Population tailored modification of tuberculosis specific interferon-gamma release assay

Kata Horvati a, Szilvia Bősze a, Hannah P. Gideon b, Bernadett Bacsa a, Tamás G. Szabó c,d, Rene Goliath b, Molebogeng X. Rangaka b, Ferenc Hudecz a,e, Robert J. Wilkinson b,f,g, Katalin A. Wilkinson b,f,*

a MTA-ELTE Research Group of Peptide Chemistry, Eötvös L. University, Budapest, Hungary
b Clinical Infectious Disease Research Initiative, Institute of Infectious Disease and Molecular Medicine, University of Cape Town, Cape Town, South Africa
c Department of Genetics, Cell- and Immunobiology, Semmelweis University, Budapest, Hungary
d Department of Laboratory Medicine, Semmelweis University, Budapest, Hungary
e Department of Organic Chemistry, Eötvös L. University, Budapest, Hungary
f The Francis Crick Institute Mill Hill Laboratory, London NW7 1AA, UK
g Department of Medicine, Imperial College London W2 1PG, UK

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Summary
Objectives: Blood-based Interferon-Gamma Release Assays (IGRA) identify Mycobacterium tuberculosis (MTB) sensitisation with increased specificity, but sensitivity remains impaired in human immunodeficiency virus (HIV) infected persons. The QuantIFERON-TB Gold In-Tube test contains peptide 38–55 of Rv2654c, based on data indicating differential recognition between tuberculosis patients and BCG vaccinated controls in Europe. We aimed to fine map the T cell response to Rv2654c with the view of improving sensitivity.

Methods: Interferon-gamma ELISpot assay was used in HIV uninfected persons with latent and active tuberculosis to map peptide epitopes of Rv2654c. A modified IGRA was tested in two further groups of 55 HIV uninfected and 44 HIV infected persons, recruited in South Africa.

Results: The most prominently recognised peptide was between amino acids 51–65. Using p51-65 to boost the QuantiFERON-TB Gold In-Tube assay, the quantitative performance of the modified IGRA increased from 1.83 IU/ml (IQR 0.30–7.35) to 2.83 (IQR 0.28–12.2; p = 0.002) in the HIV uninfected group. In the HIV infected cohort the percentage of positive responders increased from 57% to 64% but only after 3 months of ART (p = ns).

* Corresponding author. Room N2.09.B3, Wernher Beit North Building, Institute of Infectious Disease and Molecular Medicine, Faculty of Health Sciences, Observatory 7925, South Africa. Tel.: +27 21 650 6906; fax: +27 (021) 406 6796.
E-mail address: katalin.wilkinson@uct.ac.za (K.A. Wilkinson).

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Introduction

Despite the worldwide availability of vaccination, diagnosis and treatment, tuberculosis (TB) is still a major global health problem and it is estimated that more than one-third of the world’s population is infected by *Mycobacterium tuberculosis* (MTB). Exposure to MTB may result in latent TB infection (LTBI) which is defined by the absence of clinical TB symptoms but carries about 5–10% lifetime risk of developing active TB and comprises a significant reservoir of future cases of active disease, particularly in countries with high HIV burdens. The estimated incidence of TB is 9 million each year, among which 1.5 million cases die. HIV infected patients are about 30 times more likely to develop active TB and HIV-TB co-infection is responsible for one fifth of all TB related deaths.

The currently used vaccine — *Mycobacterium bovis* Bacille Calmette Guérin (BCG) — does not protect adults against TB. Moreover, BCG vaccination renders the TB diagnosis more difficult as the commonly used tuberculin skin test (TST), based on the administration of purified protein derivative (PPD), may give false-positive results due to previous BCG immunization.

Proteins encoded at loci missing from BCG have become of interest for TB diagnosis. Such regions of difference (RD), by definition, are absent from all BCG vaccine strains. Comparative genomic studies have identified 11 regions of difference of which RD1 contains important immunodominant proteins such as ESAT-6 (Rv3875) and CFP-10 (Rv3874). These proteins showed reliable diagnostic potential in T-cell based IFN-γ release assays (IGRAs) and are used as antigens in FDA approved and commercially available blood tests such as QuantiFERON-TB Gold In-Tube (QFT) and T-SPOT.TB. ESAT-6 and CFP-10 are early secreted proteins which form a heterodimeric complex and contribute to the virulence of MTB. However, genes for ESAT-6 and CFP-10 were found not only in MTB but also in virulent *M. bovis*, and in several non-tuberculous mycobacteria. Thus, infection with such ESAT-6 or CFP-10 expressing mycobacterial species can result in a false-positive response in blood tests based on ESAT-6 and CFP-10 antigens.

