Chronic pulmonary aspergillosis commonly complicates treated pulmonary tuberculosis with residual cavitation

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Chronic pulmonary aspergillosis (CPA) complicates treated pulmonary tuberculosis (TB), with high 5-year mortality. We measured CPA prevalence in this group.

398 Ugandans with treated pulmonary TB underwent clinical assessment, chest radiography and Aspergillus-specific IgG measurement. 285 were resurveyed 2 years later, including computed tomography of the thorax in 73 with suspected CPA. CPA was diagnosed in patients without active TB who had raised Aspergillus-specific IgG, radiological features of CPA and chronic cough or haemoptysis.

Author-defined CPA was present in 14 (4.9%, 95% CI 2.8–7.9%) resurvey patients. CPA was significantly more common in those with chest radiography cavitation (26% versus 0.8%; p<0.001), but possibly less frequent in HIV co-infected patients (3% versus 6.7%; p=0.177). The annual rate of new CPA development between surveys was 6.5% in those with chest radiography cavitation and 0.2% in those without (p<0.001). Absence of cavitation and pleural thickening on chest radiography had 100% negative predictive value for CPA. The combination of raised Aspergillus-specific IgG, chronic cough or haemoptysis and chest radiography cavitation had 85.7% sensitivity and 99.6% specificity for CPA diagnosis.

CPA commonly complicates treated pulmonary TB with residual chest radiography cavitation. Chest radiography alone can exclude CPA. Addition of serology can diagnose CPA with reasonable accuracy.

This article has supplementary material available from erj.ersjournals.com

Received: May 08 2018 | Accepted after revision: Dec 19 2018

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Introduction

Tuberculosis (TB) affected an estimated 10.4 million people in 2016, with 1.3 million attributable deaths in those without HIV and a further 374 000 deaths in HIV co-infected persons [1]. Just 57% of pulmonary TB cases reported to the World Health Organization (WHO) were bacteriologically confirmed. Chronic pulmonary aspergillosis (CPA) both complicates and mimics treated pulmonary TB [2, 3]. The prevalence of CPA in patients with treated TB and the contribution of misdiagnosed CPA to TB prevalence estimates are unclear.

The European Respiratory Society (ERS) published the first guidelines on CPA diagnosis and management in 2016 [4]. Chronic cavitary pulmonary aspergillosis (CCPA) is the most common form of CPA [5, 6]. It presents with progressive cavitation and symptoms including weight loss, fatigue, cough, breathlessness, chest discomfort and haemoptysis. The 5-year mortality of CCPA is 38–80% [3, 7]. However, surgery is curative in selected patients with localised disease and has been safely delivered in resource-poor settings [8–11]. Treatment with oral azole drugs can prevent clinical and radiological progression [12–14]. Chronic fibrosing pulmonary aspergillosis (CFPA) complicates untreated CCPA, with destruction of the lung [15]. Figure 1 illustrates the subtypes of CPA.

Sputum culture for Aspergillus is often negative in CPA [2]. Measurement of Aspergillus-specific IgG is central to CPA diagnosis [4, 16], but until recently had been inadequately validated for use in this context [17]. Both tests are infrequently available in areas of high TB prevalence [18].

Our recent studies demonstrate that Siemens Immulite Aspergillus-specific IgG is the best performing commercial assay for CPA diagnosis [5]. It has a sensitivity of 93.4% for UK CPA cases and specificity of 98.7% against healthy controls or 97.9% against Ugandan controls with treated pulmonary TB at a cut-off of 20 mg·L⁻¹, regardless of HIV status [19].

The global prevalence of CPA secondary to TB has been estimated at 1.2 million cases [20]. CPA has a global distribution, including over 200 reported cases in Africa [2, 3, 8, 11, 12, 21–26]. Prevalence estimates are based on the results of a 1968–1970 survey describing aspergilloma in 20% of 544 British patients with residual lung cavities after TB treatment [27]. That study was performed prior to computed tomography (CT) scanning and the HIV pandemic. Subsequent studies have demonstrated that aspergilloma is only present in a third of CPA cases [2].

We aimed to undertake the first accurate measurement of CPA prevalence. We conducted a community-based, cross-sectional prospective survey with systematic CT scanning of the thorax in an area of high TB prevalence. The prevalence of CPA in patients with and without HIV infection and chest radiography cavitation was compared, and the annual rate of CPA development between two surveys 2 years apart described.

Current CPA diagnostic criteria are arbitrary and based on expert opinion [4]. We aimed to conduct the first assessment of associations between raised Aspergillus-specific IgG and individual clinical or radiological features.
Pulmonary TB mostly occurs in resource-poor settings, where access to CT scanning of the thorax and Aspergillus-specific IgG testing is limited [18, 20]. We describe the accuracy of CPA diagnosis by chest radiography and clinical assessment, with or without serology, in comparison to CT-confirmed disease.

Methods

Adults aged ≥16 years, who were in the final month of pulmonary TB treatment or completed treatment in 2005 or later, were recruited in Gulu (Uganda) from October 2012 to January 2013. Patients were recruited from Gulu Regional Referral Hospital TB and HIV clinics or by Gulu District Health Office staff. Eligible patients had smear-positive TB or complete resolution of symptoms on therapy. We aimed to recruit 50% of patients with HIV co-infection at the time of TB treatment. Patients with no recorded HIV status who declined testing were ineligible.

Patients underwent structured clinical assessment. “Severe dyspnoea” represents a Medical Research Council dyspnoea score of 4 or 5. Chest radiography was performed at St Mary's Hospital at Lacor, Gulu. Serum was stored at −80°C at the Joint Clinical and Research Centre (JCRC) in Gulu. Immulite 2000 Aspergillus fumigatus-specific IgG testing (Siemens, Berlin, Germany) was performed at Christie Hospital (Manchester, UK) in July 2014. Raised levels are ≥20 mg·L⁻¹. CD4 testing was performed by FACSCalibur flow cytometry (Becton Dickinson, Franklin Lakes, NJ, USA) at the JCRC, if no recent result was documented (<6 months).

