**A simple breath test for Tuberculosis using ion mobility: A pilot study**

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**Summary**

Tuberculosis (TB) remains one of the world’s major health burdens with 9.6 million new infections globally. Though considerable progress has been made in reduction of TB incidence and mortality, there is a continuous need for lower cost, simpler and more robust means of diagnosis. One method that may fulfil these requirements is in the area of breath analysis. In this study we analysed the breath of 21 patients with pulmonary or extra-pulmonary TB, recruited from a UK teaching hospital (University Hospital Coventry and Warwickshire) before or within 1 week of commencing treatment for TB. TB diagnosis was confirmed by reference tests (mycobacterial culture), histology or radiology. 19 controls were recruited to calculate specificity; these patients were all interferon-gamma release assay negative (T.SPOT®.*TB*, Oxford Immunotec Ltd.). Whole breath samples were collected with subsequent chemical analysis undertaken by Ion Mobility Spectrometry. Our results produced a sensitivity of 81% and a specificity of 79% for all cases of TB (pulmonary and extra-pulmonary). Though lower than other studies analysing pulmonary TB alone, we believe that this technique shows promise, and a higher sensitivity could be achieved by further improving our sample capture methodology.

**Introduction**

Major advances have been made in TB diagnosis, treatment and mortality since the turn of the century, and the Millennium Development Goal to halt and reverse TB incidence by 2015 has partially been met. Despite this, in 2014 there were still an estimated 9.6 million new TB cases, with 1.5 million deaths [1]. Whilst the majority of those affected are from low- and middle-income countries, the significant burden of co-infection with TB and HIV (Human Immunodeficiency Virus), multi-drug resistance, alongside widespread migration, has resulted in TB remaining a global health concern. A new ambition to end the TB epidemic by 2030 has been set as part of the post-2015 global TB strategy [1]. To achieve this Sustainable Development Goal, the vision of the World Health Organisation (WHO) is for everyone to have access to innovative tools and services required for rapid diagnosis, treatment and care. In line with this, breath analysis for TB diagnosis (within minutes) as opposed to standard microbiological culture, which can take up to 2 months, is attractive.

Breath analysis utilises the detection of volatile organic compounds (VOCs), gaseous chemicals that are the endogenous products of metabolism of individual microbes and whole organ systems. It has many advantages for diagnostics – it is rapid, non-invasive, acceptable to patients, and sample volumes are both unlimited and can be given at will [2]. Importantly, analysis of breath is thought to allow access to chemical processes not only in the lung, but also in other organ systems. Detected VOCs may be produced directly in the lung, or reach lung tissue via blood through the internal metabolic pathways of respiration. This allows a wholesale comparison of chemical differences in health and a number of diseased states, including infection and malignancy.

For this reason, many researchers have undertaken chemical analysis of breath by various analytical means, with GCMS (Gas Chromatography Mass Spectrometry) being the most popular. The body of literature on GCMS has identified the key gas-phase biomarkers and supports the hypothesis for breath as a potential tool for TB diagnosis [3,4]. However, GCMS is a laboratory-based technique, with equipment being bulky, having a high unit cost, and requiring trained staff and considerable infrastructure. Therefore others have looked into alternative gas analysis techniques with varying success. The electronic nose, first proposed in the 1980s, does not attempt to identify specific chemicals in a sample (as with GCMS), but instead analyses a sample as a whole (like the biological nose) using an array of non-selective gas sensors and a pattern recognition algorithm [5]. The technique can give an immediate result, uses air as the carrier gas, and can be made portable, simple to use, and at a unit-cost which is appropriate to the medical profession. To this end, there have been a number of published studies using different electronic nose instruments for the detection of TB biomarkers in both breath and sputum [6-8]. Though the studies show promise, these instruments have limitations. Firstly, some are based on sensors which rely on a chemical interaction between the surface of the sensor and the complex chemical sample. The surface layers succumb to long-term drift and are affected by changes in sample humidity and temperature. This means regular re-training of equipment is required to optimise performance. In addition, sensor manufacturing is such that each instrument is different and requires every instrument to be trained. An alternative approach is to use GC separation followed by QCM (Quartz Crystal Microbalance) detection. This instrument circumvents the aforementioned problems, but the requirement for a continuous helium supply restricts its use in resource-limited settings.

