Introduction and objectives. NICE guidance advises neuroradiology to investigate CNS signs in patients with miliary tuberculosis (TB). The aims of our study were to describe our population of patients with tuberculomas in the presence of miliary disease and identify any clues to the best radiological modality.

Methods. The radiology and clinical history was retrospectively reviewed for all patients treated for miliary tuberculosis at one centre between 01/01/2009 and 31/12/2013.

Results. 53/1650 (3.2%) of patients during this period were diagnosed with miliary (disseminated) tuberculosis. 27/53 (50.9%) underwent further neuroimaging. 10/53 (18.9%) miliary TB patients had evidence of tuberculomas on neuroimaging (M:F 6:4, age range 22-81). 2/10 had evidence of tuberculomas on both CT (2/2 with contrast) and MRI, 5/10 had a negative CT (2/5 with contrast) but an MRI result which revealed tuberculomas. 3 patients did not have a CT scan (MRI only). All 10 patients were HIV negative and had fully sensitive TB, 9/10 had neurological signs which warranted the subsequent neuroimaging.

Conclusion. Tuberculomas are seen in a fifth of patients with miliary tuberculosis. Based on our findings, guidelines should be adapted to suggest that both use of contrast and MRI should be utilised preferentially.

Introduction. Despite efforts to improve Tuberculosis (TB) services, disease rates remain high (UK national average 14.4 per 100,000). We believe one of the ongoing challenges is engaging patients in attending outpatient clinics for care. However, there is no current UK data evaluating poor attendance to TB clinic.

Aim. To identify reasons for patient’s not attending TB clinic, in order to implement service improvements and increase patient engagement.

Methods. We conducted a prospective study reviewing the number of Did Not Attends (DNAs) to our TB clinic over a six-week period (April to June 2014). We evaluated data, usually obtained from patients who are contacted after they DNA, and cross referenced this with the trust electronic database. Data obtained included patient demographics, stage of TB treatment, route of referral, reasons for non-attendance and accessibility to clinic.

Results. 63 of 385 patients (16% - 42 males, 21 females) did not attend their TB clinic appointments compared to 15% for non-TB respiratory appointments in this time. 64% were contactable (25 males, 15 females). Median age was 32 (range 17–78 years), which included 16 ethnicities and seven languages. 62.5% were follow-up appointments and 37.5% were new. 27.5% had TB previously. Stage of TB treatment included: completed (17.5%), current (25%), none (57.5%). Referral route included GP (40%), hospital (32.5%) and contact tracing (27.5%). 59% were aware of their appointment but were unable to attend due to other engagements. 41% stated they had not received a letter informing them of their appointment, 13% of these patients had relocated to another area and not updated their address. 8% of patients highlighted problems with transport leading to difficulties accessing the clinic.

Conclusion. Communication to inform patients about appointments needs to be improved by both the referring and TB services. Utilising information technology and community links may improve patient education and therefore engagement with services. Experiencing the patient’s journey will highlight further areas for development.
recurrence patients had mean age = 42 yrs (range 18–83 yrs) and 6 were males. The sites of relapse were; pulmonary in 6 cases, 3 patients had intracranial tuberculomas, 2 patients had bony TB, 3 patients had TB lymphadenitis. The mean time to relapse/recurrence was 41 months (range 2–96 mo). All 14 patients responded favourably to re-treatment.

The true relapse rate of TB treated at the centre was 0.4%. The age, gender and ethnicity of the relapse cases were similar to the overall TB case-mix.

Conclusions Our true relapse/recurrence rate of TB is very low, and had no obvious risk factors. We cannot determine retrospectively whether these were recurrence or reinfection, but strain typing (DNA fingerprinting) could differentiate these.

M42 INCREASING COMPLEXITY OF TREATING TB IN OLDER PATIENTS
J Barrett, GA O’Hara, A Nundoll, N Price, H Milburn, RAM Breen. Guys and St Thomas; NHS Foundation Trust, London, UK
10.1136/thoraxjnl-2014-206260.430

Introduction Older adults remain an important reservoir of tuberculosis (TB) infection in the UK. Waning cellular immune responses, more frequent co-morbidities such as diabetes and malignancy, and increased polypharmacy may all modulate clinical presentation, treatment tolerability and ultimately outcomes when compared to younger individuals with TB. We sought to investigate this in our population.