MTB sensitization was traditionally described with a model of binary distribution between active TB and latent TB. Evidence supports that the interaction between MTB and the host immune system represents a spectrum of immune responses, bacterial load, metabolic activity and stages of infection ranging from sterilizing immunity to TB disease. Such heterogeneity may explain the fact that antigens that are targets of the immune response in latent TB are frequently found as target in active TB and no adequate differentiation between latent and active disease has been revealed.

Insufficient sensitivity of IGRA tests could be further enhanced with the use of more antigens derived from different immunodominant proteins of MTB. Previous studies have demonstrated that Rv2654c can improve specific immune-based diagnosis of TB infection especially in the BCG-vaccinated population and one peptide (p3855) has been included in the commercially available QuantiFERON-TB Gold In-Tube (QFT) test. Rv2654c is an 81 amino acid alanine-rich protein, encoded in the RD11 region, highly specific for MTB and absent from most of the atypical mycobacteria. We performed a systematic epitope mapping of the Rv2654c protein and found that the most immunogenic peptide recognized by MTB sensitized individuals in South Africa is different from that previously described by Aagaard and co-workers. We describe the evaluation of a modified QuantiFERON-TB Gold In-Tube test in HIV infected and uninfected cohorts.

Materials and methods

MHC-binding predictions

The amino acid sequence of Rv2654c protein was downloaded from the TubercuList website (http://tuberculist.epfl.ch/index.html). In silico identification of immunogenic regions of the Rv2654c protein was based on predicting the MHC-binding affinity of 15-mer peptides overlapping by 14 amino acid residues, using the MHC II binding prediction tool and evaluated based on the consensus percentile ranks. As suggested by Paul and colleagues, 15-mer peptides were sorted based on the median value of the percentile ranks predicted for each representative allele and the top scoring ~20% were identified as possible immunodominant epitopes.

Synthesis and analytical characterization of peptides

Peptides were produced on solid phase with an automated peptide synthesizer (Syro-I, Biotage, Uppsala, Sweden) using standard Fmoc/tBu strategy. Peptides used in this study contained an amide group at the C-terminus for stability purposes. After cleavage, crude peptides were purified by semi-preparative HPLC and analysed by mass spectrometry, analytical HPLC and amino acid analysis (Table 1). Mass spectrometric analyses were performed on a Bruker Esquire 3000 + ion trap mass spectrometer (Bruker, Bremen, Germany) equipped with electrospray ionization (ESI) source. Samples were dissolved in a mixture of acetonitrile/water = 1/1 (v/v) containing 0.1% acetic acid and introduced by a syringe pump with a flow rate of 10 μL/min. The homogeneity of the compounds was checked by...
analytical HPLC using a laboratory-assembled Knauer HPLC system (Bad Homburg, Germany) using 1 mL/min flow rate at room temperature. The gradient elution system consisted of 0.1% TFA in water (eluent A) and 0.1% TFA in acetonitrile/water = 80/20 (v/v) (eluent B). Amino acid analyses were performed on a Sykam Amino Acid 5433H analyzer (Eresing, Germany) equipped with an ion exchange separation column and post-column derivatisation. Prior to analysis samples were hydrolysed with 6 M HCl in sealed and evacuated tubes and post-column derivatisation. Prior to analysis samples were hydrolysed with 6 M HCl in sealed and evacuated tubes at 110 °C for 24 h. For post-column derivatisation the ninhydrin-method was used. Peptides were stored dry until reconstitution as 1 mg/mL stock solutions in 0.5% DMSO containing PBS buffer.

### Selection and description of participants

The University of Cape Town’s Faculty of Health Sciences Human Research Ethics Committee approved this study (REC 296/2007, REC 245/2009). Written informed consent was obtained from all participants.