FAIMS (Field Asymmetric Ion Mobility Spectrometry) is a technique which is finding favour with researchers in the medical profession [9-10]. Like an electronic nose it does not attempt to identify specific chemicals within samples, but instead measures the mobility (or movement) of ionised molecules in high electric fields (kVolts). Pattern recognition techniques can then be applied to identify underlying trends within the mobility data, which can be correlated to the presence, or lack, of a disease process. Due to their incredibly high sensitivity (parts per billion or lower), reproducible results, use of air as the carrier gas, and portability, instruments using FAIMS have found favour with the military for the detection of chemical warfare agents. As these instruments measure a physical rather than chemical property of a molecule, they suffer far less from drift and instrument-to-instrument variability [11]. In this paper we report on the use of FAIMS for the analysis of breath in patients with confirmed TB compared with healthy controls.

**Methodology**

A commercial FAIMS unit was used (Lonestar, Owlstone UK) employing an ultra-violet (UV) ionisation source. Although this does not ionise the same range of chemicals as Ni63 (a radionuclide and the most commonly used ionisation method), units with a UV source can be used in UK hospitals without a special license and are easily transported from site to site. In a FAIMS unit, the complex mixture of gases contained within the sample (in our case breath) is pushed into the instrument where it is then ionised. The ionised sample then passes between two “separator” plates, to which an oscillating asynchronous waveform is applied (typically GHz). Onto these plates is applied a high positive potential for a short period of time, followed by a small negative potential for a longer period of time. Importantly, the applied amplitude x time are equal in both cases. When in use, the ratio of short/high potential to longer/smaller negative potential (described as the “dispersion field”) is cycled from 1 up to the systems maximum value. In the electric field generated by these potentials, ions are attracted to, repelled by or remain unaffected, depending upon the mobility difference between high- and low-field regime. If an ion then touches one of the plates it loses its charge, thus only ions with specific mobility exit the plates, still with their charge, and are detected. By applying a direct current (DC) voltage (called the compensation voltage) to one of the plates, it is possible to remove this attraction and repulsion, allowing more ions to be detected. This compensation voltage is cycled between a positive and negative potential and therefore for a certain compensation voltage, only chemicals with a specific mobility will be detected. The mobility of an ion depends on the mass, charge, size and shape, however as the field and the velocity are constant and we can relate the mobility to the velocity using the equation:

|  |  |
| --- | --- |
|  | (1) |

where and are the velocity and mobility of ion respectively and is the electric field strength. Since velocity changes is not proportional to the electric field intensity variations, at higher electric fields the ion mobility can be expressed by the following equation.

|  |  |
| --- | --- |
|  | (2) |

where and are the high-field mobility and low-field mobility respectively. and are compound specific values which account for high-field mobility effect, is carrier gas density number and is the electric field strength. More details of this process can be found in [12,13]. Typical breath outputs using a FAIMS instrument are shown in [9].

We recruited 23 adults (aged over 18 years) with clinically suspected pulmonary or extra-pulmonary TB (by constitutional symptoms or abnormal chest X-ray) from a UK teaching hospital (University Hospital Coventry and Warwickshire) over 6 months (March to September 2014). They were recruited from rapid-access TB clinics or in-patient admissions, before or within 1 week of commencing anti-TB medication. Diagnosis was later confirmed by culture, histology or radiology in 21 cases (average age 47 (18-86), 5 smokers). 2 cases had an alternative infective diagnosis (1 bacterial pneumonia, 1 *Staphylococcus aureu*s chronic hip osteomyelitis). 17 healthy controls were recruited for comparison, all interferon-gamma release assay negative (T.SPOT®.*TB*, Oxford Immunotec Ltd.). These 2 suspected TB cases with alternative final diagnosis were added to the healthy control group to make a total of 19 controls. (average age 38 (24-63), 5 smokers). A summary of cases by site of infection and culture result is shown in Table 1. Scientific and ethical approval was obtained from the local Research & Development Department and the Warwickshire Ethics Committee (reference number: 09/H1211/38). Written informed consent was obtained from all participants.

Participants were requested to refrain from eating and drinking for 2 hours before breath capture. Whole breath samples were collected, with participants breathing through a standard mouthpiece into a 3L tedlar bag repeatedly until the bag was full. Whole breath was used for simplicity, ease of capture, and to allow disposal of bags after single use for infection control purposes. Samples were analysed within 2 hours of collection using the Lonestar instrument (Owlstone UK). In our case, this was fitted with an ATLAS sampling system and a pump on the exhaust to pull the sample through the instrument. The bag output was connected in parallel to a split flow box containing a mass flow controller fed by filtered air. By controlling the speed of the pump and setting the mass flow controller to a specific flow, the flow out of the bag remained constant at 200 ml/min with the flow through the machine at 2L/min. The Lonestar was setup with a dispersion field cycled between 0 and 100% in 51 steps, and the compensation voltage between -6V and +6V in 512 steps. This allowed two complete output cycles to be collected.

