AUDIT, RESEARCH AND GUIDELINE UPDATE

US Cystic Fibrosis Foundation and European Cystic Fibrosis Society consensus recommendations for the management of non-tuberculous mycobacteria in individuals with cystic fibrosis: executive summary

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ABSTRACT
Non-tuberculous mycobacteria (NTM) are ubiquitous environmental organisms that can cause chronic pulmonary infection, particularly in individuals with pre-existing inflammatory lung disease, such as cystic fibrosis (CF). Pulmonary disease (PD) caused by NTM has emerged as a major threat to the health of individuals with CF, but remains difficult to diagnose and problematic to treat. In response to this challenge, the US Cystic Fibrosis Foundation (CFF) and the European Cystic Fibrosis Society (ECFS) convened a panel of 19 experts to develop consensus recommendations for the screening, investigation, diagnosis and management of NTM-PD in individuals with CF. PICO (population, intervention, comparison, outcome) methodology and systematic literature reviews were employed to inform draft recommendations, which were then modified to achieve consensus and subsequently circulated for public consultation within the USA and European CF communities. We have thus generated a series of pragmatic, evidence-based recommendations as an initial step in optimising management for this challenging condition.

BACKGROUND
Non-tuberculous mycobacteria (NTM) are increasingly being isolated from the sputum of adults and children with cystic fibrosis (CF) both in North America and Europe. Estimates of the prevalence of NTM in the CF population have ranged from 1.3% in the earliest study reported in 19841 to 32.7% in a review of patients with CF over age 40 in Colorado.2

The NTM species most commonly identified in individuals with CF from North America and Europe are the slow growing Mycobacterium avium complex (MAC, including M. avium, M. intracellulare and M. chimaera), which can be found in up to 72% of NTM-positive sputum cultures, and the rapid growing M. abscessus complex (comprising the subspecies M. abscessus subsp abscessus (M. a. abscessus), M. a. bolletii and M. a. massiliense (the latter currently classified as part of M. a. bolletii)), which in many centres has now become the most common NTM isolated from individuals with CF.

There has been a rise in the prevalence of NTM-positive cultures in respiratory samples from individuals with CF over the last three decades, which probably reflects a true increase in the frequency of NTM infection. A number of CF studies (eg, Renna et al3) show year-on-year increases in NTM-positive cultures with no change in surveillance intensity or culture methodology.

Possible reasons for increased NTM-positive cultures in individuals with CF include: increases in environmental exposure to NTM through more permissive temperature settings of home water heaters and more contact with shower aerosols, increased antibiotic usage creating more NTM permissive lung niches, greater chronic use of medications that might impair host immunity to NTM and/or spread of NTM through person-to-person transmission.4

NTM can cause progressive inflammatory lung damage, a condition termed ‘NTM pulmonary disease’ (NTM-PD), which is defined by the presence of specific microbiological, clinical and radiological features.5 However, it has become clear that NTM can also transiently, intermittently or permanently reside within the lungs of CF individuals without causing NTM-PD, thus representing asymptomatic infection and creating considerable difficulties in deciding how best to screen for and diagnose NTM.

Further challenges exist in knowing how best to identify NTM in respiratory samples, when and how to initiate treatment for NTM-PD and how NTM may impact individuals under consideration for lung transplantation. As a consequence, the Cystic Fibrosis Foundation (CFF) and European Cystic Fibrosis Society (ECFS) sought to generate a consensus recommendations document to support and standardise the management of NTM infection in children and adults with CF, permitting
prospective evaluation of current best practice and forming a foundation for future research programmes.

METHODS

The CFF and the ECFS invited experts to participate in the statement development process. The 19 member committee consisted of professionals with expertise in CF and NTM and included adult and paediatric CF physicians, lung transplant physicians, microbiologists, infectious disease specialists and a parent of an individual with CF. The committee convened in May 2012 and divided into five subgroups, each responsible for a specific topic: epidemiology and risk factors, screening, microbiology, clinical criteria for the diagnosis of NTM pulmonary disease should be used in individuals with CF, drug susceptibility testing should be performed in accordance with CLSI guidelines, and drug intolerance and drug-related toxicity occur frequently and changes in antibiotic therapy are often required.

