an induration of <15 mm (medians, 2.22 and 0.91 IU/ml IFN-γ, respectively; P = 0.052).

The participants who had been deployed were rather homogeneous, with the main destination of deployment being Afghanistan for 163/169 (96.4%) participants and with the durations of deployment being <4 months for 47/168 (27.8%) participants and 4 to 6 months for 108/168 (63.9%) participants. There was no effect between the size of the TST induration, the duration or destination of the deployment, the level of contact with the local population, performance of a medical task, or visiting a local hospital during deployment and the QFT-Git result (data not shown).
FIG. 2. The mean QFT-Git result for subjects with consistently positive results or with one negative value between low-positive values \((n = 9)\) was \(3.76 \pm 3.3\) IU/ml IFN-\(\gamma\) (A), which was significantly higher than the mean of \(0.72 \pm 0.33\) IU/ml for nine subjects with reversion to negativity (B) \((P = 0.017)\). The corresponding median IFN-\(\gamma\) values were 2.3 and 0.66 IU/ml, respectively. Dotted line, cutoff value for a positive response (0.35 IU/ml). Note that the \(y\) axes in the two panels have different scales.

differences of which were not statistically significant, possibly due to the low overall number of positive results.

**Preventive treatment.** All TST-positive participants were offered isoniazid preventive treatment, and two of these individuals declined treatment. Liver function elevations of <3 times, 3 to 5 times, 5 to 10 times, and ≥10 times the upper limit of normal occurred in 40 (21%), 7 (4%), 4 (2%), and 2 (1%) subjects, respectively. Treatment was discontinued in 11 participants after a median of 2 months (range, 2 weeks to 4 month) because of abnormal liver function test results, according to guidelines (values greater than or equal to five times the upper limit of normal or greater than or equal to three times the upper limit of normal in association with complaints) or subjective complaints without objective signs. For the subjects who did or who did not complete 6 months of isoniazid treatment, there was no significant difference in the proportion with a positive QFT-Git result at the inclusion (18/178 [10.1%] and 1/13 [7.7%], respectively) or at 6 months (9/155 [5.8%] and 0/11 [0%], respectively). The number of subjects who did not complete 6 months of preventive treatment was, however, too small for a robust statistical analysis. No subject had or developed active TB during the follow-up period.

**DISCUSSION**

The results of this study showed a 6-fold lower rate of positive QFT-Git results among military personnel with TST conversion in the after deployment/other risk factors group (6.9%) than in recruits (41.2%), confirming and extending the findings of a previous study by Franken et al., who observed a positive QFT-Git result in 11.5% and 44.4% of the individuals in these groups, respectively (13). The very low rate of positive QFT-Git results after deployment would be consistent with the idea that most positive TST results in this group were false positive due to exposure to NTM, while the higher rate of positive QFT-Git responses in recruits reflected previous infection with *M. tuberculosis*.

Surprisingly, the QFT-Git results were negative for 42/43 subjects with a TST induration of ≥15 mm after deployment. In these BCG-naïve participants, this represented documented conversion, since all had had a negative TST result upon entry into the army. If the conversion observed would have been caused by actual infection with *M. tuberculosis* during deployment, most should have had a positive QFT-Git result, since this assay is highly sensitive for the detection of recent infection. However, most QFT-Git results were negative. False-positive TST results due to BCG mostly do not have indurations that exceed 15 mm (39). However, there are no solid data on the TST induration sizes due to NTM when the test is performed shortly after exposure, as would be the case in our participants. Therefore, our data raise the hypothesis that exposure to NTM can cause large TST induration responses immediately after a period of exposure.

Pseudopseudemics of TST conversion among deployed U.S. military personnel have been analyzed, and exposure to NTM was considered one of the possible explanations (25), but no tests were done to support that idea. More specific diagnostic tests such as IGRAstr may be therefore of potential value for the screening of army personnel. Only a few studies that used TB-specific IGRAstr to test military personnel have been published, and apart from the present study and the previously published study by Franken et al. (13), none of these were done in a setting of TST screening after deployment. Two prior studies included military recruits. The study by Mazurek et al. compared the first- and second-generation Quantiferon assays, only the latter of which is based on *M. tuberculosis*-specific antigens, in with 856 U.S. navy recruits, 5.1% of whom had TST indurations of ≥10 mm, while only 0.6% had a positive QFT-Git result (27). The other study used a TB-specific in-house enzyme-linked immunospot (ELISPOT) assay to test 100 mostly BCG-vaccinated Chinese military recruits, and positive TST and positive ELISPOT assay results were obtained for 41% and 21% of the subjects, respectively, which reflects the higher prevalence of true LTBI in China and shows that IGRAstr may contribute to the more accurate diagnosis of LBIs in a setting with a high level of endemicity of TB and routine BCG vaccination (41).

Two other studies of military personnel were done as part of contact investigations. Among mostly BCG-vaccinated Swiss military personnel who were contacts of a colleague with cavitating pulmonary TB, QFT-Git was used as the primary screening tool and resulted in 34/168 (20.2%) positive results and a good correlation between the QFT-Git result and the level of exposure (19). In a similar study among South Korean military camp contacts of soldiers with active pulmonary TB, the QFT-Git result was positive for 25/175 (14.3%) of the...
individuals (10). These studies show that the risk of LTBI can be substantial in a military setting with actual exposure to known smear-positive cases, albeit it is still lower than the 30 to 40% positive TST results found among close contacts in a civilian setting. This confirms that IGRA results can be valuable when the TST result is unreliable due to BCG vaccination.