The data from these experiments were processed using a previously developed algorithm described in [14]. In brief, each sample run generated 52,224 data points in a 2D matrix of dispersion fields and compensation voltages. First, a 2D wavelet transform (using Daubechies D4 wavelets) was applied to each sample run. This serves both as a data compression step, but also as a matched filter which can preferentially extract the diffuse ‘peaks’ in the data corresponding to different sets of chemical species. This has the effect of concentrating the signal in a relatively small number of wavelet coefficients, thus greatly simplifying and improving the subsequent analysis steps.

A threshold was then applied to the variance (across samples) of each wavelet coefficient, removing those coefficients that vary by only a very small amount (or not at all). This is known from (from prior work with FAIMS) to remove only uninformative and/or noise-dominated coefficients. This thresholding step therefore greatly reduces the dimensionality of the data, whilst only discarding data which are unlikely to aid disease prediction.

A 10-fold cross-validation was then applied, using 90% of the data as a training set, and the remaining 10% as a test set. Within each fold, a supervised feature selection step was performed to identify which features in the training set are the best for disease prediction. For this, a Wilcoxon rank sum test was used for each feature in turn, selecting those with the lowest p-value (using a training set only). Two features were extracted, which we have found to be sufficient in other studies. Predictions were then produced (disease/control) for the test set using just the two selected features.

Four different classifiers were used for prediction: Sparse Logistic Regression, Gaussian Process Classifier, Random Forest and Support Vector Machine. Of these, the Gaussian Process Classifier generated the best classification results. The Gaussian Process classifier outputs a predicted probability of a sample being from a TB patient, from which a receiver operator curve (ROC) and performance data were calculated.

**Results and Discussion**

Figure 1 shows the predictive probability using a Gaussian Process Classifier, showing a difference between the two groups (TB patients and controls). It was noted that different analysis techniques resulted in exactly the same sensitivity and specificity of 81% and 79% respectively. Figure 2 shows the ROC using the same analytical method. This result has an area under the curve of 0.92 (confidence interval at 95%, 0.84 to 1). Interestingly, just analysing the pulmonary TB patients gives a very similar result AUC = 0.91 (0.81 - 1).

The sensitivity and specificity reported here are not as high as previously reported using an electronic nose (sensitivity 93.5%, specificity 85.3%) [15]. In this study, however, Bruins *et al*. only included those with pulmonary TB, which may serve to reduce the number of false negatives thereby increasing sensitivity. Moreover they also excluded cases with HIV; we included 3 such cases. Despite HIV-TB co-infection being a common global scenario, there is a paucity of information on breath analysis in this context and whether, as for atypical clinical presentations of TB, it may also alter breath results.

Nevertheless, using this relatively crude sampling system, the results indicate that FAIMS shows promise for TB detection. Breath capture is limited by cross-reactivity with oro-pharyngeal microbiota, thus potentially reducing the diagnostic yield in TB. However, we are already in the process of improving our sample capture system and we believe that improved results are achievable.

**Conclusions**

The results indicate that FAIMS, and Ion Mobility Spectrometry (IMS) in particular, has potential as a technology that can be applied to breath testing for clinical diseases including TB. We recognise that we were limited by small sample sizes, sampling of whole breath rather than end-tidal, and undertaking the study in optimal UK hospital conditions as opposed to a low- or middle-income field setting. It is also difficult to determine whether the chemicals being detected were directly associated with the metabolism of the TB pathogen, or the body’s general response to infection. However, there is considerable potential for utilising IMS as a simple, rapid, cheap breath test for stratifying suspected TB patients. The advantages of IMS in terms of sensitivity, ease of setup with minimal training and minimal instrumental drift and variability, may provide a solution for targeting TB treatment in a global context. We are now looking to further study IMS technology in a more rigorous way, testing its efficacy for different breath capture techniques and in a variety of clinical circumstances.