Patients were traced and reassessed (resurvey) between October 2014 and January 2015, with repeat chest radiography and Aspergillus-specific IgG. GeneXpert IV Mycobacterium tuberculosis nucleic acid amplification testing (Cepheid, Sunnyvale, CA, USA) was performed at Gulu Regional Referral Hospital for all patients who could provide a sputum sample. CT scanning of the thorax was performed using a Dual Slice scanner (GE, Boston, MA, USA) at the Kampala Imaging Centre (Uganda) on those with Aspergillus-specific IgG >10 mg·L⁻¹ or report of a fungal ball on chest radiography by any radiologist in the initial survey. Digital CT images were reported using Osirix X software (Pixmeo, Bernex, Switzerland). Two radiologists, blinded to clinical and serological findings, provided radiological reports. The senior respiratory radiologist (R.S.) at the UK National Aspergillosis Centre (Manchester, UK) provided a decisive report where initial reports differed.

In the first survey, probable CCPA was diagnosed when three criteria were met: 1) cough or haemoptysis for ≥1 month, 2) cavitation or a fungal ball on chest radiography and 3) raised Aspergillus-specific IgG. Probable simple aspergilloma was diagnosed in patients with a fungal ball on chest radiography and raised Aspergillus-specific IgG, but no chronic cough or haemoptysis. All “probable CPA” is the combination of patients with probable CCPA or probable simple aspergilloma.

CPA was definitively diagnosed after the resurvey. A modified version of the DENNING et al. [21] 2003 criteria was used. CCPA cases had all of: 1) cough or haemoptysis for ≥1 month, 2) raised Aspergillus-specific IgG, 3) absence of positive GeneXpert test for M. tuberculosis and 4) either paracavitary fibrosis or a fungal ball on CT of the thorax or progressive cavitation (either new cavitation or deterioration of pre-existing cavitation) on serial chest radiographs.

CFPA was diagnosed in patients with clinical and serological features of CCPA and complete destruction of at least two lobes of a lung on CT of the thorax. Simple aspergilloma was diagnosed in patients with a fungal ball on CT of the thorax and raised Aspergillus-specific IgG, but no cough or haemoptysis. The combined totals of CPA, CFPA and simple aspergilloma are presented as “all CPA”. A seronegative fungal ball was reported in patients with a fungal ball on CT of the thorax, but normal Aspergillus-specific IgG.

New CPA, developing during the 2-year follow-up period, was diagnosed in those who had normal Aspergillus-specific IgG in the initial survey, but CPA on resurvey, on the basis of resurvey criteria where CT scanning was performed and original survey criteria where it was not. Those patients diagnosed with CPA in the absence of CT of the thorax were reported as “probable CPA” in the final prevalence results.

Ethical approval

Ethical approval was granted by The University of Manchester Research Ethics Committee (7/6/12-11424), Gulu University Faculty of Medicine Institutional Review Board (GU/IRC/04/07/12), and the Uganda National Council for Science and Technology (HS 1253).

Statistical analysis

Statistical analysis was performed using SPSS version 20 (IBM, Armonk, NY, USA). The study was designed to measure the prevalence of CPA with an accuracy of ±2.3% based on a predicted prevalence of 6%. It had 80% power to detect an absolute difference of 9% in CPA prevalence between HIV-positive and
HIV-negative groups with 5% significance. Frequencies of categorical variables were compared by Fisher’s exact test throughout, unless otherwise stated. Mean results were compared by the t-test and median results by the Mann–Whitney U-test. 95% confidence intervals for disease prevalence were calculated by the Jeffreys method.

In additional post hoc analyses we assessed associations between potential clinical and environmental risk factors and CPA diagnosis or raised Aspergillus-specific IgG. We also assessed associations between individual symptoms and radiological findings and CPA diagnosis or raised Aspergillus-specific IgG. Odds ratios for associations between clinical or radiological features and CPA were calculated by bimodal regression analysis.

We calculated the sensitivity, specificity, and positive and negative predictive values of various combinations of symptoms, radiological results and Aspergillus-specific IgG testing in relation to final CPA diagnosis after CT of the thorax. The accuracy of chest radiography for diagnosis of CT-proven pathology was described in the same manner.

Results

Patient characteristics and initial survey results are shown in table 1. Median time from completion of TB treatment to recruitment was 29 months. Low CD4 counts were uncommon in HIV-positive patients. Cough and cavitation on chest radiography were more common in HIV-negative patients in the initial survey. 11 (2.8%) patients met the criteria for probable CPA, comprising nine (2.3%) with CCPA and two (0.5%) with simple aspergilloma.

