SUMMARY OF RECOMMENDATIONS AND GOOD PRACTICE POINTS

Asthma

**Recommendations**

- Oral macrolide therapy could be considered to reduce exacerbation frequency in adults (50–70 years), with ongoing symptoms despite >80% adherence to high-dose inhaled steroids (>800 μg/day) and at least one exacerbation requiring oral steroids in the past year. This recommendation reflects the population within the AMAZES RCT which represents the highest quality evidence of macrolide therapy leading to a significant reduction in exacerbations. (Conditional)

- Treatment with azithromycin should be considered for a minimum of 6–12 months to assess evidence of efficacy in reducing exacerbations. (Conditional)

- Oral macrolide therapy should not be offered as a way to reduce oral steroid dose; in some individuals, this may result as a consequence of a reduction in exacerbations or symptoms. (Strong)

**Good practice points**

- Optimisation of other asthma therapies including establishing good adherence to inhaled therapies should be performed before considering a trial of oral macrolide therapy.

- Referral to a respiratory specialist or specialist asthma service should be considered prior to initiation of macrolide therapy aimed at reducing exacerbation frequency.

- For safety purposes, an ECG should be performed prior to initiation of macrolide therapy to assess QTc interval. If QTc is >450 ms for men and >470 ms for women, this is considered a contraindication to initiating macrolide therapy. Baseline liver function tests should also be measured.

- Patients should be counselled about potential adverse effects before starting therapy including gastrointestinal upset, hearing and balance disturbance, cardiac effects and microbiological resistance.

- Microbiological screening of sputum before and during macrolide therapy may be clinically helpful in patients who are able to expectorate sputum. This would allow monitoring for development of resistance and detect changes in microbial growth to direct appropriate antibiotic therapy if required. However, the resource implications of this approach have not been assessed.

- If oral macrolide therapy is considered, justification for ongoing treatment should be guided by clinical response based on specific outcome measures including exacerbation frequency, symptoms and quality of life assessed at baseline.

- A risk/benefit profile should be considered in each individual if significant side effects from oral macrolide therapy develop. If gastrointestinal side effects occur at the higher dose of azithromycin (500 mg thrice weekly), a dose reduction to azithromycin 250 mg thrice weekly could be considered if macrolide therapy has been of clinical benefit.

- Liver function tests should be checked 1 month after starting treatment and then every 6 months. An ECG should be performed 1 month after starting treatment to check for new QTc prolongation. If present, treatment should be stopped.

- Symptom improvement with macrolide treatment may be minimal and not consistent across all people with asthma. If macrolide therapy is considered for symptom reduction, this should be for a defined period (6–12 months) and stopped if no symptomatic improvement is seen. Use of a validated symptom score, such as the ACQ, may be useful to help make this assessment less subjective.

- If the desired clinical outcome is achieved, the possibility of breaks in therapy may be considered to reduce treatment burden for patients. It is unclear whether this may also reduce antimicrobial resistance rates.

Please see quick reference guide in online supplementary file 1.

**Bronchiectasis**

**Recommendations**

- Long-term macrolide treatment could be offered to reduce exacerbations in those with high exacerbation rates (ie, 3 or more per year). (Strong)

- The dosing regimens with the greatest supportive evidence, when using macrolides to reduce exacerbation rates, are azithromycin 500 mg three times a week, azithromycin 250 mg daily and erythromycin ethylsuccinate 400 mg twice a day. A starting dose of azithromycin 250 mg three times a week could be used to...
minimise side effect risk with subsequent titration according to clinical response. (Conditional)

- When using macrolides to reduce exacerbation rates, therapy should be offered for a minimum of 6 months. (Strong)
- Macrolides can be considered with the aim of improving quality of life but may require a long period of therapy (eg, 1 year) for significant effects. (Conditional)

**Good practice points**

✓ Therapies should be optimised in accordance with BTS Bronchiectasis Guidelines before considering long-term macrolide therapy (eg, airway clearance techniques and attendance at pulmonary rehabilitation courses).
✓ Macrolides should only be started following discussion and shared decision-making between the patient and a respiratory specialist.
✓ For safety purposes, an ECG should be performed prior to initiation of macrolide therapy to assess QTc interval. If QTc is >450 ms for men and >470 ms for women, this is considered a contraindication to initiating macrolide therapy. Baseline liver function tests should also be measured.
✓ Patients should be counselled about potential adverse effects before starting therapy including gastrointestinal upset, hearing and balance disturbance, cardiac effects and microbiological resistance. Microbiological assessment of sputum should be performed before therapy, including investigation for NTM. Macrolide monotherapy should be avoided if an NTM is identified. When evaluating for NTM infection, macrolides should not be used for 2 weeks before microbiological testing.
✓ Accurate assessment of baseline exacerbation rate should be determined before starting long-term macrolides for bronchiectasis.
✓ Liver function tests should be checked 1 month after starting treatment and then every 6 months. An ECG should be performed 1 month after starting treatment to check for new QTc prolongation. If present, treatment should be stopped.
✓ Subsequent follow-up at 6 months and 12 months should determine whether benefit is being derived from therapy. If there is no benefit, treatment should be stopped.
✓ Even if benefit is seen, consideration should be given to stopping treatment for a period each year, for example, over the summer. Such a drug holiday may help with reducing the development of resistance while maintaining efficacy because the vicious cycle has been broken.

See quick reference guide in online supplementary file 1.

**Chronic obstructive pulmonary disease**

**Recommendations**

- Long-term macrolide therapy could be considered for patients with COPD with more than three acute exacerbations requiring steroid therapy and at least one exacerbation requiring hospital admission per year to reduce exacerbation rate. (Conditional)
- Long-term macrolide therapy could be considered for a minimum of 6 months and up to 12 months to assess the impact on exacerbation rate. (Conditional)

**Good practice points**

✓ Non-pharmacological and pharmacological therapies should be optimised prior to considering long-term macrolide therapy. This includes smoking cessation, optimised inhaler technique, optimised self-management care plan, airway clearance techniques and attendance at pulmonary rehabilitation courses.
✓ Macrolides should only be started following discussion and shared decision-making between the patient and a respiratory specialist.
✓ For safety purposes, an ECG should be performed prior to initiation of macrolide therapy to assess QTc interval. If QTc is >450 ms for men and >470 ms for women, this is considered a contraindication to initiating macrolide therapy. Baseline liver function tests should also be measured.
✓ Patients should be counselled about potential adverse effects before starting therapy including gastrointestinal upset, hearing and balance disturbance, cardiac effects and microbiological resistance.
✓ Microbiological assessment of sputum should be performed before therapy, including investigation for NTM. Macrolide monotherapy should be avoided if an NTM is identified. Repeat assessments are recommended with clinical decline or during exacerbations to monitor resistance patterns.
✓ Accurate assessment of baseline exacerbation rate should be determined before starting long-term macrolides for patients with COPD and a CT scan should be considered to exclude a possible diagnosis of bronchiectasis.
✓ A risk:benefit profile should be considered in each individual if significant side effects from oral macrolide therapy develop. If gastrointestinal side effects occur at the higher dose of azithromycin (500 mg thrice weekly), a dose reduction to azithromycin 250 mg thrice weekly could be considered if macrolide therapy has been of clinical benefit.
✓ Liver function tests should be checked 1 month after starting treatment and then every 6 months. An ECG should be performed 1 month after starting treatment to check for new QTc prolongation. If present, treatment should be stopped.
✓ Subsequent follow-up at 6 and 12 months should determine whether benefit is being derived from therapy by using objective measures such as the exacerbation rate, CAT score or Quality of Life as measured by a validated assessment tool such as SGRQ. If there is no benefit treatment should be stopped.
✓ It is not necessary to stop prophylactic azithromycin during an acute exacerbation of COPD unless another antibiotic with potential to affect the QT interval has also been prescribed.

See quick reference guide in online supplementary file 1.

**Bronchiolitis obliterans (including post transplantation)**

**Recommendations**

- Low-dose, long-term azithromycin (250 mg thrice weekly) could be considered to prevent the occurrence of BOS post lung transplantation. (Conditional)
- Low-dose azithromycin (250 mg alternate days for a trial period of 3 months) could be considered to treat BOS occurring in lung transplant recipients. (Conditional)

**Use of macrolides in other respiratory conditions**

**Cough**

**Recommendations**

- Long-term macrolide antibiotics should not be used to manage patients with unexplained chronic cough. (Conditional)

**Organising pneumonia**

There is insufficient evidence to make a recommendation.
Safety issues
Gastrointestinal effects

Good practice points

✓ Prior to initiating low-dose macrolide therapy, patients should be warned of the possibility of gastrointestinal side effects.
✓ Gastrointestinal side effects may be ameliorated by dose reduction although this may also reduce clinical efficacy.
✓ Clinicians should carefully consider the risk-to-benefit balance when considering therapy for those with pre-existing gastrointestinal symptomatology.

Cardiac effects

Good practice points

✓ Prior to initiating low-dose macrolide therapy, patients should be asked if they have a history of heart disease, previous low serum potassium measurements, a slow pulse rate, a family history of sudden death or known prolonged QT interval. Patients with such a history should not receive low-dose macrolide therapy without careful consideration and counselling of the increased risk of adverse cardiac effects.
✓ Prior to initiating low-dose macrolide therapy, a drug history looking for agents that might prolong the QTc interval should be sought (see online supplementary appendices 3 and 4). Patients taking such agents should not receive low-dose macrolide therapy.
✓ Prior to initiating low-dose macrolide therapy, an ECG should be performed to exclude a prolonged QTc interval defined as >450 ms for men and >470 ms for women (see methodology in online supplementary appendices 3 and 4). Patients with a prolonged QTc interval should not receive low-dose macrolides.
✓ One month after initiating low-dose macrolide therapy, a second ECG should be performed to exclude the development of a prolonged QTc interval. Patients who develop a prolonged QTc interval on low-dose macrolides should stop the macrolide.
✓ If any new drug that could potentially prolong QTc time is started or if dose increases are made, repeat ECG assessment.