For epitope mapping of Rv2654c, patients with active TB or latent TB infection (LTBI) were recruited at Ubuntu Clinic, site B Khayelitsha, South Africa. All were of Xhosa ethnicity. Active TB was defined by smear and/or culture positivity for MTB from one or more sputum specimens. LTBI was defined by an IFN-γ ELISpot response to ESAT-6 or CFP-10 of 20 spot-forming cells (SFC)/10⁶ PBMC above background in the absence of clinical symptoms or radiographic abnormality and with negative sputum smear and culture for M. tuberculosis. All subjects recruited for epitope mapping underwent HIV counselling and testing, and positivity was an exclusion criterion, together with pregnancy and age under 18 years.

For the evaluation of the boosting effect of peptide 51-65 on the QuantiFERON-TB Gold In-Tube test, 55 HIV uninfected and 50 HIV infected persons were recruited at Ubuntu Clinic, Site B Khayelitsha, South Africa. HIV infected patients were recruited at enrolment into the antiretroviral treatment (ART) programme based on their eligibility (CD4⁺ T cell count of 250 cells/mL or less), according to South African national guidelines at the time. 30 mL venous blood was collected for immunological analysis, as well as viral load and CD4⁺ T cell count measured at day 0, 1 month, 3 months and 6 months of ART. Induced sputum was collected at all time-points; six patients with positive MTB culture were referred to TB treatment and excluded from further follow up and analysis.

### Cell culture and in vitro assays

Peripheral blood mononuclear cells (PBMC) were prepared using standard Ficoll separation technique and were stored in liquid nitrogen until used. The measurement of IFN-γ by ELISpot was performed as described previously using the human Interferon-γ ELISpot⁹ kit (MABTECH, Nacka Strand, Sweden). Briefly, 2.5 × 10⁶ PBMC were plated in 200 μL of RPMI culture media supplemented with 10% FCS. Antigenic stimuli were either Rv2654c peptides at 10 μg/mL final concentration, or a pool of ESAT-6 (Rv3875, 5 μg/mL) and CFP10 (Rv3874, 5 μg/mL) derived peptides. As positive control, anti-CD3 mAb CD3-2 at 100 ng/mL final concentration was used. ELISpot plates were read on an Immunospot Series 3B Analyzer (Cellular Technology, Cleveland, OH, USA) and plates were retained for visual inspection and confirmation in the case of anomaly. Results are quoted as spot forming cells per million PBMC (SFC/10⁶ PBMC), with the background (unstimulated) values subtracted.

The QuantiFERON-TB Gold In-Tube test was performed in a Cellestis accredited laboratory according to the manufacturer’s instructions (QuantiFERON-TB Gold In-Tube, Cellestis Ltd., Carnegie, Australia). The modified (peptide boosted) QuantiFERON-TB Gold In-Tube test (QFTB) was performed in parallel, using a second set of tubes (representing Nil, Antigen, Mitogen), with peptide 51-65 added at 5 μg/mL final concentration (determined based on dose response experiments) in the laboratory. Tubes were

### Table 1  Sequence and characteristics of peptides used in this study.

| Code  | Sequence                  | Mₘₐₓ calc. | Mₘₐₓ meas. | Rₜ [min] |
|-------|---------------------------|------------|------------|---------|
| p1-20 | MSGHALAARTLLAAADELVG      | 1966.3     | 1966.1     | 31.7    |
| p11-30| LLAAADELVGGPPVEASAA       | 1821.1     | 1821.0     | 25.8    |
| p21-40| GPPYEASAALLAGDAAGWR       | 1837.0     | 1837.0     | 27.5    |
| p31-50| LGDAAGAWRTAELRAL          | 1982.3     | 1982.1     | 34.4    |
| p41-60| TAAVELARLYRVAESHG V       | 2019.3     | 2019.2     | 33.3    |
| p51-70| VRVAESHGVAALFAAATAA       | 1910.2     | 1909.8     | 32.0    |
| p61-81| AAVLF A AAAAAAADV D G D P P| 1925.2     | 1925.1     | 30.5    |
| p38-55| AWRTAAELRALVR A A          | 1922.1     | 1922.2     | 41.5    |
| p51-65| VRVAESHGVAALVL            | 1523.8     | 1523.8     | 28.4    |
| p55-70| A V SHGVAALVF A A A A A A A V | 1483.8     | 1483.8     | 30.0    |
| p61-75| AAVLF A AAAAAAADV D G D P P| 1286.7     | 1286.8     | 16.5    |
| p66-81| AA T A A A A AADV D G D P P| 1422.7     | 1422.8     | 16.6    |

**a** Average molecular mass measured by a Bruker Esquire 3000+ electrospray mass spectrometer.

**b** Retention time on an analytical RP-HPLC using an EuroSphere-100, C-18, 5 μm, 250 × 4 mm column; gradient: 5% B, 5 min; 5–60% B, 35 min.