| TABLE 1 Patient characteristics and initial survey results (2012) |
|---------------------------------------------------------------|
| Subjects | All patients | Resurveyed patients | Lost to follow-up or died | p-value | HIV-positive | HIV-negative | p-value |
| Subjects | 398 | 284 | 114 | 199 | 199 |
| HIV infection | 199 (50.0) | 135 (47.5) | 64 (56.1) | 0.149 | 199 | 199 |
| CD4 count <100 cells·µL−1 | 8 (2.0) | 3 (1.1) | 5 (4.4) | 0.066 | 8 (4.0) | 0 | 0.007 |
| Female | 155 (38.9) | 100 (35.2) | 55 (48.2) | 0.017 | 98 (49.2) | 57 (28.6) | <0.001 |
| Positive sputum smear at TB diagnosis | 303 (76.1) | 224 (78.9) | 79 (69.3) | 0.051 | 134 (67.3) | 169 (84.9) | <0.001 |
| On TB treatment | 10 (2.5) | 6 (2.1) | 4 (3.5) | 0.386 | 10 (5.0) | 0 | <0.001 |
| <1 year from TB treatment | 86 (21.6) | 56 (19.7) | 30 (26.3) | 0.386 | 83 (41.7) | 84 (42.7) | 0.662 |
| 1–3 years from TB treatment | 141 (35.4) | 105 (37.0) | 36 (31.6) | 0.386 | 60 (30.2) | 81 (40.7) | 0.007 |
| >3 years from TB treatment | 160 (40.2) | 116 (40.8) | 44 (38.6) | 0.386 | 93 (46.7) | 67 (33.7) | 0.007 |
| Traditional "grass-thatch" home | 371 (93.2) | 269 (94.7) | 102 (89.5) | 0.777 | 183 (92) | 188 (94.5) | 0.426 |
| Dampness reported in home | 119 (29.9) | 77 (27.1) | 42 (36.8) | 0.069 | 62 (31.2) | 57 (28.6) | 0.662 |
| Subsistence farmer | 373 (93.7) | 269 (94.7) | 104 (91.2) | 0.252 | 180 (90.5) | 193 (97) | 0.012 |
| Frequently cooks on open charcoal stove | 81 (20.4) | 57 (20.1) | 24 (21.1) | 0.891 | 45 (22.6) | 36 (18.1) | 0.319 |
| Smoking tobacco | 39 (9.8) | 30 (10.6) | 9 (7.9) | 0.462 | 16 (8.1) | 26 (13.1) | 0.042 |
| Age (mean [range]) years | 42 (16–83) | 42 | 41 | 0.332 | 40 (16–73) | 44 (16–83) | <0.001 |
| CD4 count in HIV-positive persons (median [range]) cells·µL−1 | 424 (14–1400) | 424 (59–1400) | 439 (14–869) | 0.664 |
| Cough* | 130 (32.7) | 92 (32.4) | 38 (33.3) | 0.906 | 51 (25.6) | 79 (39.7) | 0.004 |
| Haemoptysis* | 14 (3.5) | 11 (3.9) | 3 (2.6) | 0.755 | 5 (2.5) | 9 (4.5) | 0.415 |
| Fatigue* | 191 (48.0) | 138 (48.6) | 53 (46.5) | 0.740 | 96 (48.2) | 95 (47.7) | 1 |
| Fevers* | 99 (24.9) | 74 (26.1) | 25 (21.9) | 0.442 | 50 (25.1) | 49 (24.6) | 1 |
| Severe dyspnoea* | 90 (22.6) | 57 (20.1) | 33 (28.9) | 0.064 | 43 (21.6) | 47 (23.6) | 0.719 |
| Night sweats* | 131 (32.9) | 98 (34.5) | 33 (28.9) | 0.345 | 54 (27.1) | 77 (38.7) | 0.019 |
| Chest pain* | 214 (53.8) | 163 (57.4) | 51 (44.7) | 0.026 | 99 (49.7) | 115 (57.8) | 0.131 |
| Cavititation | 65 (16.3) | 46 (16.2) | 19 (16.7) | 0.882 | 24 (12.1) | 41 (20.6) | 0.029 |
| Paracavitary fibrosis | 34 (8.5) | 24 (8.5) | 10 (8.8) | 1 | 14 (7.0) | 20 (10.1) | 0.370 |
| Pleural thickening | 58 (14.6) | 44 (15.5) | 14 (12.3) | 0.530 | 23 (11.6) | 35 (17.6) | 0.118 |
| Fungal ball | 15 (3.8) | 9 (3.2) | 6 (5.3) | 0.383 | 5 (2.5) | 10 (5.0) | 0.292 |
| Aspergillus IgG >20 mg·L−1 | 21 (5.3) | 15 (5.3) | 6 (5.3) | 1 | 7 (3.5) | 14 (7.0) | 0.177 |

Data are presented as n or n (%), unless otherwise stated. TB: tuberculosis. #: present for >1 month; ¶: two-sided Chi-squared test. p-values calculated by Fisher’s exact test, except where indicated. Mean results compared by the t-test and median results by the Mann–Whitney U-test. p<0.05 considered significant.
At the 2-year follow-up, 285 (71.6%) patients were resurveyed and 19 (4.8%) were confirmed to have died between surveys. 94 (23.6%) could not be traced. Rates of loss to follow-up were similar in those with or without probable CPA (27.3% versus 24.0%; p=0.731). No patient with probable CPA died between surveys.

Results for resurvey patients are compared in table 2. While overall symptom frequency fell between surveys, frequency of haemoptysis and raised Aspergillus-specific IgG increased. Radiological findings consistent with CPA and elevated Aspergillus-specific IgG were more common in males (p=0.012–0.041), but symptoms consistent with CPA were not (p=0.270–0.830) (supplementary table S1).

CT of the thorax was performed in 73 (25.6%) resurvey patients, including 32 out of 46 (69.6%) with chest radiography cavitation in the original survey. CT of the thorax results and accuracy of chest radiography for identification of CT-proven pathology are shown in table 3. Two (0.7%) resurvey patients had a positive GeneXpert test out of 26 samples tested. Neither met diagnostic criteria for CPA.

Definite CPA was diagnosed in 14 (4.9%) resurvey patients. Full prevalence results are shown in table 4. Two patients had a seronegative fungal ball. Another two who did not undergo CT of the thorax developed probable CPA between surveys, comprising one with a new fungal ball on chest radiography and one with chest radiography cavitation, plus progressive paracavitary fibrosis and pleural thickening and new haemoptysis. In total, 18 (6.3%) resurvey cases had definite or probable CPA. There were an additional eight (2.8%) patients with radiological features of CPA, other than a fungal ball and characteristic symptoms, but negative Aspergillus-specific IgG.