Otoxicity

Good practice point

✓ Prior to initiating low-dose macrolide therapy, patients should be asked if they have a history of hearing or balance difficulties. Such patients should be made aware of the potential for a further, almost always reversible, deterioration.

Other side effects

Good practice points

✓ Prior to initiating low-dose macrolide therapy, baseline liver function tests (LFTs) should be checked.
✓ LFTs should be checked after 1 month of treatment and then every 6 months thereafter for the duration of therapy.

Antimicrobial resistance

Good practice points

✓ The risks associated with increasing antimicrobial resistance should be discussed with patients prior to starting low-dose macrolide therapy. Patients should understand the risk that there may not be an effective antibiotic for them, or someone else, when needed in the future.
✓ Prior to initiating low-dose macrolide monotherapy, patients should be asked if they have a history of previous or current NTM infection or disease. Current NTM infection should be managed with reference to BTS guidance and precludes low-dose macrolide monotherapy. Successfully treated NTM disease should not preclude low-dose macrolide monotherapy.
✓ If there is any clinical suspicion of possible NTM disease, patients should be screened via examination of sputum samples prior to starting therapy. If positive for recognised potential pathogenic species, low-dose macrolide prophylaxis is contraindicated.

SECTION 1: INTRODUCTION

Aim of the guideline
The aim of this guideline is to examine the evidence for the use of long-term, low-dose macrolide agents in the therapy of adult respiratory diseases and to develop guidance for clinicians in such use of these agents.

Intended users of the guideline and target patient populations
These guidelines will be of interest to UK-based clinicians caring for adults with respiratory disease including respiratory physicians, acute/general medicine physicians and respiratory specialist nurses. The guidelines may also be of interest to GPs, community matrons and practice nurses, hospice staff and community respiratory teams, physiotherapists, microbiologists, pathologists, pharmacists, haematologists and lung transplant teams.

Scope of the guideline
The Guideline Development Group (GDG) has examined the use of macrolides in adults (>16 years) where the duration of treatment exceeds that usually employed to treat an acute infection and the dosage is less than that usually employed to treat an acute infection. Such usage is considered to be exerting an effect through mechanisms other than a direct antibacterial one, commonly described as immunomodulatory mechanisms. We have characterised this as long-term, low-dose usage and have examined such usage (often in comparison with conventional therapy) in the following conditions:

• Asthma
• Bronchectasis
• COPD
• Bronchiolitis obliterans
• Chronic cough
• Organising pneumonia
• Diffuse panbronchiolitis.
The use of macrolides as antibacterial agents to treat respiratory infection is excluded.

Limitations of the guideline
Healthcare providers need to use clinical judgement, knowledge and expertise when deciding whether it is appropriate to apply recommendations for the management of patients. The recommendations cited here are a guide and may not be appropriate for use in all situations. The guidance provided does not override the responsibility of healthcare professionals to make decisions appropriate to the circumstances of each patient, in consultation with the patient and/or their guardian or carer.

The GDG recognises that none of the macrolide antibiotics are licensed in the UK for long-term low-dose usage as immunomodulatory agents at the time of guideline development.

Members of the GDG
The GDG was chaired by two respiratory consultants—Dr David Smith (DS) and Dr Ingrid Du Rand (IDR). The GDG had a wide membership and included colleagues from respiratory medicine, pharmacy and microbiology. A patient representative was on the group for the duration of the process. Those on the group were not required to be BTS members. Mrs Joan McCarthy was the lay representative.

A full list of members can be seen at online supplementary appendix 2.

Acknowledgements
The GDG is grateful to the Standards of Care Committee for assistance during the development of the guideline, and particular thanks are due to Dr Toby Capstick and Dr Mike Crooks for their input and advice in relation to certain sections of the guideline.

SECTION 2: METHODOLOGY OF GUIDELINE PRODUCTION
Establishment of guideline development group
The GDG was convened in June 2016, with the first meeting taking place in October 2016. The full GDG met three times during the development of the guideline and kept in close contact by teleconference/WebEx and email throughout the process.

Methodology
This is the first BTS guideline to use the GRADE approach as part of the process of guideline development. Previous guidelines have used the SIGN methodology. BTS has made this change reflecting common practice in guideline development internationally across all medical specialities. The advantages of the GRADE approach are described in detail in the GRADE handbook and the BTS GRADE guideline production manual. GRADE specifies an approach to framing questions, choosing outcomes of interest and rating their importance, evaluating the evidence, and incorporating evidence with considerations of values and preferences of patients and society to arrive at recommendations.

The methodology used to write the guideline adheres strictly to the criteria as set by the AGREE II collaboration, which is available online (www.agreetrust.org/resource-centre/agree-ii/). The British Thoracic Society Standards of Care Committee guideline production manual is available online (https://www.brit-thoracic.org.uk/quality-improvement/guidelines/).

Table 1 Summary of outcome measures

<table>
<thead>
<tr>
<th>Population</th>
<th>Intervention</th>
<th>Control</th>
<th>Outcomes</th>
</tr>
</thead>
</table>
| Adults with asthma               | Long-term, low-dose macrolides | Placebo | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |
| Adults with bronchiectasis       | Placebo      |         | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |
| Adults with COPD                 | Placebo      |         | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |
| Adults with bronchiolitis obliterans | Placebo  |         | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |
| Adults with chronic cough        | Placebo      |         | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |
| Adults with organising pneumonia | Placebo      |         | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |
| Adults with diffuse panbronchiolitis | Placebo |         | ► Quality-of-life measures  
► Symptom improvement/symptom score  
► Exacerbation rates  
► Hospital admission rate  
► Disease progression and changes in lung function tests  
► Mortality  
► Exercise capacity/tolerance  
► Sputum volume/colour/character and microbiological resistance/dysbiosis  
► Drug monitoring/side effects/toxicity |

Summary of key questions, outcomes and literature search
Clinical questions were formulated in the PICO (Patient, Intervention, Comparison and Outcome) format (see table 1). The key questions are summarised below.

1. Should long-term, low-dose macrolides be used in the treatment of adults with asthma?
2. Should long-term, low-dose macrolides be used in the treatment of adults with bronchiectasis?
3. Should long-term, low-dose macrolides be used in the treatment of adults with chronic obstructive pulmonary disease?
4. Should long-term, low-dose macrolides be used in the treatment of adults with bronchiolitis obliterans?
5. Should long-term, low-dose macrolides be used in the treatment of adults with chronic cough?
6. Should long-term, low-dose macrolides be used in the treatment of adults with organising pneumonia?
7. Should long-term, low-dose macrolides be used in the treatment of adults with diffuse panbronchiolitis?

The following patient-centred outcomes were identified by the group when the scope of the guideline was agreed:

- Quality-of-life measures.
- Symptom improvement/symptom score.
- Exacerbation rates.
- Hospital admission rate.
- Disease progression and changes in lung function tests.
- Mortality.
- Exercise capacity/tolerance.
- Sputum volume/colour/character and microbiological resistance/dysbiosis.
- Drug monitoring/side effects/toxicity.

The PICO framework was used to define the scope of the guideline and formed the basis of the literature search. The initial search was completed in February 2017 by York University. Systematic electronic database searches were conducted in order to identify all papers which might potentially be included in the guideline. For each question, the following databases were searched: Cochrane Database of Systematic Reviews (CDSR), Database of Abstracts of Reviews of Effects (DARE), Health Technology Assessment Database (HTA), Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE and MEDLINE In-Process, EMBASE and PubMed. The search strategy is available for review in online supplementary appendix 1.

Appraisal of the literature
The literature search was run in 2016. The search was limited to papers published in English. The search identified a total of 5225 abstracts; after initial screening, this was reduced to 1152.

Criteria formulated for initial screening of the abstracts were:

- Whether the study addressed the clinical question.
- Whether the study addressed the clinical question.
The full list of abstracts was retained and is kept in an archive. The literature search was repeated in December 2017 to capture additional published evidence while the guideline was in development prior to finalising the draft document. Additional references were included from critical appraisal of the literature and review of existing evidence base as appropriate.

Letters, conference papers and news articles were then removed—518 abstracts were rejected at this stage. IDR and DS read the remaining abstracts, marked those considered relevant to the scope of the guideline and allocated each relevant abstract to a clinical question(s). In total, 634 abstracts were allocated to clinical question(s).

The literature search was updated in late 2017 and identified 138 additional abstracts that were reviewed and 14 papers were selected to be critically appraised.

GDG members were allocated to work on individual questions in small groups.

Each abstract was read and at least two members agreed whether the paper was relevant, possibly relevant or irrelevant to the particular clinical question. Papers were excluded at the title/abstract sift if the following applied:

► If the paper did not address the clinical question and at least one of the outcomes concerned.
► If it was a case series of less than 20 patients—however, this was not an absolute cut-off. Professional judgement was applied such that some smaller case series were considered if evidence was sparse, and indeed some case reports of more than 20 patients were excluded if there was higher quality evidence available.
► If the language of the full paper was not English.

Full papers were obtained for all relevant, or possibly relevant, abstracts. Each paper was read in full and critically appraised by at least two members of the GDG to confirm relevance to the clinical question and the presence of at least one outcome of interest.