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Table 1. Summary of cases by site of TB infection and culture result

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **TB site** | **Sample type** | **Culture positive** | **Culture negative\*** | **TOTAL** |
| Pulmonary | Sputum | 5 | 0 | **5** |
| Bronchoalveolar lavage fluid† | 2 | 2 | **4** |
| Lymph node | Tissue / pus | 5 | 1 | **6** |
| Spine / psoas abscess | Pus | 4 | 0 | **4** |
| Joint | Fluid | 1 | 0 | **1** |
| Other‡ |  | 0 | 1 | **1** |
|  |  |  | **TOTAL** | **21** |
| \* Histological or radiological diagnosis † 1 patient: lung tissue ‡1 patient: testicular TB | | | | |

Figure 1: Classification probablilities for the Control and TB groups. Applying a Wilcoxon rank-sum test to the sets of classification probabilities from the two groups, we get a p-value of 2.89\*10-6, showing that there is a highly statistically significant difference between the control and TB groups.

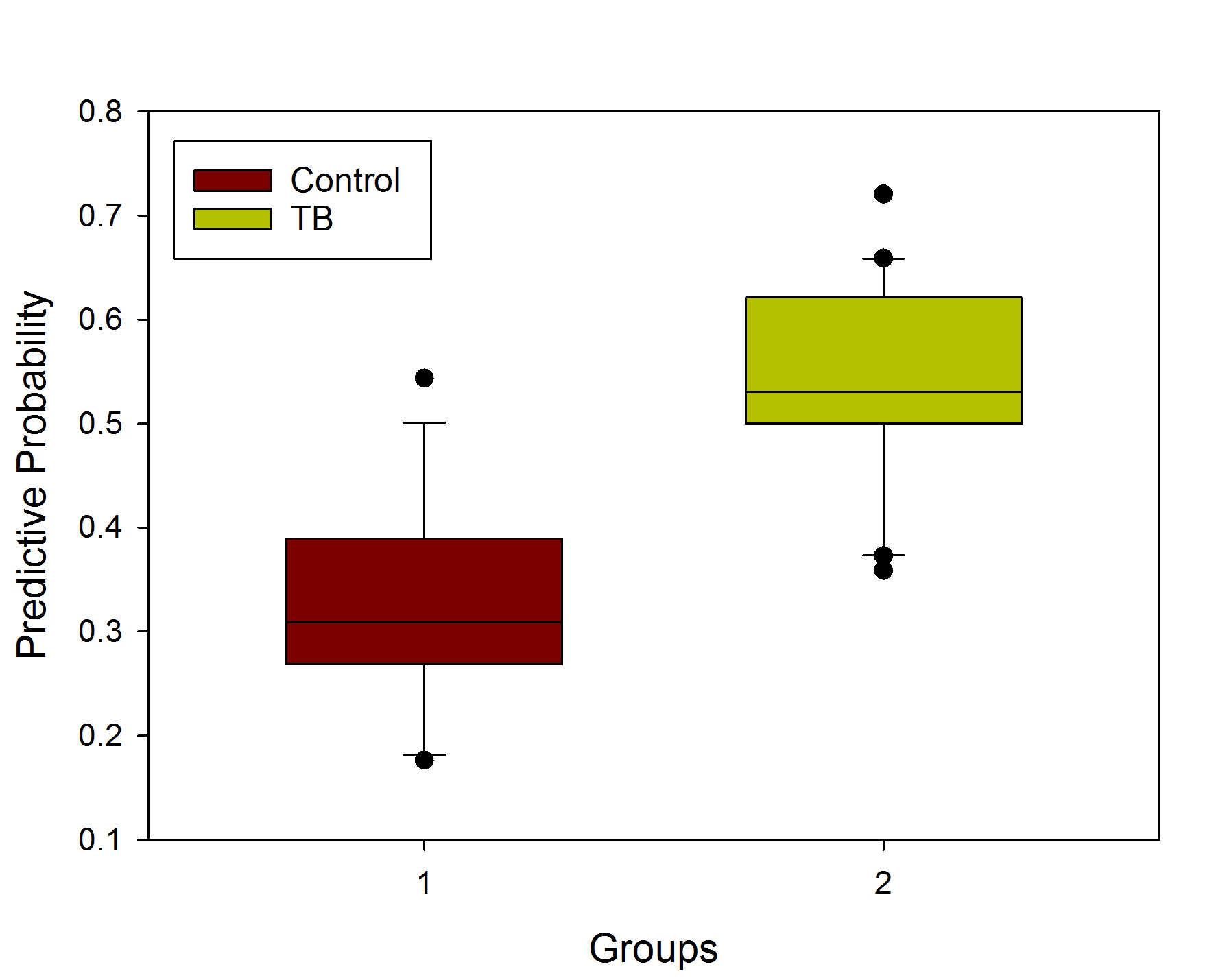


Figure 2: Receiver Operator Curve (AUC = 0.92; 95% CI: 084, 1) for breath analysis of patients with TB