CPA was present in 12 out of 46 (26.1%) resurvey patients with chest radiography cavities and two out of 239 (0.8%) patients with no such cavities (p<0.001). 12 (85.7%) of the 14 CPA cases had chest radiography cavities, all present in both surveys. Fungal ball was identified in nine (64.3%) CPA cases. Two (14.3%) CPA cases without a fungal ball had destroyed lung and one (7.1%) had progressive cavitation. Two (14.3%) CPA

| TABLE 2 Results for resurvey patients |
|--------------------------------------|
| **Initial** | **Resurvey** | **p-value** | **Resurvey** | **p-value** |
| Subjects | 284 | 284 | 135 | 150 |
| Cough\# | 92 [32.4] | 94 [33.0] | 0.929 | 34 [25.2] | 60 [40.0] | 0.008 |
| Haemoptysis\# | 11 [3.9] | 21 [7.4] | 0.1 | 11 [8.1] | 10 [6.7] | 0.657 |
| Fatigue\# | 138 [48.6] | 89 [31.2] | <0.001 | 40 [29.6] | 49 [32.7] | 0.610 |
| Fevers\# | 74 [26.1] | 26 [9.1] | <0.001 | 12 [8.9] | 14 [9.3] | 0.453 |
| Severe dyspnoea\# | 57 [20.1] | 3 [1.1] | <0.001 | 3 [2.2] | 0 | 0.105 |
| Night sweats\# | 98 [34.5] | 51 [17.9] | <0.001 | 22 [16.3] | 29 [19.3] | 0.539 |
| Chest pain\# | 163 [57.4] | 76 [26.7] | <0.001 | 31 [23.0] | 45 [30.0] | 0.227 |
| Cavitation | 46 [16.2] | 50 [17.5] | 0.737 | 20 [14.8] | 30 [20.0] | 0.277 |
| New cavitation | 4 [1.4] | 3 [2.2] | 0.737 | 1 [0.7] | 1 | 0.346 |
| Deterioration of pre-existing cavitation | 3 [1.1] | 3 [2.2] | 0 | 0 | 0.104 |
| All progressive cavitation | 7 [2.5] | 6 [4.4] | 0.737 | 1 [0.7] | 1 | 0.055 |
| Paracavitary fibrosis | 24 [8.5] | 36 [12.6] | 0.133 | 15 [11.1] | 21 [14.0] | 0.481 |
| New paracavitary fibrosis | 12 [4.2] | 4 [3.0] | 0.737 | 8 [5.3] | 5 | 0.387 |
| Deterioration of pre-existing paracavitary fibrosis | 6 [2.1] | 3 [2.2] | 0.737 | 3 [2.0] | 1 | 0.055 |
| All progressive paracavitary fibrosis | 18 [6.3] | 7 [5.2] | 0.737 | 11 [7.3] | 1 | 0.627 |
| Pleural thickening | 44 [15.5] | 50 [17.5] | 0.573 | 20 [14.8] | 30 [20.0] | 0.277 |
| New pleural thickening | 6 [2.1] | 3 [2.2] | 0.737 | 3 [2.0] | 1 | 0.055 |
| Deterioration of pre-existing pleural thickening | 10 [3.5] | 3 [2.2] | 0.737 | 7 [4.7] | 4 | 0.343 |
| All progressive pleural thickening | 16 [5.6] | 6 [4.4] | 0.737 | 10 [6.7] | 7 | 0.453 |
| Fungal ball | 9 [3.2] | 13 [4.6] | 0.515 | 4 [3.0] | 9 [6.0] | 0.265 |
| New fungal ball | 4 [1.4] | 2 [1.5] | 0.737 | 2 [1.3] | 1 | 0.055 |
| Deterioration of pre-existing fungal ball | 3 [1.1] | 1 [0.7] | 0.737 | 2 [1.3] | 1 | 0.055 |
| All progressive fungal ball | 7 [2.5] | 3 [2.2] | 0.737 | 4 [2.7] | 1 | 0.055 |
| Any radiological progression | 34 [11.9] | 15 [11.1] | 0.737 | 19 [12.7] | 19 [12.7] | 0.718 |
| *Aspergillus* IgG >20 mg·L\(^{-1}\) | 15 [5.3] | 28 [9.8] | 0.056 | 9 [6.7] | 19 [12.7] | 0.111 |

Data are presented as n or n (%), unless otherwise stated. \#: present for \(\geq 1\) month. All p-values calculated by Fisher’s exact test. p<0.05 considered significant.
## TABLE 3 Radiology results for the computed tomography (CT) of the thorax group

|                  | CT: radiologist 1 | CT: radiologist 2 | Final CT result | HIV-positive | HIV-negative | p-value | Chest radiography result | Sensitivity% | Specificity% | PPV% | NPV% |
|------------------|-------------------|-------------------|-----------------|--------------|--------------|---------|--------------------------|--------------|--------------|------|------|
| Subjects         | 73                | 73                | 73              | 26           | 47           |         | 73                       |              |              |      |      |
| Fungal ball      | 12 (16.4)         | 7 (9.6)           | 11 (15.1)       | 3 (11.5)     | 8 (17)       | 0.736   | 9 (12.3)                 | 36.4         | 91.9         | 44.4 | 89.1 |
| Cavitation       | 41 (56.2)         | 32 (43.8)         | 35 (47.9)       | 9 (34.6)     | 26 (55.3)    | 0.142   | 32 (43.8)                | 77.1         | 86.8         | 84.4 | 80.5 |
| Paracavitary fibrosis | 24 (32.9)     | 30 (41.1)         | 28 (38.4)       | 6 (23.1)     | 22 (46.8)    | 0.077   | 20 (27.4)                | 50           | 86.7         | 70   | 73.6 |
| Pleural thickening | 49 (67.1)        | 64 (87.7)         | 48 (65.8)       | 14 (53.8)    | 34 (72.3)    | 0.129   | 31 (42.5)                | 60.4         | 92           | 93.5 | 54.8 |
| Nodule(s)        | 68 (93.2)         | 58 (79.5)         | 62 (84.9)       | 20 (76.9)    | 42 (89.4)    | 0.183   |                          |              |              |      |      |
| Bronchiectasis   | 64 (87.7)         | 44 (60.3)         | 54 (74.0)       | 16 (61.5)    | 38 (80.9)    | 0.096   |                          |              |              |      |      |
| Destroyed lung   | 3 (4.1)           | 3 (4.1)           | 3 (4.1)         | 1 (3.8)      | 2 (4.3)      | 1       |                          |              |              |      |      |

Data are presented as n or n (%), unless otherwise stated. PPV: positive predictive value; NPV: negative predictive value. Nodules, bronchiectasis and destroyed lung were not reported for chest radiography. #: of chest radiography for identification of CT of the thorax findings. All p-values calculated by Fisher’s exact test. p<0.05 considered significant.
cases met radiological case criteria for CPA due to paracavitary fibrosis alone. Both also had pleural thickening (figure 2). Eight (57.1%) CPA cases had progressive chest radiography changes. All eight resurvey patients with probable CPA in the original survey met CT of the thorax criteria for CPA on resurvey.