Each paper was then appraised by outcome(s), to generate a best estimate of the effect on each outcome and an index of the uncertainty associated with that estimate where possible. An evidence profile entry was completed for each outcome which included grading of the quality of the evidence. The type of evidence available for each outcome varied from systematic reviews through to case series; for each outcome, the highest quality evidence available was included. GRADEPro was used to generate the evidence profiles, published online on the BTS website where they are available for review (see online supplementary appendix 2). GRADEPro (https://gradepro.org/) is an easy-to-use, all-in-one web-based electronic guideline development tool to support guideline development using the GRADE process and methodology. In addition, it provides an electronic platform to present evidence tables, considered judgement and assessment of the evidence base.

The GRADE approach to rating the quality of evidence begins with the study design (table 2) and then, through a process of considered judgement, applies five reasons to possibly rate down the quality of evidence and three reasons to possibly rate up the quality (table 3).

The GDG reviewed each clinical question during the regular meetings and consensus was reached.

In assessing the evidence the guideline development group combined low and very low evidence into one category (Low) as the body of evidence was limited.

**Development of recommendations**

Having generated evidence profiles for each of the clinical questions the GDG as a whole then considered the importance of each of the outcomes for each clinical question and proceeded to grade the overall body of evidence for critical and important outcomes.

The GDG went on to decide on the direction and strength of recommendations considering the quality of the evidence, the balance of desirable and undesirable outcomes and the values and preferences of patients and others. GRADE specifies two categories of strength of a recommendation as shown in table 4.

Good practice points (GPPs) were developed by consensus in areas where there was no quality evidence but the GDG felt that some guidance based on the clinical experience of the GDG might be helpful to the reader. These are indicated as shown below.

| √ | Recommended best practice based on the clinical experience of the guideline development group |

From the outset, it was acknowledged that there would be little high-quality evidence for some of the clinical questions identified. In this instance, low-grade evidence was considered, along with expert opinion via consensus at the meetings.

Cost-effectiveness was not considered in detail as in-depth economic analysis of recommendations falls outside of the scope of the BTS guideline production process. However, the GDG

<table>
<thead>
<tr>
<th>Table 2 Categories of evidence</th>
<th>Characteristics</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Based on consistent results from well-performed randomised controlled trials</td>
<td>Further research is very unlikely to change the estimate of the effect</td>
</tr>
<tr>
<td>Moderate</td>
<td>Based on randomised controlled trials where there is evidence of bias or from other well-conducted study types (eg, well-executed observational studies)</td>
<td>Further research is likely to have an impact on the estimate of the effect</td>
</tr>
<tr>
<td>Low</td>
<td>Based on observational evidence or from controlled trials with several serious limitations</td>
<td>Further research is likely to have an important impact</td>
</tr>
<tr>
<td>Very low</td>
<td>Based on case studies or expert opinion</td>
<td>Estimates of effect are far from certain and more research is needed</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 3 Decreasing and increasing the grade of evidence</th>
<th>Decrease grade if</th>
<th>Increase grade if</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serious or very serious limitation to study quality</td>
<td>Magnitude of the treatment effect is very large and consistent</td>
<td></td>
</tr>
<tr>
<td>Important inconsistencies in results</td>
<td>Evidence of a large dose-response relation</td>
<td></td>
</tr>
<tr>
<td>Some or major uncertainty about directness of the evidence</td>
<td>All plausible confounders/biases would have decreased the magnitude of an apparent treatment effect</td>
<td></td>
</tr>
<tr>
<td>Imprecise or sparse data (relatively few participants and/or events)</td>
<td>High probability of reporting bias</td>
<td></td>
</tr>
</tbody>
</table>

*Each quality criterion can reduce the quality by one or, if very serious, by two levels. See BTS GRADE guideline production manual for further details (https://www.brit-thoracic.org.uk/quality-improvement/guidelines/).
Table 4  Explanation of the terminology used in BTS recommendations

<table>
<thead>
<tr>
<th>Strength</th>
<th>Benefits and risks</th>
<th>Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong. It is recommended and so ‘offer’</td>
<td>Benefits appear to outweigh the risks (or vice versa) for the majority of the target group</td>
<td>Most service users would want to or should receive this intervention</td>
</tr>
<tr>
<td>Conditional. It is suggested and so ‘consider’</td>
<td>Risks and benefits are more closely balanced, or there is more uncertainty in likely service users values and preferences</td>
<td>The service users should be supported to arrive at a decision based on their values and preferences</td>
</tr>
</tbody>
</table>

Drafting the guideline

The GDG corresponded regularly by email and meetings of the full group were also held in the period between July 2016 and early 2019. The guideline was discussed at an open session at the BTS Winter Meeting in December 2018. A revised draft guideline document was circulated to all the relevant stakeholders for consultation in December 2018 followed by a period of online consultation. The BTS SOCC reviewed the draft guideline in March 2018 and March 2019.

Review of the guideline

This guideline topic was chosen to inform practice in a growing clinical area, but also to act as a pilot for the use of GRADE methodology for all future BTS guidelines. The GDG recognise that the topic area would include a range of diseases with differing levels of evidence and thus act as a useful learning exercise for the introduction of GRADE methodology. It is not proposed to update this guideline as a distinct entity—the intention is for macrolide use to be encompassed in future disease-specific guideline updates.

Declarations of interests

BTS Declarations of Interest forms have been completed by all members for each year they were part of the GDG. Details of these forms can be obtained from BTS Head Office. “Declarations of Interests” was a standing item at each GDG meeting.

Stakeholders

Stakeholders were identified at the start of the process. All stakeholder organisations were notified when the guideline was available for public consultation and a list is published in online supplementary appendix 2.

Other available guidance

The GDG were aware of parallel workstreams within the BTS (eg, the BTS guideline for bronchiectasis in adults) and in other organisations (eg, NICE) either complete or in progress at the time this guideline began development. Some of those in progress have subsequently been published and overlap with this guideline. The majority of these guidelines are disease specific rather than focusing on a single drug group. The pool of evidence available for these guidelines is the same, but the sample drawn from the literature will differ. Therefore the methodology applied and the resources available to individual organisations are also variable. It is thus not surprising that the recommendations sometimes differ between contemporaneous documents. We have included a list of recently published guidelines and Cochrane reviews which overlap with this document.

BTS Guideline for Bronchiectasis in Adults 2019

NICE CF diagnosis and management Oct 2017
NICE Asthma diagnosis, monitoring and management Nov 2017
NICE Chronic obstructive pulmonary disease (acute exacerbation): antimicrobial prescribing Dec 2018
NICE Bronchiectasis (acute exacerbation): antimicrobial prescribing Dec 2018
NICE COPD guideline 2018
GOLD guideline COPD 2019
ERS/ATS COPD Exacerbations 2017
Cochrane Prophylactic antibiotic therapy for COPD 2018
Cochrane Prolonged antibiotics for non-cystic fibrosis bronchiectasis in children and adults 2017
Cochrane Antibiotics for exacerbations of asthma 2018
Cochrane Macrolide antibiotics for bronchiectasis 2018
Cochrane Macrolides for chronic asthma 2015
GINA 2018 Global Strategy for Asthma Management and Prevention

Within each of sections 4, 5 and 6, a summary of other guideline recommendations for that disease has been included.

SECTION 3: INTRODUCTION TO THE USE OF MACROLIDES IN RESPIRATORY CARE

The term macrolide describes compounds with a macrocyclic lactone ring of 12 or more elements. Within this definition are a variety of drugs including antibiotics, antifungals, prokinetic agents and immunosuppressants. The most commonly used macrolide antibiotics have 14-membered or 15-membered lactone rings and include the first macrolide, erythromycin, which was launched commercially in 1952. Further 14-membered macrolides include clarithromycin and roxithromycin; the 15-membered azithromycin became available in the 1980s. Macrolide antibiotics are active orally, have excellent tissue penetration and antimicrobial activity against many Gram-positive bacteria, some Gram-negative organisms and some ‘atypical’ respiratory pathogens such as legionella and mycoplasma together with some mycobacterial species. This makes them a popular choice for respiratory tract infections.

Early studies with macrolide antibiotics in asthma in short antimicrobial courses and in longer, lower-dose regimens were the first to suggest a possible non-antimicrobial mode of action, possibly as a steroid-sparing agent.

Diffuse panbronchiolitis (DPB) was recognised as a distinct clinical entity in Japan in 1969 having a poor prognosis, 25% mortality at 10 years without Pseudomonas aeruginosa infection or >75% with P aeruginosa. The first report of a dramatic response to low-dose, long-term treatment with erythromycin appeared in 1987 and led to further confirmatory studies of macrolide efficacy in DPB. This acted as a catalyst for the subsequent exploration of the non-antimicrobial effects of macrolides.
The mechanisms behind the immunomodulatory effects of macrolides have been widely researched. They include alterations in airway secretions through effect on ion transport and mucus production. Changes in the inflammatory process occur through effects on cytokine production, adhesion molecule expression and the functions of inflammatory cells, airway epithelial cells and other cells. Sublethal effects on bacteria include disruption of biofilms, interference with quorum sensing and reduced bacterial adherence, toxin production and mobility.12

The recognition of these immunomodulatory properties has led to the investigation of potential benefit from low-dose, long-term macrolides in a number of other chronic inflammatory airway diseases including cystic fibrosis, bronchiectasis, asthma, Chlamydiae, rhinosinusitis and bronchiolitis obliterans. In some of these conditions (eg, cystic fibrosis), the evidence base supporting the use of macrolides is large but in others (eg, COP) it is small.