Microbiology

- All NTM isolates from individuals with CF should undergo molecular identification.
- All NTM isolates should be identified to the species level, except for *M. intracellulare*, *M. avium* and *M. chimaera*, where identification can be limited to *M. avium complex (MAC)*, and *M. abscessus complex*, which should be sub-specified.
- The MAC bacteriology susceptibility testing should be performed on an isolate recovered prior to initiation of treatment. Clarithromycin susceptibility testing should also be performed on subsequent isolates if the patient a) fails to culture convert after six months of NTM treatment; b) recultures *M. avium complex* after initial culture conversion while on NTM treatment; or c) recultures *M. avium complex* after completion of NTM treatment.
- *M. abscessus complex*, susceptibility testing should include at least clarithromycin, cefoxitin and amikacin (and preferably also ticarcillin, imipenem, minocycline, moxifloxacin and linezolid).
- Drug susceptibility testing should be performed in accordance with CLSI guidelines.

Drug Intolerance

- Other CF pathogens and co-morbidities should be considered as potential contributors to a patient’s symptoms and radiological features when determining the clinical significance of NTM positive cultures.

NTM treatment should be considered for individuals with CF who have ATS/IDSA defined NTM pulmonary disease.

Treatment of *M. abscessus complex* pulmonary disease should involve an intensive phase followed by a continuation phase.

The initial phase should include a daily oral macrolide (preferably azithromycin) in conjunction with 3-12 weeks of intravenous amikacin and one or more of the following: intravenous ticarcillin, imipenem or cefoxitin, guided but not dictated by drug susceptibility testing. The duration of intensive phase therapy should be determined by the severity of infection, the response to treatment and the tolerability of the regimen.

The continuation phase should include a daily oral macrolide (preferably azithromycin) and inhalated amikacin, in conjunction with 2-3 of the following additional oral antibiotics: minocycline, clofazimine, moxifloxacin and linezolid, guided but not dictated by drug susceptibility testing.

- Individuals with *M. abscessus complex* pulmonary disease should be managed in collaboration with experts in the treatment of NTM and CF as drug intolerance and drug-related toxicity occur frequently and changes in antibiotic therapy are often required.
- Microbiology with a macrolide or other antimicrobial should never be used in the treatment of *M. abscessus complex* pulmonary disease.
- The same antibiotic regimen should be used for treatment of all species within the *M. avium* complex.
- Clarithromycin-sensitive *M. avium* complex pulmonary disease should be treated with a daily oral antibiotic regimen containing a macrolide (preferably azithromycin), rifampin and ethambutol.
- Clarithromycin-resistant *M. avium* complex pulmonary disease should be managed in collaboration with experts in the treatment of NTM and CF.
- Individuals with CF receiving NTM treatment should have expectorated or induced sputum samples sent for NTM culture every 4-8 weeks throughout the initial course of treatment to assess the microbiological response.
- A schedule for detecting drug toxicity (including hearing loss, visual loss, renal impairment and liver function test abnormalities) should be set in place at the time of NTM treatment initiation and implemented throughout treatment based on the specific drugs prescribed.

Transplantation

- The presence of current or previous respiratory tract samples positive for NTM should not preclude individuals being considered for lung transplantation.
- Individuals with CF who have NTM pulmonary disease and are being evaluated for transplantation should commence treatment prior to transplant listing.
- Individuals with CF receiving NTM treatment with sequential negative cultures may be eligible for transplant listing.
- Individuals with CF who have completed treatment for NTM pulmonary disease with apparent eradication of the organism may be eligible for transplant listing.

Drug Intolerance

- Other CF pathogens and co-morbidities should be considered as potential contributors to a patient’s symptoms and radiological features when determining the clinical significance of NTM positive cultures.

1. Other CF pathogens and co-morbidities should be considered as potential contributors to a patient’s symptoms and radiological features when determining the clinical significance of NTM positive cultures.

2. NTM treatment should be considered for individuals with CF who have ATS/IDSA defined NTM pulmonary disease.

3. Treatment of *M. abscessus complex* pulmonary disease should involve an intensive phase followed by a continuation phase.

4. The intensive phase should include a daily oral macrolide (preferably azithromycin) in conjunction with 3-12 weeks of intravenous amikacin and one or more of the following: intravenous ticarcillin, imipenem or cefoxitin, guided but not dictated by drug susceptibility testing. The duration of intensive phase therapy should be determined by the severity of infection, the response to treatment and the tolerability of the regimen.