Seven (2.5%) resurvey patients developed CPA between surveys, comprising five (1.8%) diagnosed by CT of the thorax and two (0.7%) diagnosed only by chest radiography criteria. Six (86%) of the new CPA cases had chest radiography cavitation on initial survey. The annual rate of CPA development was 1.25% in all patients, 6.5% in the 46 resurvey patients who had chest radiography cavitation on initial survey and 0.2% in the 239 resurvey patients who did not (p<0.001). The incidence of new CPA was similar in all "time from TB therapy" groups. Five (71.4%) of those who developed new CPA did so within 5 years of completion of TB therapy. Chest radiography progression in a new case of CPA is shown in figure 3.

| Subjects                        | All patients  | HIV-positive | HIV-negative | p-value |
|---------------------------------|---------------|--------------|--------------|---------|
| CCPA                            | 10 (3.5 (1.8–6.1)) | 2 (1.5 (0.3–4.7)) | 8 (5.3 (2.6–9.8)) | 0.108 |
| CFPA                            | 3 (1.1 (0.3–2.8)) | 1 (0.7 (0.1–3.4)) | 2 (1.3 (0.3–4.2)) | 1 |
| Simple aspergilloma             | 1 (0.4 (0–1.6)) | 1 (0.7 (0.1–3.4)) | 0 (0 (0–1.7)) | 0.474 |
| All definite CPA                | 14 (4.9 (2.8–7.9)) | 4 (3.0 (1–6.9)) | 10 (6.7 (3.5–11.5)) | 0.177 |
| Seronegative fungal ball        | 2 (0.7 (0.1–2.2)) | 1 (0.7 (0.1–3.4)) | 1 (0.7 (0.1–3.1)) | 1 |
| Probable CPA in non-CT group    | 2 (0.7 (0.1–2.2)) | 2 (1.5 (0.3–4.7)) | 0 (0 (0–1.7)) | 0.223 |
| All definite and probable CPA   | 18 (6.3 (3.9–9.6)) | 7 (5.2 (2.3–9.9)) | 11 (7.3 (4–12.3)) | 0.478 |

Data are presented as n or n [% (95% CI)], unless otherwise stated. CCPA: chronic cavitary pulmonary aspergillosis; CFPA: chronic fibrosing pulmonary aspergillosis; CT: computed tomography. All p-values calculated by Fisher’s exact test. p<0.05 considered significant.

FIGURE 2 Radiological presentations of chronic pulmonary aspergillosis (CPA) in Gulu. a) Chronic cavitary pulmonary aspergillosis (CCPA) with multiple cavities and a fungal ball. b) CCPA with a large cavity, paracavitary fibrosis and adjacent pleural thickening, but no fungal ball. c) Chronic fibrosing pulmonary aspergillosis with complete destruction of the whole right lung with multiple cavities and surrounding fibrosis.
No associations were found between CPA diagnosis and potential clinical and radiological risk factors (supplementary table S2). Associations between individual symptoms and raised Aspergillus-specific IgG or CPA are shown in table 5. Cough and haemoptysis were significantly more common in those with raised Aspergillus-specific IgG, but other symptoms were not.

Fungal ball, cavitation, paracavitary fibrosis and pleural thickening were all significantly more common in those with raised Aspergillus-specific IgG, but bronchiectasis and nodules were not. Every patient with destroyed lung had raised Aspergillus-specific IgG and CPA. All forms of chest radiography progression were more common in CPA, except progressive cavitation.

The accuracy of combinations of symptoms and tests for CPA diagnosis is shown in table 6. All CPA cases had either cavitation or pleural thickening on chest radiography, but no combination of chest radiography findings and clinical assessment produced a positive predictive value >40%. Addition of Aspergillus-specific IgG measurement allowed CPA diagnosis with a positive predictive value of 92.3% and a negative predictive value of 99.3%.

Discussion
We have performed the first community-based survey of persons with treated pulmonary TB to measure CPA prevalence. It is the first CPA prevalence survey to achieve accurate diagnosis through systematic CT of the thorax and use of an Aspergillus-specific IgG assay validated for the study population.

The finding that CPA complicates 4.9–6.3% of all treated pulmonary TB cases and 26% of cases with chest radiography cavitation has profound implications for TB management programmes. It suggests that testing for CPA with Aspergillus-specific IgG should be routine in all patients with recurrent cough, haemoptysis or weight loss following treatment of pulmonary TB, regardless of HIV status or time from completion of TB therapy. Those with raised levels should have imaging, preferably with CT of the thorax, to confirm the diagnosis. As CPA mimics the presentation of pulmonary TB, testing should also form part of routine investigations in any patient where empirical treatment for “smear-negative TB” is being considered.

Patients with chest radiography cavities had an annual rate of new CPA of at least 6.5%, a condition associated with 38–80% 5-year mortality. This suggests that all patients should have chest radiography at the end of pulmonary TB treatment and that those with cavities should be monitored for CPA to facilitate early treatment. We identified new cases of CPA occurring up to 9 years after completion of TB therapy. The frequency of new CPA occurring beyond this point is unknown.