Alongside this increase in the longer-term use of macrolides as immunomodulatory agents, concerns regarding the safety of these drugs used in this way have appeared.23 Gastrointestinal side effects are unsurprisingly common but rarely serious; however, ototoxicity and effects on myocardial tissue are potentially more harmful. Alterations in the microbiome of individuals and populations and the rapid emergence and persistence of macrolide-resistant organisms have also prompted discussion.24 It should also be noted that the rising rates of antimicrobial-resistant pathogens is of global concern. Any consideration of the benefits of an individual patient receiving an antibiotic needs to be carefully balanced against the risk of increased resistance—both for that individual patient but also for other individuals, both now and in the future.

It was against this background of expanding use of macrolides in inflammatory respiratory diseases that BTS decided to commission a guideline aimed at supporting best practice in a developing arena.

SECTION 4: ASTHMA

Introduction

The majority of asthma treatments aim to reduce airway inflammation with subsequent symptomatic improvement. Pathologically, asthma is characterised by airway inflammation and bronchial hyper-responsiveness. Airway inflammation can be predominantly neutrophilic or eosinophilic in nature.15 Despite an increase in therapeutic options and clear guidance on stepwise management, asthma symptoms can be difficult to manage and exacerbations remain a predominant feature for many. Adherence to treatment can be challenging. Newer therapeutic options, including biologics, can be expensive, require frequent monitoring and are limited to specific clinical phenotypes.15 25

There has been significant interest in the potential role of oral macrolide therapy as an option in asthma. It is relatively inexpensive, easy to adhere to and has a physiologically relevant mechanism of action which may reduce airway inflammation in asthma. There is also evidence suggesting increasing asthma severity may be linked to chronic colonisation with Chlamydiae and Mycoplasma pneumoniae. It has therefore been postulated that benefit from macrolide therapy may also be through its antibacterial effects on these and other lung organisms.76

As a consequence, there have been a number of studies over several decades examining the response to macrolide therapy in asthma which vary considerably in design and outcome measures. The first study examining the role of macrolides in asthma was their use as a steroid-sparing agent in 1970.17 Studies since have examined their role in reducing airway inflammation and bronchial hyper-responsiveness, impact on symptoms and quality of life and most recently on their ability to reduce exacerbation frequency.

Evidence base

Examining the evidence base for macrolide therapy in asthma is particularly challenging because of the significant heterogeneity in study designs, outcome measures and study populations, as well as the heterogeneity of the asthma population itself. A Cochrane review of macrolide therapy in asthma identified significant publication bias, poor study design and lack of consistency in outcome measures as the key barriers to making recommendations based on the available evidence.14 Considering the size of the clinical population and burden of disease in this population,15 there is a relative paucity of high-quality evidence regarding macrolide use in asthma. There are relatively few studies with small study populations which are often underpowered to detect appropriate changes in the outcome measures used. The study populations are also varied limiting the applicability of their results to the wider asthma population and the ability to determine where macrolide therapy should sit within an asthma treatment regime.

Despite this, across multiple outcome measures statistical significance was achieved in many studies, but the magnitude of the changes seen were very small. The clinical significance of these changes varies from unknown to unlikely to be of any clinical benefit. This is reflected in the recommendations made in this guideline. No evidence of the impact of macrolide therapy on mortality, exercise capacity, disease progression or sputum production in people with asthma was found therefore no recommendations in regard to these outcomes can be made in this guideline. In all cases, macrolides should be considered after current therapy has been optimised and adherence established.

Other recent guidance

Recommendations on macrolide therapy are not included in current NICE/BTS/SIGN asthma guidelines.4 25 NICE guidance suggests referral to an asthma specialist and consideration of the addition of a trial of ‘an additional drug’ (which could include a macrolide) if asthma is uncontrolled in adults (aged 17 and over) on a moderate maintenance ICS dose with a LABA (either as MART or a fixed-dose regimen).

Quality of life (QOL)

QOL was the most consistently assessed outcome measure in studies of macrolide therapy in asthma. All but one study27 used the well-validated Asthma Quality of Life Questionnaire (AQLQ) score. This score was first validated in 1992 and has a minimal important difference (MID) of 0.5.28 This is critical to appraising this body of evidence, as in similarity with the symptomatic improvements seen, after treatment with macrolides there is a consistent small but statistically insignificant (p>0.05) improvement in QOL seen across the majority of studies.14 27 28–36

In two studies comparing macrolide therapy with placebo,33 34 this change reached the MID of the AQLQ. The first used 8 weeks’ treatment with clarithromycin34 and the second was the open-label treatment arm of a study of 48 weeks’ treatment with azithromycin.34

It is therefore clear that treatment with macrolides may result in measurable improvements in QOL for people with asthma, but the clinical impact of these changes remains unknown and may be very small.
Symptoms

Thirteen studies used symptom scoring as an outcome in response to macrolide therapy.\textsuperscript{14, 27, 29–31, 34–41} There was considerable variation in the scoring systems used, from the validated Asthma Control score (ACQ) to simplified unvalidated scoring systems, created specifically for an individual study. Overall, 10 of the studies demonstrated a reduction in symptoms. Three studies\textsuperscript{29, 31, 33} demonstrated an increase in symptoms, of which only one used a validated and reproducible symptom score. Although the majority of studies demonstrated an improvement, the actual changes were minimal, unlikely to be of clinical significance and reached statistical significance in only three cases.\textsuperscript{37, 38, 41} None of the studies using the ACQ score demonstrated a change which would meet the ACQ minimal important difference (MID) (0.5).\textsuperscript{42}

Further work is needed to see if these improvements may be of greater significance over more prolonged treatment periods, and whether symptomatic improvement is related to the impact of macrolide therapy on exacerbation rate.

Exacerbations

Exacerbations of asthma are defined as an increase in symptoms, accompanied by a reduction in peak flow rate (PEFR), of graded severity dependent on the degree of clinical and PEFR deterioration.\textsuperscript{4, 25} Exacerbation rate is the definitive outcome measure in regard to the clinical efficacy of macrolides in other respiratory diseases, and the major purported driver for their use in cystic fibrosis, bronchiectasis and COPD.\textsuperscript{14} Two studies\textsuperscript{30, 40} have been specifically designed to address this question in asthma. In other studies, exacerbations have been measured as a secondary outcome, with differing definitions between studies and insufficient statistical power to detect a reduction in exacerbation frequency. A meta-analysis of these studies showed exacerbations requiring steroids in 19.6% (31/158) of those treated with macrolides and 24.2% (32/132) of those not receiving macrolide treatment.\textsuperscript{14} However, this reduction was not sufficient to reach statistical significance (p>0.05).

Of the two studies\textsuperscript{30, 40} specifically designed to determine a reduction in exacerbation frequency in response to azithromycin, the AMAZES trial demonstrated a significant reduction in exacerbations (p<0.0001),\textsuperscript{40} but the AZIZAST trial did not (p=0.68).\textsuperscript{30} This may reflect the duration of treatment (AMAZES=48 weeks; AZIZAST=6 months), dose effect (AMAZES=500 mg thrice weekly; AZIZAST=250 mg thrice weekly) or differing study populations. AMAZES recruited a larger population (n=420) on high-dose inhaled steroids with an average of one exacerbation in the previous year, whereas the AZIZAST population was smaller (n=109) with higher inhaled steroid doses and two exacerbations in the previous year (see table 5). Both studies also had slightly differing definitions of an exacerbation. These studies are the highest quality evidence found for the purposes of this guideline and were suitably powered to detect a change in exacerbation frequency.

The AZIZAST study demonstrated no significant difference between exacerbation rates with 48% (26/54) of the placebo group and 47% (26/55) of the azithromycin group experiencing an exacerbation (relative risk 0.98, 95% CI 0.68 to 1.43, p=0.68).\textsuperscript{30} In AMAZES, 61% (127/203) of the placebo group experienced an exacerbation compared with 44% (94/213) of the azithromycin group giving an incidence rate ratio of 0.59 (95% CI 0.47 to 0.74, p<0.0001)\textsuperscript{40} (see table 5).

Both studies assessed whether the predominant type of lung inflammation impacted the effectiveness of macrolide therapy. In a predefined subgroup analysis of the AZIZAST population, those with non-eosinophilic asthma showed a significant reduction in exacerbations in response to azithromycin treatment, with exacerbations occurring in 33% (9/27) of the azithromycin group compared with 62% (18/29) of the placebo group (relative risk 0.54, 95% CI 0.29 to 0.98, p=0.037).\textsuperscript{30} In AMAZES, both those with eosinophilic asthma and non-eosinophilic asthma demonstrated a reduction in exacerbations, with a slightly greater reduction seen in the eosinophilic group. The incidence rate ratio in the eosinophilic group was 0.52 (95% CI 0.29 to 0.94, p=0.030) and 0.66 (95% CI 0.47 to 0.93, p=0.019) in the non-eosinophilic group.\textsuperscript{40} This difference in outcomes may be partly explained by the differing definition of phenotypes between the two studies. In AZIZAST, non-eosinophilic asthma was defined by a fraction of exhaled nitric oxide (FeNO) lower than the upper limit of normal and a blood eosinophilia ≤200/μL.\textsuperscript{30} In AMAZES, non-eosinophilic asthma was defined by a baseline sputum eosinophil count <3% or blood eosinophil count <300/μL if sputum unavailable.\textsuperscript{40}

Overall, this evidence suggests that macrolide therapy may vary in its effectiveness between differing asthma phenotypes and that clinical benefit may be determined by more than the predominant type of inflammation. Further work is needed to investigate the impact of macrolides on exacerbation rate in differing asthma phenotypes and populations. However, it

<table>
<thead>
<tr>
<th>Table 5</th>
<th>Comparison between the AZIZAST and AMAZES trials</th>
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<td><strong>Population</strong></td>
<td><strong>Intervention</strong></td>
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<tr>
<td>Brusselle et al (AZIZAST)\textsuperscript{30}</td>
<td>18–75 years&lt;br&gt;GOLD step 4/5&lt;br&gt;&gt;1000 μg fluticasone or equivalent&lt;br&gt;2 exacerbations requiring OCS in past 12/12&lt;br&gt;FeNO in normal limits</td>
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<tr>
<td>Gibson et al (AMAZES)\textsuperscript{40}</td>
<td>18 years or older&lt;br&gt;Diagnosis of asthma (post-bronchodilator reversibility &gt;12%&lt;n&gt;airway hyper-responsiveness or PEFR variability &gt;12%)&lt;n&gt;&lt;br&gt;Partial loss of asthma control—ACQ≥0.75</td>
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ACQ, Asthma Control Questionaire; GOLD, Global Initiative for Chronic Obstructive Lung Disease; OCS, Oral corticosteroids; PEFR, Peak Expiratory Flow Rate.
improvement. This is illustrated by the results of studies examining the impact of oral macrolide therapy since the 1990s.