**c** Gradient: 5% B, 5 min; 5–90% B, 45 min.

**d** Phenomenex Jupiter C-4, 5 μm, 250 × 4 mm column, gradient: 5% B, 5 min; 5–70% B, 35 min.
incubated overnight and IFN-γ measured as IU/ml, using the commercial assay. The results were interpreted similarly, using the manufacturer’s criteria for assay positivity (≥0.35 IU/mL above nil).

**In silico HLA-binding study**

In an attempt to relate epitope predictions to our laboratory findings, binding affinities were also corrected for the allele frequencies observed in the Xhosa population. Since the most detailed description of HLA types in the Xhosa population is based on serological testing, while prediction tools are genotype-based, serotype frequencies were interpreted as allele frequencies according to the conversion matrix provided as Supplementary Table 1.

To estimate the portion of the Xhosa population expected to present the given epitope, we used a model, where an individual’s antigen presenting cells (APCs) express both maternal and paternal HLA-DR and HLA-DQ genes. The relative number of individuals expressing a given combination of HLA loci was computed using the Hardy–Weinberg equation. Competition between peptide epitopes was simulated according to two different scenarios, described in the supplementary information in more detail. Based on these simulations, only the best scoring peptides in a particular subpopulation were regarded as ‘MHC-binder’ peptides. The sum of all fractions of the population, where the peptide scored as a ‘binder’, was regarded as the portion of the whole population that is able to present the peptide epitope. Results were visualized with the help of the Matplotlib Python module.

In addition to the Xhosa population, HLA frequencies described in the Danish population (a source of donors for the p38-55 peptide in a previous study) were also included in the analysis. The frequencies described by Lindblom and colleagues were downloaded from the Allele Frequency Net Database (allelfrequencies.net). In this case, a more simplified model was used for simulating the population, omitting HLA-DR3/DR4 and HLA-DQ loci.

**Statistical analysis**

Data were analysed using GraphPad Prism v6 software (San Diego, CA, USA). For analysis of statistical significance (p < 0.05) the Mann–Whitney U test was used for unpaired data and the Wilcoxon matched pairs test was used for paired data. Bonferroni correction was used to correct p values for multiple comparisons where applicable.

**Results**

**Identification of highly immunogenic regions by MHC-binding predictions**

The median value of the consensus percentile rank scores of binding to supertype-defining HLA II alleles was obtained from predictions by the IEDB webservice, for each 15-mer peptide. As reflected by the median score for MHC binding, the highest binding peptides of the Rv2654c protein were found to be in the middle section (starting amino acid positions between 36 and 40) and the C-terminal section with starting positions between 58 and 63 (Fig. 1).

**Epitope mapping of Rv2654c protein**

To validate and refine the *in silico* prediction, synthetic 20-mer peptides, spanning the complete sequence of the Rv2654c protein and overlapping by 10 amino acids, were prepared and analysed *in vitro* (Table 1). A pool of overlapping peptides, covering the sequence of Rv2654c, was used as a substitute for the recombinant protein, as previous studies have demonstrated antigenic equivalence of the T-cell response between the recombinant proteins and the corresponding synthetic peptide mixtures for ESAT-6, CFP-10 and TB10.4.

Altogether, PBMC from 34 individuals were used, comprising 24 individuals with LTBI (median age 27 years, 14 female, 10 male) and 10 patients with active TB (median age 27 years, 4 female, 6 male). Since all individuals were MTB sensitized, and there was no differential recognition of the peptides between LTBI and active TB patients in line with published literature, results were combined for analyses (Table 2). The frequency of recognition of the C-terminal peptides p51-70 and p61-81 was comparable to that of the Rv2654c pool in the combined groups (29% and 35% compared to 32% respectively). We therefore selected the C-terminal 51–81 region of the protein for further characterization, using shorter peptides.