We have probably underestimated the frequency of CPA. 10 patients had a combination of symptoms and radiological features suggestive of CPA despite normal A. fumigatus-specific IgG, including two with a
| Symptoms | All patients | Aspergillus-specific IgG-positive | Aspergillus-specific IgG-negative | p-value | OR (95% CI) CPA present | CPA absent | p-value | OR (95% CI) |
|----------|-------------|---------------------------------|---------------------------------|--------|-----------------------|-----------|--------|----------------|
| Cough    | n=285       | 15 (53.6)                       | 79 (30.7)                       | 0.015  | 2.60 (1.18–5.72)      | 11 (78.6) | 83 (30.6) | <0.001 8.30 (2.26–30.55) |
| Haemoptysis | 21 (7.4)  | 8 (28.6)                        | 13 (5.1)                        | <0.001 | 7.51 (2.78–20.24)     | 7 (50.0)  | 14 (5.2) | <0.001 18.36 (5.65–59.60) |
| Fatigue  | 89 (31.2)   | 9 (32.1)                        | 80 (31.1)                       | 0.912  | 1.05 (0.45–2.42)      | 6 (42.9)  | 83 (30.6) | 0.336 1.70 (0.57–5.05)   |
| Fevers   | 26 (9.1)    | 3 (10.7)                        | 23 (8.9)                        | 0.729  | 1.22 (0.34–4.35)      | 2 (14.3)  | 24 (8.9) | 0.371 1.71 (0.36–8.12)   |
| Night sweats | 51 (17.9) | 6 (21.4)                        | 45 (17.5)                       | 0.607  | 1.25 (0.49–3.35)      | 4 (28.6)  | 47 (17.3) | 0.287 1.91 (0.57–6.34)   |
| Chest pain | 76 (26.7) | 11 (39.3)                       | 65 (25.3)                       | 0.112  | 1.91 (0.85–4.29)      | 7 (50.0)  | 69 (25.5) | 0.06 2.93 (0.99–8.64)    |

| Chest radiography (resurvey) | All patients | Aspergillus-specific IgG-positive | Aspergillus-specific IgG-negative | p-value | OR (95% CI) CPA present | CPA absent | p-value | OR (95% CI) |
|------------------------------|-------------|---------------------------------|---------------------------------|--------|-----------------------|-----------|--------|----------------|
| Cavitation                   | n=285       | 17 (60.7)                       | 33 (12.8)                       | <0.001 | 10.49 (4.52–23.34)    | 12 (85.7) | 38 (14.0) | <0.001 36.79 (7.92–170.87) |
| Fungal ball                  | 13 (4.6)    | 5 (17.9)                        | 8 (3.1)                         | <0.001 | 6.77 (2.05–22.38)     | 4 (28.6)  | 9 (3.3)  | 0.002 11.64 (3.06–44.31) |
| Paracavitary fibrosis        | 36 (12.6)   | 7 (25.0)                        | 29 (11.3)                       | 0.038  | 2.62 (1.02–6.7)       | 4 (28.6)  | 32 (11.8) | 0.085 2.99 (0.88–10.09)   |
| Pleural thickening           | 50 (17.5)   | 16 (57.1)                       | 34 (13.2)                       | <0.001 | 8.74 (3.81–20.07)     | 10 (71.4) | 40 (14.8) | <0.001 14.44 (4.32–48.28) |

| Chest radiography progression | All patients | Aspergillus-specific IgG-positive | Aspergillus-specific IgG-negative | p-value | OR (95% CI) CPA present | CPA absent | p-value | OR (95% CI) |
|-------------------------------|-------------|---------------------------------|---------------------------------|--------|-----------------------|-----------|--------|----------------|
| Cavitation                    | n=285       | 4 (14.3)                        | 3 (1.2)                         | 0.002  | 14.06 (2.97–66.51)    | 1 (7.1)   | 6 (2.2)  | 0.301 3.38 (0.38–30.21) |
| Fungal ball                   | 7 (2.5)     | 5 (17.9)                        | 4 (1.6)                         | 0.023  | 7.56 (1.60–35.70)     | 2 (14.3)  | 15 (5.5) | 0.05 6.64 (1.17–34.81)   |
| Paracavitary fibrosis         | 16 (6.3)    | 5 (17.9)                        | 13 (5.1)                        | 0.008  | 4.06 (1.33–12.41)     | 3 (21.4)  | 15 (5.5) | 0.05 4.64 (1.17–34.81)   |
| Pleural thickening            | 16 (5.6)    | 5 (17.9)                        | 11 (4.3)                        | 0.003  | 4.84 (1.55–15.14)     | 3 (21.4)  | 14 (4.8) | 0.037 5.39 (1.34–21.71) |
| Any progressive change        | 34 (11.9)   | 12 (42.9)                       | 22 (8.6)                        | <0.001 | 8.01 (3.37–19.06)     | 7 (50.0)  | 27 (10.0) | <0.001 9.04 (2.95–27.71) |

| CT of the thorax | All patients | Aspergillus-specific IgG-positive | Aspergillus-specific IgG-negative | p-value | OR (95% CI) CPA present | CPA absent | p-value | OR (95% CI) |
|------------------|-------------|---------------------------------|---------------------------------|--------|-----------------------|-----------|--------|----------------|
| Fungal ball      | n=73        | 9 (42.9)                        | 2 (3.8)                         | <0.001 | 18.75 (3.58–98.28)    | 9 (64.3)  | 2 (3.4)  | <0.001 51.3 (8.61–305.47) |
| Cavitation       | 35 (47.9)   | 16 (17.6)                       | 19 (36.5)                       | 0.002  | 5.56 (1.76–17.59)     | 14 (100)  | 21 (35.6) | <0.001 14.55 (5.50–50.27) |
| Paracavitary fibrosis | 28 (38.4)  | 16 (17.6)                       | 12 (23.1)                       | <0.001 | 10.67 (3.23–35.19)    | 14 (100)  | 14 (23.7) | <0.001 8.83 (1.55–50.27) |
| Pleural thickening | 48 (65.8)  | 19 (19.5)                       | 29 (55.8)                       | 0.005  | 7.53 (1.59–35.73)     | 14 (100)  | 34 (57.6) | 0.003 6.64 (1.17–34.81)   |
| Nodule(s)        | 62 (84.9)   | 16 (17.6)                       | 46 (88.5)                       | 0.185  | 0.42 (0.11–1.56)      | 10 (71.4) | 52 (88.1) | 0.116 0.337 (0.08–1.37)   |
| Bronchiectasis   | 54 (74.0)   | 16 (17.6)                       | 38 (73.1)                       | 0.784  | 1.17 (0.36–3.82)      | 12 (85.7) | 42 (71.2) | 0.265 2.43 (0.49–12.02)   |
| Destroyed lung   | 3 (4.1)     | 3 (14.3)                        | 0                               | 0.021  | 3 (21.4)              | 0         | 0       | 0.006 1.00 (0.05–20.00)   |