5. The continuation phase should include a daily oral macrolide (preferably azithromycin) and inhalated amikacin, in conjunction with 2-3 of the following additional oral antibiotics: minocycline, clofazimine, moxifloxacin and linezolid, guided but not dictated by drug susceptibility testing.

6. Individuals with *M. abscessus complex* pulmonary disease should be managed in collaboration with experts in the treatment of NTM and CF as drug intolerance and drug-related toxicity occur frequently and changes in antibiotic therapy are often required.

7. Microbiology with a macrolide or other antimicrobial should never be used in the treatment of *M. abscessus complex* pulmonary disease.

8. The same antibiotic regimen should be used for treatment of all species within the *M. avium* complex.

9. Clarithromycin-sensitive *M. avium* complex pulmonary disease should be treated with a daily oral antibiotic regimen containing a macrolide (preferably azithromycin), rifampin and ethambutol.

10. Clarithromycin-resistant *M. avium* complex pulmonary disease should be managed in collaboration with experts in the treatment of NTM and CF.

11. Individuals with CF receiving NTM treatment should have expectorated or induced sputum samples sent for NTM culture every 4-8 weeks throughout the initial course of treatment to assess the microbiological response.

12. A schedule for detecting drug toxicity (including hearing loss, visual loss, renal impairment and liver function test abnormalities) should be set in place at the time of NTM treatment initiation and implemented throughout treatment based on the specific drugs prescribed.

Figure 1  Cystic Fibrosis Foundation and European Cystic Fibrosis Society recommendations on non-tuberculous mycobacteria (NTM) management in cystic fibrosis (CF).

microbiology, treatment and transplantation. Each subgroup developed topic-specific questions using the PICO format (population, intervention, comparison, outcome). Questions were reviewed and approved by the entire committee before systematic literature searches were conducted.

The members of each subgroup used the PICO questions to guide literature searches in PubMed. Searches were limited to English language and the period 1984–2013. Subgroup members also searched for topic relevant guidelines through searches of the ATS website, the IDSA website, the Clinical Laboratory Standards Institute website and the UK CF Trust website.

After reviewing relevant literature and existing guidelines, subgroup members drafted recommendation statements. In October 2012, a second meeting was convened, and subgroups finalised draft recommendation statements. The committee also voted to set 80% agreement of all 19 members as the threshold for acceptance of a recommendation statement.

Each subgroup submitted final draft questions for entry into an electronic survey tool (Survey Monkey) for the purposes of anonymous voting and comment by all members. A project coordinator administered the survey, and committee members were asked to rate each statement on a scale of 0 (completely disagree) to 9 (completely agree), with 80% or between 7 and 9 being considered ‘good’ agreement. Space for entering free text was also provided after each statement to allow members to cite literature in support of their opinions or suggested revisions. All committee members were required to vote on each statement regardless of their role or expertise. Multiple rounds of voting and revisions to the statements were conducted, and for each round committee members were requested to complete their voting within 3 weeks. The committee chairs reviewed the results from each round and updated the statements based on comments entered by respondents for subsequent rounds.

A draft of the recommendations was presented at the 2013 North American Cystic Fibrosis Conference and the ECFS Meeting. In addition, the committee solicited feedback from the CF communities in the USA and Europe, which included physicians, nurses, physical and respiratory therapists, parents and individuals with CF. Comments collected from this process were considered by the committee in the development of the final recommendation statements.

RESULTS

Three rounds of voting were conducted to achieve 80% consensus for each statement. Fifty-three statements were included in the first round of voting and 50 statements in the second and third rounds. Final statements are shown in figure 1.

DISCUSSION

The management of individuals with CF infected with NTM is extremely challenging. The limited amounts of published research and clinical trial data provide inadequate evidence to base management decisions on how best to screen, diagnose, detect and treat NTM-PD. As a response to this urgent clinical need, the CF Foundation and ECFS formed a committee of clinicians, scientists and infectious disease experts to develop recommendations to guide and assist clinicians in the management of NTM-PD in individuals with CF. The committee believe these recommendations should serve as a benchmark for current medical care while providing a framework to inform the development of clinical, translation and basic research studies to generate robust evidence to base future iterations of these management guidelines leading to better outcomes for individuals with CF infected with NTM.

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Competing interests

None declared.

Patient consent

Obtained.

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