The only published studies that used IGRA to test military personnel following deployment to a region with a higher prevalence of TB are the present study and the study by Franken et al. (13) mentioned above. Both were limited to non-BCG-vaccinated Dutch military personnel, thereby excluding BCG as a cause of false-positive TST results. The main difference between the studies was the destination of deployment. As IGRA is highly sensitive for the detection of a recent LTBI, it is reasonable to assume that only the observed 6.9% positive QFT-Git results are for most of those with true LTBI. The low rate of positive QFT-Git results indicates that most positive TST results in this setting were probably not caused by recent LTBI and could be related to exposure to NTM; yet all individuals with TST conversion were offered and mostly completed preventive treatment with isoniazid. Significant elevations in liver function test levels were frequent, and side effects were the cause of the discontinuation of treatment for 5.8% of the subjects. On the basis of the results of this and the previous study (13), presumably mostly unjustified treatment of most individuals, together with the considerable costs, risks, and disadvantages of treatment, we believe that it is justified to reconsider the cost-benefit ratio of preventive treatment in military personnel.

Although the negative and positive predictive values of IGRA remain to be proven definitively, the findings presented in several recent publications show that progression to active TB occurred only in subjects with a positive IGRA result at the time of screening, while TB did not occur in those with negative TST results (1, 11, 16). A study for which the association was less clear was conducted with exposed and mostly treated children in Turkey, where reinfections may be more common (6). Several national guidelines now advocate the results of IGRA, either as a replacement of or as adjunct to the TST, for clinical decision making (26, 28). In the Netherlands, a preliminary guideline was issued in 2008. In brief, that guideline advocates the use of a two-step approach, in which a positive TST result can be followed by confirmation by an IGRA for clinical decision making (18).

Because the results of the present study showed even lower rates of positive QFT-Git results than the previous study, the results prompted a change in treatment policy at the Central Military Hospital in November 2008, with preventive treatment being offered exclusively to military personnel with documented TST conversion as well as a positive QFT-Git result, while those with a negative QFT-Git result are monitored for 2 years by the use of a visit for clinical examination and chest radiography every 6 months. The benefit of restricting preventive treatment to less than 10% of subjects with TST conversions represents considerable benefit in terms of avoided costs, inconvenience, and side effects. If the negative predictive value proves to be sufficiently high, the monitoring of subjects with negative QFT-Git results may not be necessary, resulting in additional logistical and financial savings. This policy could most likely be extrapolated to BCG-vaccinated subjects, since the risk of a false-positive TST result is even higher in that group.

Thus far, the role of IGRA during follow-up is not clear, and it has been suggested that repeated IGRA are not useful at all (12, 17, 36). Reports on the kinetics of IGRA are limited in number, and the results were, in part, contradictory, which may be related to differences in the study populations, the assays used, and the endemicity of TB and the risk of reinfection. A list of studies that reported the results of repeat testing by IGRA in a setting of active or latent TB infection is provided as supplemental material to this article. In patients treated for active TB, the quantitative results of follow-up IGRA decreased as a rule, but the rates of reversion to a negative result varied from 10% to 71% (20, 24, 35–37). Follow-up IGRA for subjects with latent M. tuberculosis infection showed more variable results, but several studies reported an inverse association between the baseline IGRA response and the chance of reversion, which is in accordance with the results of the present study (9, 12, 15, 17, 24, 29, 30, 40). In our study, an initial negative QFT-Git result remained negative for 163/165 subjects. Half of the individuals with initially positive results reverted to negative, which was limited to baseline positive values comprising a moderate increase (<1.2 IU/ml IFN-γ in the present study) and which mainly occurred in subjects in the after deployment/other risk factors group. This pattern has been observed previously (12, 29–31), indicating that the result of a follow-up IGRA is unlikely to revert to negative when a positive result exceeds a certain value, even though a quantitative decrease even of high values was frequently observed during follow-up, but that subjects with low-positive results have a considerable chance of reverting to having a negative result if the exposure was recent. The participants who were screened after deployment and who had a positive QFT-Git result were probably recently infected, while for recruits with positive QFT-Git results, the interval between infection and testing was unknown but was presumably longer than that for individuals screened after deployment. The available data indicate that a positive result remains positive for longer periods for roughly 40% of subjects with LTBI (5, 23, 24), which is in agreement with the findings of our study for recruits.

The reason why QFT-Git results revert to negative over time in over half of the subjects may be caused by the natural kinetics of the immune response, with a change from large numbers of circulating effector cells to low numbers of memory cells that cannot respond with IFN-γ production during the 24 h of incubation used for the commercial IGRA (23). In addition, there may be an effect of preventive treatment, but this has thus far not been demonstrated unequivocally. The present study included a limited number of positive results, and almost all individuals were treated, which did not allow a robust analysis of the effect of treatment. In the past, a documented positive TST result in an individual who thereafter needed to be screened repeatedly for LTBI implicated that future screening was possible only by chest radiography, which is sensitive for the detection of only active pulmonary TB and not LTBI. A documented negative QFT-Git result at the time of screening or reversion of a positive QFT-Git result to negative after 6 months strongly suggests that QFT-Git might be used to detect a later reactivation, although some variation around the threshold value was observed, and it cannot be
concluded definitively that a negative result will remain negative thereafter. However, follow-up testing is probably not useful if the initial QFT-Git result exceeds a certain threshold value.

In conclusion, this study confirms that the majority of TST conversions after deployment are not caused by actual infection with _M. tuberculosis_ but could be related to exposure to NTM. This supports a change in policy in which only subjects with TST conversion in association with a positive IGRA result should be treated, on the condition that they have adequate follow-up. This change in policy is under investigation. In the presence of a positive IGRA result, a follow-up IGRA was not useful when the initial test result was in the higher range, while initially low or moderately positive test results frequently reverted to negative with potential use of eventual later screening and detection of reinfection.

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REFERENCES