Data are presented as n (%), unless otherwise stated. CPA: chronic pulmonary aspergillosis; CT: computed tomography. #, these features formed part of the CPA case definition. All p-values calculated by Fisher’s exact test. Odds ratios were calculated by bimodal regression analysis. p<0.05 considered significant.
fungal ball on CT of the thorax. These cases probably represent CPA cases caused by another species, as Aspergillus fumigatus-specific IgG may have low sensitivity in this group [17]. One patient also had HIV with a CD4 count <200 cells·µL⁻¹. The accuracy of Aspergillus-specific IgG in this group is poorly described.

Our study has limitations. Convenience sampling allows recruitment bias, although this was minimised through active recruitment of eligible patients. Our case definition was written prior to the publication of the ERS guidelines [4]. These guidelines require either radiological or symptomatic evidence of disease duration >3 months for CCPA/CFPA. All our CCPA/CFPA patients met these criteria as 10 out of 13 (77.0%) had symptoms for >3 months and three had chest radiography evidence of CPA in both surveys. The guidelines also require either a fungal ball on CT of the thorax or the combination of cavitation and radiological progression. 13 out of 14 (93%) CPA cases met these criteria.

Subacute invasive pulmonary aspergillosis typically kills within a few months and is present in 2–11% of AIDS autopsies [28–32]. We used a 1-month symptom duration definition to allow inclusion of these patients, but we will not have identified patients who developed aspergillosis and died outside the survey periods.

Existing consensus case definitions for CPA are arbitrary and based on expert opinion. We have assessed associations between individual symptoms and raised Aspergillus-specific IgG and CPA for the first time. It may be appropriate to restrict future case definitions to cough and haemoptysis. Pleural thickening was present in most CPA cases and was significantly more common in those with CPA or raised Aspergillus-specific IgG. It may be appropriate to include CT of the thorax cavitation plus adjacent pleural thickening as radiological evidence of CPA in future case definitions. Progressive cavitation was the only form of chest radiography progression not associated with CPA diagnosis. Every patient with destroyed lung had raised Aspergillus-specific IgG and CPA, mandating investigations for CPA in this group.

Aspergillus colonisation of bronchiectatic airways is unlikely to be the main cause of raised Aspergillus-specific IgG in this population as there was no association between bronchiectasis and raised Aspergillus-specific IgG. The absence of any association between nodules and raised Aspergillus-specific IgG supports the current recommendation that biopsy is required to diagnose Aspergillus nodule disease [4, 33].

Prior studies demonstrate the presence of aspergillosis in other post-TB populations. The 1968–1970 study found aspergilloma in 20% of 544 British treated TB patients with residual cavities [27]. New Aspergillus-specific antibodies developed in 9.8% and aspergilloma in 5.4% of patients during 2 years of follow-up, indicating an annual rate of aspergilloma development of 2.7%. Aspergillus-specific antibodies were found in 8.3% of 350 predominantly treated TB patients in Brazil [34], 23–25% of 77 Indian treated TB patients [35, 36], 19% of 226 Japanese patients with active or treated TB [37] and 9% of 101 HIV-positive patients completing pulmonary TB treatment in Kampala [38]. Chest radiography plus serology surveys have found probable CPA in 8.7% of 208 patients completing TB therapy in Nigeria [22] and 11.3% of 124 HIV-negative patients with active or treated TB in Iran [23].

These prior studies are not directly comparable to ours as they took place in select clinical populations, used unvalidated or insensitive serology methods, lacked systematic CT imaging of the thorax and

**TABLE 6 Accuracy of test combinations for chronic pulmonary aspergillosis (CPA) diagnosis in the absence of computed tomography of the thorax**

| Test                                      | n  | Sensitivity % | Specificity % | PPV % | NPV % |
|-------------------------------------------|----|---------------|---------------|-------|-------|
| Serology                                  | 28 | 100           | 94.8          | 50.0  | 100*  |
| Symptoms                                  | 101| 92.9          | 67.5          | 12.9  | 99.5  |
| Serology plus symptoms                    | 18 | 92.9          | 98.2          | 72.2  | 99.6  |
| Chest radiography fungal ball             | 13 | 28.6          | 96.7          | 30.8  | 96.3  |
| Chest radiography cavitation              | 50 | 85.7          | 86.0          | 24.0  | 99.1  |
| Chest radiography pleural thickening      | 50 | 71.4          | 85.2          | 20.0  | 98.3  |
| Chest radiography cavitation or pleural thickening | 73 | 100           | 78.2          | 19.2  | 100   |
| Chest radiography cavitation plus symptoms| 30 | 85.7          | 93.4          | 40.0  | 93.4  |
| Chest radiography cavitation or pleural thickening plus symptoms | 41 | 92.9          | 89.7          | 31.7  | 99.6  |
| Serology plus chest radiography cavitation| 17 | 85.7          | 98.2          | 70.6  | 99.3  |
| Serology plus chest radiography cavitation plus symptoms | 13 | 85.7          | 99.6          | 92.3  | 99.3  |
| Serology plus chest radiography cavitation or pleural thickening plus symptoms | 17 | 92.3          | 98.5          | 76.5  | 99.6  |

PPV: positive predictive value; NPV: negative predictive value. *: positive serology was a mandatory diagnostic criterion for CPA in this study.
sometimes included non-TB patients in mixed cohorts. Nonetheless, aspergillosis was present at clinically important levels in every population studied and all the studies were consistent with CPA prevalence at least as high as our findings. The combination of our data and these prior studies demonstrates that CPA complicating pulmonary TB represents an unrecognized global public health issue.