For the detailed C-terminal analysis, 15-mer peptides were prepared (Table 1), in order to fine map the 51–81 amino acid region. As control, the p38-55 peptide was also prepared as this peptide is included in the QuantiFERON-TB Gold In-Tube test (coded as TB7.7 p4). In the combined group of 19 MTB sensitized individuals (representing a subset of the previously studied n = 10 active TB patients, and n = 9 LTBI, selected based on availability of frozen PBMC remaining after the first set of experiments), the highest IFN-γ response was seen to p51-65, with a median of 27 spot forming cells/million PBMC (SFC/10⁶, IQR 12–85), recognized by 67% of all PBMC samples (Fig. 2). This was significantly higher than the response to the previously described p38-55 peptide (median 8 SFC/10⁶, IQR 0–12; pcorr = 0.002, Mann–Whitney U test with Bonferroni correction), with only 11% of PBMC samples giving a positive response. These results suggest that the most frequently recognized epitope of Rv2654c protein in this population is present within the 51–65 amino acid region.

**Predicted epitope promiscuity in different test populations**

Since the method suggested by Sinu and colleagues does not take differences in HLA frequencies in different populations into account, we related epitope predictions and our *in vitro* experimental findings to the population tested. In order to achieve this, MHC binding scores of the IEDB epitope prediction tool were corrected for the serotype frequencies of the Xhosa population following a stringent model. We found that only the region near the C-terminal end of the protein was immunogenic enough to show up in the analysis (Fig. 3A). If binding only to HLA-
Sensitivity was calculated as a ratio of the number of true positive responders over the number of false negative plus true positive.

Median spot number and interquartile range.

Effected (30 female, 14 male) patients. The median CD4 count was 26 (30 female, 25 male) and 33 for the HIV inactive TB. The median age of the HIV uninfected participants, none of whom had any signs and/or symptoms of active TB. The median CD4 T-cell count in the HIV uninfected cohort was 817 and 197 in the HIV infected group.

In the HIV uninfected group, the quantitative performance of the QFT test increased significantly from a median of 1.83 IU/ml (background — nil, IQR 0.30–7.35) to 2.83 (IQR 0.28–12.2), p = 0.002, Table 3. There was a statistically non-significant change in the percentage of responders from 73% to 75%, and in sensitivity from 95% to 98%, due to the peptide boost.

In the case of the HIV infected cohort, the quantitative performance of the QFT and modified QFTB did not differ at day 0 of ART, with medians of 1.17 (IQR 0.26–7.14) and 1.17 IU/ml (IQR 0.05–6.11) respectively (Table 4). However, 3 months into antiretroviral treatment (ART) the frequency of positive responders increased from 57% to 64% (statistically not significant) with 5 patients changing from LTBI to ESAT-6/CFP-10 response. The cut-off for positive response was 20 SFC/10^6.

Table 2 Recognition of single peptides and Rv2654c peptide pool by individuals with LTBI and active TB.

| Peptide Pool | LTBI n = 24 | Active n = 10 | Combined n = 34 |
|--------------|-------------|--------------|----------------|
|              | No. pos. resp. | Sensitivity % (95% CI) | Median SFC/10^6 (IQR) | No. pos. resp. | Sensitivity % (95% CI) | Median SFC/10^6 (IQR) | No. pos. resp. | Sensitivity % (95% CI) | Median SFC/10^6 (IQR) |
| p1-20        | 6/24         | 25 (9.8–46.7) | 5 (0–19) | 2/10 | 20 (2.5–55.6) | 3 (0–15) | 8/34 | 24 (10.8–41.2) | 4 (0–17) |
| p11-30       | 1/24         | 4 (0.1–21.1)  | 0 (0–8)  | 1/10 | 10 (0.3–44.5) | 2 (0–6)  | 2/34 | 6 (0.7–19.7)  | 0 (0–8)  |
| p21-40       | 4/24         | 17 (4.7–37.4) | 4 (0–10) | 2/10 | 20 (2.5–55.6) | 9 (0–28) | 6/34 | 18 (6.8–34.5) | 4 (0–11) |
| p31-50       | 6/24         | 25 (9.8–46.7) | 6 (0–18) | 2/10 | 20 (2.5–55.6) | 9 (0–19) | 8/34 | 24 (10.8–41.2) | 7 (0–14) |
| p41-60       | 4/24         | 17 (4.7–37.4) | 4 (0–8)  | 1/10 | 10 (0.3–44.5) | 2 (0–9)  | 5/34 | 15 (5.0–31.1) | 4 (0–8)  |
| p51-70       | 7/24         | 29 (12.6–51.1) | 4 (0–23) | 3/10 | 30 (6.7–65.3) | 9 (0–26) | 10/34 | 29 (15.1–47.5) | 4 (0–21) |
| p61-81       | 7/24         | 29 (12.6–51.1) | 8 (0–20) | 5/10 | 50 (18.7–81.3) | 18 (5–46) | 12/34 | 35 (19.8–53.5) | 10 (0–24) |
| Rv2654c pool | 7/24         | 29 (12.6–51.1) | 6 (0–35) | 4/10 | 40 (12.2–73.8) | 10 (0–24) | 11/34 | 32 (17.4–50.5) | 8 (0–24) |