Methods Retrospective study of all adults over 60 diagnosed with TB during a five year period at one hospital trust. Case-note and electronic record review established baseline disease features, co-morbidities, pre-morbid immune suppression including HIV status, TB-related outcomes and death. A randomly selected control group of identical size, containing adults aged 16–59 who were treated for TB during the same period, was used for comparison.

Results Forty-eight cases aged >60 years at TB diagnosis were identified. The case and control groups are described in the Table. Multi-lobar pulmonary disease was significantly more common in the >60 year old, as was diabetes, other significant co-morbidities and non-HIV immune suppression. Whilst treatment regimen discontinuation or alteration was more common in the >60 year old group (7 (14%) versus 3 (6%) if 60 years old but none (0%) of the younger group; whilst deaths after completion of TB treatment have been observed in 8 (17%) and 1 (2%) of cases respectively (no post-treatment death was related to TB in either group).

Abstract M42 Table 1

<table>
<thead>
<tr>
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<th>&lt;60 years old</th>
<th>&gt;60 years old</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age at diagnosis (range)</td>
<td>34 (16–57 years)</td>
<td>71 (60–88 years)</td>
</tr>
<tr>
<td>Male</td>
<td>24 (54%)</td>
<td>33 (69%)</td>
</tr>
<tr>
<td>Pulmonary disease</td>
<td>34 (70%)</td>
<td>33 (69%)</td>
</tr>
<tr>
<td>multi-lobar</td>
<td>9 (26%)</td>
<td>17 (52%)</td>
</tr>
<tr>
<td>Culture-confirmed TB</td>
<td>38 (79%)</td>
<td>36 (75%)</td>
</tr>
<tr>
<td>HIV-infected</td>
<td>6 (13%)</td>
<td>0</td>
</tr>
<tr>
<td>Diabetes</td>
<td>0</td>
<td>13 (27%)</td>
</tr>
<tr>
<td>Significant immune suppression</td>
<td>8 (17%)</td>
<td>29 (60%)</td>
</tr>
<tr>
<td>Significant co-morbidities</td>
<td>14 (35%)</td>
<td>36 (75%)</td>
</tr>
</tbody>
</table>

COPD: co-morbidities, deficiencies and interventions

M137 CAN STEROID INSENSITIVITY IN COPD PATIENTS BE RESTORED USING VITAMIN D?
D Mukherjee, D Parekh, R Dancer, M Ungurs, H Khorya, AM Turner. Queen Elizabeth Hospital Research Laboratories, University of Birmingham, Birmingham, UK
10.1136/thoraxjnl-2014-206260.432

Background NICE guidelines for TB diagnosis recommend that sputum is obtained for culture for all suspected cases of pulmonary TB, and biopsies for all cases extrapulmonary TB. As results can take 6 weeks, treatment initiation decisions are frequently made without microbiological confirmation.

Aim This study set out to examine the accuracy of clinical diagnoses in a high incidence area, and the basis for these decisions.

Methods The data entered onto the national TB database was used to obtain a list of patients for whom no culture results had been recorded. Clinic letters, laboratory records and imaging were examined to determine whether samples had been sent for culture, how diagnoses were made in the event of negative results, and if alternative diagnoses were concluded.

Results Of 323 patients on the database, 7% had no samples sent for culture. There were 109 culture negative patients, of whom 13 (4% all cases) had alternative diagnoses. A combination of relevant history and imaging was the most commonly used method of diagnosis when culture was negative (47%). Histology was used in 17% patients and Mantoux or IGRA testing supported initiating treatment in 39% cases. The database was missing positive culture results for 102 patients, of which four were MDR TB.

Conclusions In this study, we found accurate initial clinical diagnoses, with only 4% patients subsequently obtaining alternative diagnoses. Most diagnoses were made on the basis of relevant history and imaging. Of concern are the 7% patients for whom tissue was never sent for culture. This is likely to be an underestimate when including all patients initially suspected of TB, raising the possibility of missed diagnoses. The utility of Mantoux and IGRA testing in active disease is now disputed. It is hoped with inter-specialty education regarding the importance of culture and futility of immunological based assays, the proportion of patients with suspected TB who have sputum or tissue sent for culture increases. Accurate recording of MDR-TB on the national TB database needs to be improved, to enable efficient monitoring of intervention programmes.