Steroid dose reduction
The original studies of macrolide use in asthma had the aim of reducing steroid dosage, but this has not been the stated outcome of a study using macrolide therapy since the 1990s. With the advent of newer biologic therapies and the focus on clinical phenotyping, there has been a move away from using therapies as steroid-sparing agents, towards a stratified approach to asthma therapy.

The three studies using steroid reduction as an outcome all used troleandomycin for between 2 and 52 weeks. All showed a reduction in steroid dosage, which has been confirmed in two Cochrane Reviews. However, these studies also included paediatric populations. Troleandomycin is also not widely used and is not available in the UK.

Lung function/PEFR
FEV₁ remains a key outcome measure in trials of airflow diseases. However, in asthma, FEV₁ can show considerable variation and is used as a secondary outcome measure in many studies. Peak flow was used frequently as a primary and secondary outcome, but with considerable variation in timing, rigour of performance and units of measurement.

Overall FEV₁ improved in response to oral macrolide therapy, but to a small degree which may not result in any clinical improvement. This is illustrated by the results of four meta-analyses which individually compared FEV₁ responses across 15 studies using a range of macrolides. The Cochrane meta-analysis demonstrated a 0.08 L (95% CI 0.02 to 0.14 (p=0.0097)) improvement in FEV₁ (47); the Tong et al meta-analysis showed a 0.11 L (95% CI 0.06 to 0.16 (p<0.001)) improvement; and Reiter et al showed no effect on FEV₁ (standardised mean difference (SMD) 0.05 (95% CI −0.14 to 0.25 (p=0.60)). FEV₁ (% predicted) was only analysed in the Tong et al meta-analysis, showing a SMD of 0.27% (95% CI 0.05 to 0.59 (p=0.10)). The fourth meta-analysis was confined to studies examining troleandomycin, showing a 0.06 L (95% CI −0.8 to 0.92 (p=0.88)) change in FEV₁.

Morning PEFR was the most commonly measured parameter across studies. A positive change in morning PEFR was seen in several studies, of a relatively smaller magnitude than that seen in FEV₁. The Cochrane and Tong et al meta-analyses showed a 2.22 L/min (95% CI 9.73 to 14.17 (p=0.72)) improvement and a SMD 0.25 (95% CI 0.1 to 0.39 (p=0.001)) in morning PEFR, respectively. A third meta-analysis by Reiter et al also showed a 6.7 L/min (95% CI 1.35, 12.06 (p=0.014)) improvement, but this analysis included paediatric study populations. The change in evening PEFR is more mixed across studies, with some showing a reduction rather than improvement, although with a smaller evidence base.

With the known variability of PEFR in asthma, again it is difficult to judge the clinical importance of these changes.

Inflammation
Studies examining the impact of oral macrolide therapy on inflammation in asthma have used a variety of outcome measures in both blood and sputum. In keeping with common clinical phenotypes, the impact of macrolides on both neutrophil and eosinophil inflammation has been assessed. Two meta-analyses examined the effect of macrolide therapy on sputum and blood eosinophil counts measured in three separate RCTs after treatment with roxithromycin, clarithromycin and azithromycin. Consistent reductions in blood eosinophil counts were shown (mean difference (MD) −33.5 (95% CI −30.9 to −36.11)×10⁹/mL (p<0.00001)), but only the study examining clarithromycin (MD −74×10⁹/mL) demonstrated a reduction in sputum eosinophils.

Five studies examined the impact of roxithromycin, clarithromycin and azithromycin on sputum neutrophils. One study demonstrated an increase in sputum neutrophils (MD 19.2 (95% CI −24.2 to 62.6), p>0.05) in response to azithromycin, but pooled results of the other studies, as part of a meta-analysis, demonstrated an overall reduction of −0.25% (95% CI −0.62 to 0.13, p=0.20).

Emerging biomarkers in the form of matrix metalloproteinase (MMP-9), neutrophil elastase and eosinophil cationic protein (ECP) have also been examined. Clarithromycin resulted in a reduction in sputum MMP-9 and neutrophil elastase after 8 weeks of treatment. Serum and sputum ECP were both reduced after treatment with clarithromycin and roxithromycin. FeNO is increasingly used as a biomarker of inflammatory activity in asthma and a measure of response to therapy. Three randomised trials used FeNO as an outcome measure all demonstrating a small and non-significant reduction in response to treatment with 12 weeks of azithromycin (−1.94 (95% CI −5.97 to 2.10) ppb (p=0.34)) and 26 weeks of azithromycin (MD −1.6 ppb (p=0.52)) and 16 weeks of clarithromycin (−4.6 (SE ±4.2) ppb (p=0.5)), respectively.

While there is considerable variation in study populations, treatment regimens and outcome measures, the overall picture is that macrolides do reduce airway inflammation in asthma. Further work is needed to clarify if this translates into clinically relevant improvements for patients and to determine which inflammatory biomarker may be of most use in determining response to treatment.

Bronchial hyper-responsiveness
Bronchial hyper-responsiveness has been used as an outcome measure in a number of studies of macrolide therapy in asthma to provide proof of physiological effect and potential explanation for improvement in other clinical outcome measures. The studies confirm that macrolide therapy does result in improvements in bronchial hyper-responsiveness measured by methacholine challenge test (PC20/PD20). Two studies were specifically designed to assess bronchial hyper-responsiveness as a primary outcome. When pooled, they demonstrated a 1.99 (95% CI 0.46,3.52 (p=0.011)) SMD improvement in methacholine response. A second meta-analysis of five studies demonstrated an improvement of 0.9 (95% CI 0.5,1.75 (p=0.04)). Other studies demonstrated varying magnitudes in the degree of improvement seen, likely related to differing macrolides, treatment dosages, duration of treatment, study populations and measurement differences.

While these findings support a physiological mechanism of action of macrolides through a reduction in airway inflammation and bronchial hyper-responsiveness, there seems to be a lack of clear correlation between the degree of improvement seen and any corresponding improvement in clinical outcomes. This requires further investigation in future trials.

Microbiology
Five studies specifically reported the impact of oral macrolide therapy on microbiological outcomes. Black et al conducted experiments...
an RCT of 6 weeks of roxithromycin versus placebo in 232 people with asthma and raised IgG/IgA antibody titres to Chlamydia pneumoniae.29 This demonstrated a reduction in IgG antibodies, but not IgA in response to treatment; but with no clinically significant changes in other outcome measures. Hahn et al demonstrated that high IgA but not IgG antibodies to C. pneumoniae were significantly associated with more severe asthma symptoms at the end of follow-up.33 Azithromycin resulted in improvements in IgA antibodies regardless of whether IgA antibody levels were high or low at the start of treatment.33

Sutherland et al attempted to conduct a PCR-stratified RCT based on PCR positivity for C. pneumoniae or Mycoplasma pneumoniae.31 Due to lower than anticipated numbers of PCR-positive participants, the two arms could not be successfully matched in number, but comparison of the two groups showed no significant differences in outcome between PCR positive (n=12) and PCR negative (n=80) after treatment with 16 weeks of clarithromycin.33

The AMAZES study monitored a subgroup of study participants for emergence of antimicrobial resistance to azithromycin within serial sputum samples over the 48-week study period.40 At the study end, resistance to macrolides was found in 19/39 (48.7%) receiving azithromycin and 12/42 (28.6%) receiving placebo.40 AZISAST performed a similar subgroup analysis in 46 participants demonstrating an increase in macrolide-resistant streptococci from 47.8% to 87% participants in the azithromycin group and a reduction from 39.1% to 35% of participants in the placebo group.30 Interestingly, the percentage of macrolide-resistant streptococci reduced from 73.8% to 45.9% in the 4-week washout period at the end of the study.30 Further analysis of the oropharyngeal microbiome of a subgroup from AZISAST demonstrated a fivefold increase in Streptococcus salivarius and corresponding fivefold decrease in Leptotrichia wadei during azithromycin treatment. However, azithromycin had little impact on the rest of the microbiota. In keeping with the culture-based microbiology, the microbiome had already returned to pre-treatment levels in 50% participants by the end of the 4-week washout period.49

This suggests that chronic macrolide therapy may increase antimicrobial resistance to macrolides. This result has yet to be reproduced in other large studies and the clinical impact of this remains unknown. It is also unclear whether this increase in resistance is temporary or fluctuating. However, it does suggest that monitoring for antimicrobial resistance may be of benefit in those receiving chronic macrolide therapy. Use of breaks in chronic therapy, if the desired clinical outcome is achieved, may be considered to reduce resistance, as is the practice in some bronchiectasis services. Further work is needed to examine this relationship in more depth and determine the clinical impact of increasing resistance rates.