PBMC of either LTBI (n = 24) or active TB patients (n = 10) were in vitro stimulated with 20-mer overlapping peptides spanning the entire sequence of Rv2654c protein using the ELISpot assay. The number of spot forming cells per million PBMC (SFC/10^6) are presented and compared to ESAT-6/CFP-10 response. The cut-off for positive response was 20 SFC/10^6.

a Number of positive responders/number tested.

b Sensitivity was calculated as a ratio of the number of true positive responders over the number of false negative plus true positive responders.

c Median spot number and interquartile range.
of the Rv2654c protein. It is interesting that when testing
p51-65 which was the most dominantly recognised peptide
epitope mapping resulted in the identification of peptide
region between amino acids 51
forming cells per million PBMC (SFC/10^6) are presented from
of Rv2654c compared to p38-55 peptide
peptides of Rv2654c in South Africa are in the
test. Our data indicate that more frequently recognised
can be used to induce a cytokine response in an IGRA-type
evaluated with the aim to identify antigenic peptides that
are generally more stable than full-length proteins or whole organ-
isms, and they do not require refrigerated storage.

Therefore, epitope mapping of the Rv2654c protein was
evaluated with the aim to identify antigenic peptides that
can be used to induce a cytokine response in an IGRA-type
test. Our data indicate that more frequently recognised
peptides of Rv2654c in South Africa are in the C-terminal
region between amino acids 51–81. Further detailed
epitope mapping resulted in the identification of peptide
p51-65 which was the most dominantly recognised peptide
of the Rv2654c protein. It is interesting that when testing
the 20-mer peptides, the spot counts were higher for p61-
81 as compared to p51-70, which contains the p51-65
sequence. However, the % responders overall were high
for both peptides. The strength of peptide binding is influ-
enced by the flanking regions of the epitope and it is
possible that the p51-65 sequence allows stronger binding
of the epitope region as compared to p51-70. It is also likely
that the shared sequence between p61-81 and p51-65,
amino acids 61–65 are part of the epitope core, the binding
of which is more efficient in the context of the p51-65, in
stimulating T cell responses.

This finding contrasts with publications^18,19 that identi-
ified p38-55 as the most potent antigenic peptide. Our
observation might be explained by different experimental
conditions (overnight ELISpot assay compared to a 5 day
dilution assay combined with the measurement of
IFN-γ in the supernatant by ELISA), or by the difference
in the HLA frequencies of the South African Xhosa popula-
tion, where our experiments were performed. Computational
prediction suggested that the middle region
between amino acids 30–50 of the Rv2654c protein to be
unrecognised by the Xhosa population and only the region
near the C-terminal section of the protein was predicted
to be immunogenic enough to show up in the analysis.
These results, together with the results of Sette and his
coworkers,^36 who have also found discrepancy between
their study and earlier works, highlight the necessity of tak-
ing population dependent HLA restriction in antigen recog-
nition into account.

In terms of chemical considerations, p51-65 bears
further advantages compared to p38-55, such as being
more soluble and more stable under in vitro
conditions. These characteristics make peptide p51-65 that provoked
significantly increased IFN-γ response in a whole blood
assay, a more convenient synthetic antigen.