Between 2000 and 2016, around 45 million lives were saved due to effective treatment of pulmonary TB worldwide [1]. However, 1.6–3.5 million of these would subsequently have developed CPA if this occurs within the 95% confidence interval identified in our study. Residual chest radiography cavitation is found in 20–30% of the 7.7 million patients cured of pulmonary TB each year [1]. If the annual rates of new CPA we found in those with and without residual cavitation are applied, then 112,000–160,000 people will develop CPA worldwide every year after completion of TB therapy.

This is probably an underestimate as we only diagnosed “new” CPA in those with negative baseline serology. Without baseline CT of the thorax we could not calculate the additional rate of “new” CPA in those with positive baseline serology but no radiological features. This estimate may also be low because it assumes that all patients with pulmonary TB actually had this disease and not CPA, and it omits any patients with both disorders concurrently who died during anti-TB therapy, with undiagnosed CPA.

Fungal ball is absent in most CPA cases [2]. Aspergillus-specific IgG and CT of the thorax are therefore required to reliably diagnose CPA cases. Unfortunately, both are rarely available in resource-poor settings [18]. In their absence patients with CPA meet WHO diagnostic criteria for “smear-negative TB” [39]. A recent Nigerian study found probable CPA in 19% of such patients, who were HIV-negative [22]. We found that a fungal ball on chest radiography was insensitive for diagnosing CPA. However, CPA could be excluded by chest radiography in this cohort and diagnosed with acceptable accuracy if Aspergillus-specific IgG measurement was also performed.

We therefore propose that CPA should be diagnosed in persons with treated TB in resource-poor settings if recurrent TB has been excluded by GeneXpert PCR testing and a combination of raised Aspergillus-specific IgG, chest radiography cavitation and either chronic cough or haemoptysis is present. This proposal requires validation in another population. Improved access to Aspergillus-specific IgG is necessary to diagnose CPA in resource-poor settings. Implementation of point-of-care tests, optimised for resource-poor settings, would facilitate this. A lateral flow device for Aspergillus-specific IgG has recently been commercialised in Europe (LDBio, Lyon, France).

Untreated CPA is usually fatal, but treatment is possible with relatively inexpensive oral generic antifungals, now included on the WHO Essential Medicine List [40], or surgery in selected cases. Meaningful improvements in TB-associated mortality are therefore possible by addressing CPA in a programmatic way within existing TB care systems. Public health bodies should now act to ensure equitable access to CPA diagnosis and treatment.

Acknowledgements: We would like to thank study assistants Patrick Kidega and Thomas Okumu (Gulu Regional Referral Hospital, Gulu, Uganda) for translation, phlebotomy and sample handling. Marion Sorgenor, Helen Slattery and the Gulu Manchester Link team (Manchester University NHS Foundation Trust, Manchester, UK) for provision of accommodation in Gulu for study members and for logistical and practical assistance throughout the study. Staff at Gulu Regional Referral Hospital TB and HIV clinics for providing space to assess patients, identifying potential recruits, and facilitating sample storage and testing of sputum samples. All the staff at Gulu Regional District Health Office Tuberculosis and Leprosy Control department for identifying eligible patients and assisting in recruitment. Staff at the radiology departments at St Mary’s Hospital, Lacor and Kampala Imaging Centre (Uganda) for their help with processing radiography and CT scans. Laboratory staff at Christie Hospital (Manchester, UK) for allowing access to their Siemens Immulite 2000 system to perform Aspergillus-specific IgG testing. Staff from Siemens for providing training on the Immulite 2000. Julie Morris (University of Manchester, Manchester, UK) for statistical support. University of Manchester students Andrew Mockridge, Matthew Kneale and Michael Clarke for their assistance with study set-up and recruitment.

Conflict of interest: I.D. Page reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study; grants and test kit donation from Serion and Dynamiker, test kit donation from Genesis and OLM Medical, and travel grants for conference attendance from the British Infection Association, British Society for Medical Mycology, International Society for Human and Animal Mycology, European Society for Clinical Microbiology and Infectious Diseases, and Conference for Retroviruses and Opportunistic Infections, outside the submitted work. R. Byanyima reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study. N. Onyachi reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study. C. Opira reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study. M. Richardson reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study.
the study; personal fees for talks and consultancy from Gilead Sciences Europe, MSD and Basilea, outside the submitted work. R. Sawyer reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study. A. Sharman reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study. D.W. Denning reports grants from Astellas Pharma Europe, test kit donations from Siemens, grants and accommodation for study team from Gulu Manchester Link, during the conduct of the study; has founder shares with F2G, has received personal fees for consultancy from Scynexis, Cidara, Biothergen, Quintiles, Pulmatrix, Zambon and Fujifilm, personal fees for lectures from Astellas, Dynamiker, Gilead, Merck and Pfizer, outside the submitted work; and is a long-standing member of the Infectious Disease Society of America Aspergillosis Guidelines group, the European Society for Clinical Microbiology and Infectious Diseases Aspergillosis Guidelines group, and the British Society for Medical Mycology Standards of Care committee.

Support statement: The study was funded by an investigator-driven study grant from Astellas Pharma Europe Ltd. Further financial support and accommodation for study members in Gulu was provided by the Gulu Manchester Link charity, based at Wythenshawe Hospital (Manchester, UK). Aspergillus–specific IgG test kits used in the study were donated by Siemens. No funder had any role in study design or interpretation of results. Funding information for this article has been deposited with the Crossref Funder Registry.

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https://doi.org/10.1183/13993003.01184-2018