Safety and adverse events
Reporting of adverse events in the majority of studies was of low quality with no specific assessment for the well-documented adverse effects of macrolide therapy in the majority of studies. However, overall there was little evidence of significant reactions or side effects secondary to macrolide therapy in asthma. The most commonly reported side effects were gastrointestinal, including nausea and abdominal pain. In the meta-analysis performed by Reiter et al, a significantly increased risk of nausea was seen in those receiving macrolide therapy.41 Black et al also showed that 12.4% (13/105) of the roxithromycin group developed nausea compared with 4.5% (5/112) of the placebo group.29

The AMAZES and AZISAST trials specifically assessed and reported adverse events as part of the planned study protocols.30 40 In AMAZES, the overall serious adverse event (SAE) rate in the azithromycin group was 7.5% (16/213) compared with 12.8% (26/203) in the placebo group.30 Gastrointestinal side effects were more common in those on azithromycin compared with placebo—diarrhoea (33.8% (72/213) vs 19.2% (39/203)) and abdominal pain (17.8% (38/213) vs 14.8% (30/203)).30 QTc prolongation was not seen more commonly in the azithromycin group with 0.5% (1 participant) of each group developing ECG changes during the study.40 There was also no difference in the rates of tinnitus (0.9% (2/213) vs 1% (2/203)) or hearing loss (2.8% (6/213) vs 3.4% (7/203)).30

In AZISAST, the SAE rate was identical in both placebo and azithromycin groups (11%).30 Discontinuation of treatment was more common in the placebo group (9% (5/54 vs 4% (2/53)). Diarrhoea, nausea and abdominal pain were also more common in the placebo group. Two participants in the azithromycin group developed abnormal liver function tests, but the severity of this was not reported. No participants reported any change in their hearing.30

Overall, it appears that macrolides are well tolerated in people with asthma, although the quality of evidence is low and inclusion of formal assessment for adverse events should be included in future studies. There is a slightly higher incidence of gastrointestinal side effects which did not lead to cessation of treatment. Warning of the possibility of these side effects developing should be given to patients on initiation of treatment. There is insufficient evidence on the frequency of other adverse events specifically QTc prolongation, liver function abnormalities and audiological changes to make a formal recommendation on screening for these events. In general, participants with evidence of QTc prolongation or hearing loss were excluded from study populations further limiting the applicability of this evidence to the wider asthma population.

Outcome weighting
In the creation of this guideline, a set of key outcome measures in relation to macrolide therapy in management of respiratory disease were determined. However, this guideline’s scope covers a spectrum of respiratory disease and the importance of individual outcome measures within each disease area varies. In regards to asthma, the GDG gave importance to those outcomes of most clinical relevance to respiratory physicians managing people with asthma and our recommendations reflect this. The group only made recommendations on these outcomes where the quality of evidence and magnitude of change was deemed sufficient to be of clinical utility. Therefore, recommendations are made in regard to exacerbation reduction, but not in relation to symptoms or quality of life as the changes seen in these outcomes were too small to be clinically important.

The body of evidence in regard to other outcomes, which may be of scientific or physiological relevance, is outlined in the evidence summary below, but no recommendations have been made in areas where the outcome is not used as a clinical endpoint—for example, bronchial hyper-responsiveness.

Evidence summary
Oral macrolide therapy improves quality of life in people with asthma, but changes are small and their clinical significance is uncertain. (Moderate)
Treatment with macrolides results in an improvement in asthma symptoms in some patients although this is small in magnitude and may not be applicable to all patients with asthma. (Low)

Oral macrolide therapy can reduce exacerbations of asthma in adults (50–70 years) with ongoing symptoms despite >80% adherence to high-dose inhaled steroids (>800 µg/day) and at least one exacerbation requiring oral steroids in the past year. (Low)

Azithromycin given thrice weekly at 500 mg over 48 weeks results in a reduction in asthma exacerbations. (Low)

In some individuals, oral macrolide therapy may result in a reduction in oral steroid dose, but this is not a consistent finding. (Low)

Oral macrolide therapy can result in a small improvement in lung function and PEFR in people with asthma. (Low)

Oral macrolide therapy may reduce airway inflammation in asthma. (Low)

Oral macrolide therapy may reduce bronchial hyperresponsiveness in asthma. (Low)

Two studies demonstrated that treatment with oral macrolides (azithromycin) may result in an increase in bacteria within sputum resistant to macrolides, but currently there is no evidence of this adversely affecting clinical outcomes. (Low)

Oral macrolide therapy may result in an increase in gastrointestinal side effects including abdominal pain, nausea and diarrhoea. (Low)

There is insufficient evidence to make any recommendation on the impact of oral macrolide therapy on mortality, exercise capacity, disease progression or sputum characteristics in people with asthma.

Recommendations

- Oral macrolide therapy could be considered to reduce exacerbation frequency in adults (50–70 years), with ongoing symptoms despite >80% adherence to high-dose inhaled steroids (>800 µg/day) and at least one exacerbation requiring oral steroids in the past year. This recommendation reflects the population within the AMAZES RCT which represents the highest quality evidence of macrolide therapy leading to a significant reduction in exacerbations. (Conditional)

- Treatment with azithromycin should be considered for a minimum of 6–12 months to assess evidence of efficacy in reducing exacerbations. (Conditional)

- Oral macrolide therapy should not be offered as a way to reduce oral steroid dose; in some individuals, this may result as a consequence of a reduction in exacerbations or symptoms. (Strong)

Good practice points

- Optimisation of other asthma therapies including establishing good adherence to inhaled therapies should be performed before considering a trial of oral macrolide therapy.

- Referral to a respiratory specialist or specialist asthma service should be considered prior to initiation of macrolide therapy aimed at reducing exacerbation frequency.

- For safety purposes, an ECG should be performed prior to initiation of macrolide therapy to assess QTc interval. If QTc is >430 ms for men and >470 ms for women, this is considered a contraindication to initiating macrolide therapy. Baseline liver function tests should also be measured.

- Patients should be counselled about potential adverse effects before starting therapy including gastrointestinal upset, hearing and balance disturbance, cardiac effects and microbiological resistance.

- Microbiological screening of sputum before and during macrolide therapy may be clinically helpful in patients who are able to expectorate sputum. This would allow monitoring for development of resistance and detect changes in microbial growth to direct appropriate antibiotic therapy if required. However, the resource implications of this approach have not been assessed.

- If oral macrolide therapy is considered, justification for ongoing treatment should be guided by clinical response based on specific outcome measures including exacerbation frequency, symptoms and quality of life assessed at baseline.

- A risk:benefit profile should be considered in each individual if significant side effects from oral macrolide therapy develop. If gastrointestinal side effects occur at the higher dose of azithromycin (500 mg thrice weekly) a dose reduction to azithromycin 250 mg thrice weekly could be considered if macrolide therapy has been of clinical benefit.

- Liver function tests should be checked 1 month after starting treatment and then every 6 months. An ECG should be performed 1 month after starting treatment to check for new QTc prolongation. If present, treatment should be stopped.

- Symptom improvement with macrolide treatment may be minimal and not consistent across all people with asthma. If macrolide therapy is considered for symptom reduction, this should be for a defined period (6–12 months) and stopped if no symptomatic improvement is seen. Use of a validated symptom score, such as the ACQ, may be useful to help make this assessment less subjective.

- If the desired clinical outcome is achieved the possibility of breaks in therapy may be considered to reduce treatment burden for patients. It is unclear whether this may also reduce antimicrobial resistance rates.

See quick reference guide in online supplementary file 1.

Research recommendations

Further research, over a longer period, is needed to investigate the role of long-term macrolide therapy in reducing exacerbations of asthma.

Use of validated scoring systems and creation of a core outcome set should be considered in future trials of macrolide therapy in asthma to allow more accurate delineation of clinical response and comparison between studies.

Head to head comparisons of different macrolide therapy and of differing dose regimens for the same macrolide are required to optimise the use of macrolide therapy in asthma. Research into the discrepancy between significant reductions in inflammation but minimal improvements in clinical parameters is required.

Research into the impact of macrolide therapy in different clinical phenotypes of asthma should be performed in order to assess whether greater benefit is seen in one phenotype over another.

Research into the overlap between asthma and bronchiectasis and the impact of this on response to macrolides may assist in identifying asthma phenotypes which may respond differently to macrolide therapy.
SECTION 5: BRONCHIECTASIS

Introduction

Bronchiectasis is a chronic respiratory condition in which abnormally dilated bronchi are radiologically displayed in patients with a relevant clinical syndrome. These patients suffer from multiple symptoms, including a chronic productive cough, recurrent exacerbations associated with bronchial infection, dyspnoea and fatigue. While patients with cystic fibrosis can have both the radiological changes and the clinical syndrome, the term ‘bronchiectasis’ generally refers to those who do not have cystic fibrosis and the condition is often referred to as non-cystic fibrosis bronchiectasis. Even after the exclusion of cystic fibrosis, a wide range of aetiologies are attributed to bronchiectasis, though a large proportion of cases are described as idiopathic or post-infective. Recent UK data report prevalence rates of 566/100 000 in adult women and 486/100 000 in adult men. Patients with bronchiectasis bear a significant burden of morbidity and mortality.