Using synthetic peptide p51-65 in the QuantiFERON-TB
Gold In-Tube assay resulted in significant boosting of the
quantitative performance of the QFT test in the HIV
uninfected group. This increase in the IFN-γ response is
most likely due to the presence of the additional epitope
recognised by antigen specific T cells, secreting IFN-γ. As
the IFN-γ is measured in the supernatant, the response is
additive, hence the increase. In the HIV infected cohort
the quantitative performance did not change at day 0 and 1
month into the initiation of antiretroviral treatment. How-
ever, 3 months of ART resulted in an enhanced proportion of
persons scoring positive in the boosted QFT assay, most
likely due to ART induced immune recovery and increased
CD4^+ T-cell numbers, able to recognise additional epitopes.
Although the increased recognition was not statistically
(and thus also not clinically) significant, is in accordance
with the findings of Kellar and colleagues, who have demon-
strated that the combination of ESAT-6 and CFP-10 peptide
pools with the pool of overlapping peptides representing
Rv2654c protein, has resulted in a significantly greater
cytokine and chemokine response in TB patients using
whole-blood assay. Morever, the reason for not seeing
a significant increase in median IFN-γ responses in the HIV
infected individuals could be due to the relatively short
duration of follow up (6 months) on ART. In our previous
study,^18 we demonstrated that ART is associated with an
absolute increase in effector function (which is what is being

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**Figure 2** Detailed analysis of the C-terminal 51–81 region of Rv2654c compared to p38-55 peptide. The number of spot forming cells per million PBMC (SFC/10^6) are presented from n = 19 MTB sensitized individuals. The cut-off for positive response was 20 SFC/10^6 above nil (dotted line). Significance was calculated using Mann–Whitney U test. **p < 0.005. Black lines represent the median SFC/10^6 values on both panels.**
Figure 3  Population tailored epitope prediction. The portion of the population able to present a 15-mer peptide was estimated based on simulations combining MHC binding scores of the IEDB epitope prediction tool and HLA serotype frequencies described in the Xhosa population. The 5 best scoring peptides for a given combination of HLA alleles were regarded as binders (binding to an APC), with a fair chance for antigen presentation. The number of people likely to have a given combination of HLA alleles was estimated using the Hardy–Weinberg equation. Percentage of the whole population, expected to present a given peptide is given for each possible starting position of 15-mers derived from the Rv2654 protein. Panel (A) represents the percentage of the Xhosa population, expected to present 15-mer peptides, based on both HLA-DR and HLA-DQ predictions of the IEDB tool. On panel (B), percentage of Xhosa population based only on HLA-DR prediction is presented, while on panel (C), percentage of the Danish population, based on HLA-DR predictions is represented.

| Table 3  | Recognition and boosting effect of p51-65 in the HIV negative cohort. |
|----------|---------------------------------------------------------------------|
|          | QFT performance          | QFTB performance          |
| Number tested | n = 55                   | 55                        |
| Median IU/ml (IQR) | 1.83 (0.32–6.90)      | 2.83 (0.32–11.54)        |
| Comparison of median IU/ml response | ** (p = 0.002) | ** (p = 0.002) |
| No. of positive responders | 40                   | 41                        |
| Responders % | 73                   | 75                        |
| Sensitivity % (95% CI) | 95 (83.8–99.4) | 98 (87.1–99.9) |

Blood samples of 55 HIV uninfected participant were assayed in commercially available QFT and p51-65 peptide boosted QFT (QFTB). The cut-off for positive recognition was 0.35 IU/ml above nil.

- **Median IU/ml value in QFT or QFTB assay and interquartile range.**
- **Significance was calculated using Wilcoxon matched pairs test.** **p < 0.01.**
- **Number of positive responders/number tested.**
- **Percentage of responders out of all donors tested.**
- **Sensitivity was calculated as a ratio of the number of true positive responders over the number of false negative plus true positive responders. False negativity was confirmed by ELISpot using ESAT-6/CFP-10 antigens. The difference in sensitivity statistically is not significant.**
measured by IGRA, but the proportional response decreased over 1 year of ART, and the strongest correlate of increased ART-mediated immunity was the central memory response. Thus, in the expanding CD4 T cell pool, containing antigen specific T cells that are potentially able to recognise the additional antigen, these cells are not high enough in numbers to reflect an increase in IFN-γ responses at 6 months of ART.

Overall, different ethnic populations may respond to different peptides than what is available in the commercial IGRA, but the overall effect is a modest increase in responders and not likely to change current assay procedures. While our data are preliminary and warrant further validation with different epitopes, we can conclude that it shows promise for population tailored detection of MTB sensitization and for promiscuous synthetic epitope peptides of proteins such as Rv2654c to be considered as part of more effective immunodiagnostics.

Conflict of interest

The authors declare no commercial or financial conflict of interest.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.jinf.2015.10.012.

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