The basic concept for the management of bronchiectasis is to attempt to break the vicious recurrent cycle of chronic bacterial infection, inflammation, impaired mucociliary clearance and structural lung disease. With the goal of breaking this cycle and the extrapolation of management strategies from cystic fibrosis, it is unsurprising that macrolides have been used to try to improve outcomes in bronchiectasis. As well as having broad antimicrobial effects, anti-inflammatory effects are likely. Studies in patients with bronchiectasis have suggested reductions in Th17 cell responses and evidence of reduced airway inflammation via measuring markers such as IL8, neutrophil elastase and matrix metalloproteinase. A further attractive mechanism, particularly in a condition where P. aeruginosa is an important pathogen, is the inhibition of quorum sensing (a form of chemical signalling between bacteria).

Evidence base

Davies and Wilson described early experiences of using azithromycin in 2004 and subsequent published studies of varying quality, using different regimens, have added to the available evidence. The main body of high-level evidence is based on three RCTs (see table 6). These studies were performed in the Netherlands, Australia and New Zealand and were all placebo controlled. The macrolides used were azithromycin in the Netherlands and New Zealand studies, while the Australian study used erythromycin. Audit data suggest that azithromycin is the preferred macrolide in the UK at present. The two studies which used azithromycin had different dosing regimens (250 mg daily and 500 mg three times a week). In addition, other studies of lower quality evidence have used other regimens such as azithromycin 250 mg three times a week, clarithromycin 500 mg once daily and roxithromycin 150 mg a day. The lack of consistency in the macrolide regimen used, and the absence of head-to-head comparisons, makes the optimum macrolide and dosing unclear. Also, because none of the studies were longer than a year, the longer-term benefits and risks have not been established. Subsequent to the three RCTs which provide high-quality evidence, meta-analyses have been produced. While this provides pooled data, some lower quality evidence influence the output. A further meta-analysis of individual patient data (IPD) from the three main RCTs was published after the completion of literature searches and is referred to below because of the potential importance of its findings.

Other recent guidance

The BTS bronchiectasis guidelines cover options beyond macrolides when considering long term antibiotic therapy in stable disease. They recommend macrolides as first choice for patients without P. aeruginosa with inhaled gentamicin as a second-line alternative. For patients with P. aeruginosa, the BTS guideline recommends inhaled colistin with macrolides as an alternative for those intolerant of inhaled antibiotics. Interestingly, the IPD meta-analysis by Chalmers et al suggests that macrolides are effective in patients with and without P. aeruginosa. In addition, the BTS guideline recommends a starting dose of azithromycin of 250 mg three times a week to minimise side effects. While accepting that side effects are common with macrolides, we note that dropouts from studies due to side effects are rare and the strongest evidence of efficacy lies with higher dosing regimes. We suggest a pragmatic starting regime of azithromycin 500 mg three times a week or 250 mg daily in patients with bronchiectasis. However, if there is a patient history of drug intolerances, it would be sensible to start at the lower dose of 250 mg three times a week.

Table 6  Three main randomised controlled trials of macrolides in bronchiectasis

<table>
<thead>
<tr>
<th>Country</th>
<th>Population</th>
<th>Intervention</th>
<th>Follow-up</th>
<th>Patients</th>
<th>Primary outcome (exacerbations)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMBRACE Wong</td>
<td>New Zealand</td>
<td>Mean age of 60 At least one exacerbation requiring antibiotics in last year HRCT-defined bronchiectasis</td>
<td>Azithromycin 500 mg three times a week 6 months</td>
<td>141</td>
<td>Exacerbations in first 6 months: 0.59 (treatment group) vs 1.57 (placebo group) p&lt;0.0001</td>
</tr>
<tr>
<td>BAT Altenburg</td>
<td>Netherlands</td>
<td>Mean age of 62 years At least 3 LRTIs treated with antibiotics in the last year A sputum culture isolating a respiratory pathogen in the last year Bronchiectasis defined by HRCT or bronchography</td>
<td>Azithromycin 250 mg daily 1 year</td>
<td>83</td>
<td>Exacerbations in 52-week treatment period: 0.84 (treatment group) vs 2.05 (placebo group) p&lt;0.001</td>
</tr>
<tr>
<td>BLESS Serisier</td>
<td>Australia</td>
<td>Mean age of 62 At least two exacerbations requiring antibiotics in the last year Daily sputum production Clinically stable for at least 4 weeks. HRCT-defined bronchiectasis</td>
<td>Erythromycin ethylsuccinate 400 mg twice daily 48 weeks</td>
<td>117</td>
<td>Annualised exacerbation rate: 1.29 (treatment group) vs 1.97 (placebo group) p=0.003</td>
</tr>
</tbody>
</table>

LRTI, Lower respiratory tract infection.
Exacerbations

Due to the three main RCTs using exacerbations as a primary endpoint and the subsequent meta-analyses, there is high level evidence which is in favour of using long-term macrolides to reduce exacerbations in bronchiectasis. As in other respiratory conditions, there is variability in the interpretation of what exactly constitutes an exacerbation. A consensus definition has been published; however, this was subsequent to the important studies in this field.

The previously mentioned three principal RCTs described benefit for this outcome when macrolides were used for between 6 and 12 months. The BLESS study described an incidence rate ratio of 0.57 (95% CI 0.42 to 0.77) when compared with no intervention. The BAT study reported an absolute risk reduction of 33.5% (95% CI 14.1% to 52.9%). Of interest in this study, there was an improvement in median number of exacerbations from 5 to 2 in the placebo group. Over a 6-month period, the EMBRACE study identified a rate ratio of 0.38 between the treatment group and the placebo group (95% CI 0.26 to 0.54). It is important to note that the EMBRACE study continued for a further 6 months after stopping the intervention and these benefits were maintained. However, it should also be noted that this study provided two definitions of ‘exacerbations’ and the results were not consistent, and that there was a higher baseline exacerbation rate in the control group. The meta-analysis data are consistent with an improvement in exacerbation rates.

For macrolides to be used to reduce exacerbations, there needs to be a relevant exacerbation rate at baseline to consider when implementing this therapy. There were different entry criteria for the main trials with regards to baseline exacerbation rates. While entry into the EMBRACE study required just a single exacerbation for study recruitment, the mean exacerbation rate in the preceding year was over three (3.34 in treatment group and 3.93 in control group). In the BLESS and the BAT studies, two and three exacerbations were required, respectively, in the preceding year. In the BLESS study, approximately a third had five or more exacerbations in the preceding year, but it is unclear what the mean or median baseline exacerbation rate was. In the BAT study, the median baseline exacerbation rate was 4 (and 5 in the control group). In light of the baseline exacerbation rates in these trials, and three or more exacerbations being a marker of disease severity, a threshold of three or more exacerbations per year seems appropriate as a guide. It should be acknowledged that the aforementioned IPD meta-analysis that has followed the literature search for this guideline performed multiple subgroup analyses including for different exacerbation rates. One of these subgroups was those with a preceding exacerbation rate of less than this threshold (ie, 1 or 2 per year). While this group was small (37 receiving macrolides vs 36 receiving placebo), there appeared to be a benefit with regards to exacerbations, but a trend towards decreased QOL. It is conceivable that the benefits of macrolides may be outweighed by side effects in those with lower exacerbation rates. This subgroup require further study.

While out of keeping with the BTS Bronchiectasis Guideline and some UK respiratory clinicians’ current practice, the regimens used in the three main RCTs (azithromycin 250 mg daily, azithromycin 500 mg three times a week and erythromycin ethylsuccinate 400 mg twice a day) have the greatest supportive evidence of benefit. The BTS bronchiectasis guideline suggest a lower starting dose (azithromycin 250 mg three times a week) noting the dose-related nature of side effects. Withdrawals due to side effects in the three main studies were, however, rare. It would seem sensible to start at the higher dose (azithromycin...
500 mg three times a week or 250 mg daily) unless there is a history of previous drug intolerances. 7 As well as variable dosing regimens, variable durations of therapy have also been studied. The highest quality studies used macrolide therapy for 6 or 12 months. While it is conceivable that it continues to provide benefit, there is no evidence either in favour or against the prolongation of therapy beyond 12 months.

**Exercise capacity**

Exercise capacity was measured by a 6 min walk test in both the BLESS and the EMBRACE studies. 61 63 This provides some high-level evidence via objective measures with blinded subjects and investigators. In neither study was there statistical evidence of benefit or harm from therapy. In addition, the activity component of the SRGQ was reported in the EMBRACE study 61 and no impact on this outcome was seen. On the basis of this evidence from two studies, there is no indication that macrolides influence exercise capacity or tolerance.

**Sputum characteristics**

Sputum production is a common occurrence in many patients with bronchiectasis and is considered a significant issue by patients. 65 75 There is, however, no high-quality evidence with regards to the effects of macrolide therapy on sputum. There is moderate level evidence of reduced sputum weight and sputum volume. 63 70 Estimates of the effect of therapy may be a reduction of approximately 11 mL per day (95% CI −12.70 to −8.83) based on pooled data. 63 The BLESS study reported a treatment effect of a reduction of 4.3 g per day. 63 It is unclear whether this scale of benefit is important to a patient but could be considered in discussion alongside assessment of other factors such as exacerbation rate when considering therapy.

**Safety and adverse events**

The evidence base for adverse events is quite variable within the bronchiectasis literature. While QTc prolongation is a potential concern with long-term macrolide use, there is very little evidence with regards to this adverse effect. In the BLESS study, QTc was monitored and one patient was withdrawn due to concerns with regards to prolongation, 63 however pre-therapy and post-therapy ECGs suggested that this may not have been a drug-related effect. The EMBRACE and BAT studies did not address this potential adverse event. 61 62 An adverse impact on hearing is a further concern with macrolides; however, there is also limited evidence for this potential side effect. The BLESS and EMBRACE studies did not assess it, while the BAT study just performed a post-study questionnaire (and did not detect an effect). 61 63

There is, however, considerably more evidence in these cohorts for gastrointestinal side effects. The meta-analysis pooled data show high-quality evidence for diarrhoea being an adverse event associated with long-term macrolide use. 69 71 72 The frequency of patients suffering diarrhoea was 19.3%–20.6% in the treatment groups compared with 4.1%–4.5% in the placebo groups. There was also high-level evidence for abdominal pain/discomfort when reported in the BAT study (18.6% vs 2.5%, RR 7.44) and meta-analysis data (OR 6.97). 62 71 Despite these gastrointestinal side effects, meta-analysis data assessing for withdrawal from studies due to side effects did not reveal any differences between those on macrolide or placebo (OR 1.18, 95% CI 0.33 to 4.19). 69

While not a patient-centred outcome, high-quality evidence from the BLESS study reports increased micro-organism resistance. 63 Further moderate level evidence was also added from the BAT study, though not the EMBRACE trial where routine testing was not performed. 61 62 In the BLESS study, there was a significant increase of macrolide resistance in commensal oropharyngeal streptococci with erythromycin use (27.7% vs 0.04%, p<0.001). 63 The BAT study provided a significant amount towards the meta-analysis data and in the pathogens that were tested for macrolide sensitivity, 88% became resistant compared with 26% in the placebo group. 62 69 At present, there is no evidence that the resistance seen either does or does not have an impact on clinical outcomes.

**Outcome weighting**

The available data allow assessment of the pre-specified outcomes with varying levels of evidence. Certain outcomes were excluded from our final assessment and recommendations. These included lung function, hospital admission rates, disease progression and death. Lung function was excluded despite available data, as the guideline group did not feel it is a useful measure when taken in the context of the current evidence. The trials were of too short a period for a meaningful change in lung function to be elicited despite statistically significant outcomes. For example, meta-analysis data from five studies described a statistically significant greater improvement of FEV1 of 0.02 L (mean weighted difference). 70 This was not considered a clinically meaningful change for an individual patient. Hospital admission rate data were available; however, due to such low incidence rates for this outcome in the three RCTs which reported it, no recommendations are made for this outcome. 61 63 64 The outcomes of disease progression and death are not commented on due to a lack of data. The outcomes of QOL, symptoms, exacerbation rates, exercise capacity, sputum characteristics and adverse events were all considered ‘important’ in the GRADE process.

**Evidence summary**

There is evidence of an improvement in QOL as measured by SGRQ when azithromycin 250 mg daily is used for 1 year (High). Other high-quality studies with other dosing regimens showed a trend towards benefit but not a statistically significant benefit. Meta-analysis data reported a benefit but there are concerns regarding the inclusion of a study of lower quality (Low).

There is evidence from high-quality studies of improvement in symptom scores; however, inconsistency exists which may be due to different scoring systems being used. (Moderate)

Long-term macrolide treatment reduces exacerbations in bronchiectasis. (High)

When using macrolides to reduce exacerbation rates, the dosing regimens with the greatest supportive evidence are azithromycin 250 mg daily, azithromycin 500 mg three times a week and erythromycin ethylsuccinate 400 mg twice a day. (Moderate)

Studies with other dosing regimens, including azithromycin 250 mg three times a week (as pragmatically suggested in the BTS Bronchiectasis Guideline), have also reported a reduction in exacerbation, but have a lower evidence base. (Low)

The studies with the greatest evidence for reducing exacerbations used therapy for a minimum of 6 months. (High) The impact beyond 12 months is unknown.

There is evidence for a reduction in exacerbations over 12 months when therapy is used for 6 months and then not for the subsequent 6 months. (Moderate) It is unknown what the impact is of subsequently recommencing.

Long-term macrolide therapy is not associated with improved exercise capacity. (High)
Long-term macrolide therapy may reduce sputum volume and weight. (Moderate)

Long-term macrolide therapy is associated with diarrhoea and with abdominal pain. (High)

Long-term macrolide usage can result in increased antimicrobial resistance. (High) It is unknown if this has a clinical impact.

Recommendations

► Long-term macrolide treatment could be offered to reduce exacerbations in those with high exacerbation rates (ie, 3 or more per year). (Strong)

► The dosing regimens with the greatest supportive evidence, when using macrolides to reduce exacerbation rates, are azithromycin 500 mg three times a week, azithromycin 250 mg daily and erythromycin ethylsuccinate 400 mg twice a day. A starting dose of azithromycin 250 mg three times a week could be used to minimise side-effect risk with subsequent titration according to clinical response. (Conditional)

► When using macrolides to reduce exacerbation rates, therapy should be offered for a minimum of 6 months. (Strong)

► Macrolides can be considered with the aim of improving QOL but may require a long period of therapy (eg, 1 year) for significant effects. (Conditional)

Good practice points

✓ Therapies should be optimised in accordance with BTS Bronchiectasis Guidelines before considering long-term macrolide therapy (eg, airway clearance techniques and attendance at pulmonary rehabilitation courses).

✓ Macrolides should only be started following discussion and shared decision-making between the patient and a respiratory specialist.

✓ For safety purposes, an ECG should be performed prior to initiation of macrolide therapy to assess QTc interval. If QTc is >450 ms for men and >470 ms for women, this is considered a contraindication to initiating macrolide therapy. Baseline liver function tests should also be measured.

✓ Patients should be counselled about potential adverse effects before starting therapy including gastrointestinal upset, hearing and balance disturbance, cardiac effects and microbiological resistance. Microbiological assessment of sputum should be performed before therapy, including investigation for NTM. Macrolide monotherapy should be avoided if an NTM is identified. When evaluating for NTM infection, macrolides should not be used for 2 weeks before microbiological testing.

✓ Accurate assessment of baseline exacerbation rate should be determined before starting long-term macrolides for bronchiectasis.

✓ Liver function tests should be checked 1 month after starting treatment and then every 6 months. An ECG should be performed 1 month after starting treatment to check for new QTc prolongation. If present, treatment should be stopped.

✓ Subsequent follow-up at 6 months and 12 months should determine whether benefit is being derived from therapy. If there is no benefit, treatment should be stopped.

✓ Even if benefit is seen, consideration should be given to stopping treatment for a period each year, for example, over the summer. Such a drug holiday may help with reducing the development of resistance while maintaining efficacy because the vicious cycle has been broken.

See quick reference guide in online supplementary file 1.

Research recommendations

Long-term studies of microbiological impact of prolonged macrolide therapy.

Head-to-head comparison of different dose regimens for the same macrolide (eg, 250 mg three times a week of azithromycin vs 500 mg three times a week).

Head-to-head comparisons of different macrolides (eg, azithromycin vs erythromycin).

Prolonged studies of benefit and risk beyond 12 months of use.

Studies assessing the effect of macrolides following re-introduction of therapy after a break.

Use of bronchiectasis specific QOL measures in macrolide trials.

Comparison of outcomes in patients with different baseline exacerbation rates when treated with macrolides.

Comparison of outcomes in patients with different baseline microbiological culture/microbiome profiles when treated with macrolides.

Comparison of outcomes in patients with different baseline QOL scores when treated with macrolides.

Comparison studies of long-term macrolides to other oral or inhaled prophylactic regimens.

Studies looking into the benefits of combined macrolide and inhaled antibiotic regimens.

SECTION 6: COPD

Introduction

COPD is a progressive, inflammatory disease of the airways, characterised by ongoing development of non-reversible airflow limitation and acute episodes of exacerbation. Acute exacerbations are described as episodes of acute worsening of respiratory symptoms that necessitate a change in regular medication. The natural disease course, health-related QOL, hospital admissions and mortality associated with COPD are influenced by acute exacerbations. In addition, acute exacerbations of COPD (AECOPD) are estimated to account for between 50% and 75% of the total costs for COPD treatment. Treatment interventions effective in preventing acute exacerbations are therefore of significant clinical benefit.

The mechanisms for COPD disease progression are complex and involve inflammatory and immune responses. Neutrophils, IL-17 and IL-23 are considered to play an active role in disease progression while CD8-positive T cells, neutrophils and macrophages mediate chronic inflammatory responses. Acute exacerbations of COPD are postulated to be caused by bacteria or bacteria in combination with viral infection in up to 50% of cases. Macrolide antibiotics possess both anti-inflammatory and antibacterial effects and are therefore an important potential treatment strategy to prevent AECOPD.

The potential mechanisms of action and clinical implication of long-term macrolide antibiotic therapy in COPD have been described and reviewed in several studies.

Table 7 summarises the potential effects of macrolides on disease modulators in COPD.

Evidence base

To confirm the hypothesis that macrolide antibiotics are an effective treatment strategy in COPD to prevent acute exacerbations and possibly modify disease progression, several studies have been conducted yielding varied results. The evidence base for effectiveness of macrolide antibiotics in this context is compromised by the significant heterogeneity in populations.
Nine RCTs were reviewed and included in the guideline. The published since 2000 with evidence included in meta-

<table>
<thead>
<tr>
<th>Matrix of evidence analysed for COPD</th>
<th>Table 8</th>
</tr>
</thead>
<tbody>
<tr>
<td>RCT</td>
<td></td>
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<tr>
<td>Meta-analysis j</td>
<td></td>
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<tr>
<td>Albert et al[9][3]</td>
<td>● ● ● ●</td>
</tr>
<tr>
<td>He et al[9][3]</td>
<td>● ● ● ●</td>
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<tr>
<td>Blasi et al[9][6]</td>
<td>● ● ● ●</td>
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<tr>
<td>Seemungal et al[9][4]</td>
<td>● ● ● ●</td>
</tr>
<tr>
<td>Banerjee et al[9][6]</td>
<td>● ● ● ●</td>
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<tr>
<td>Suzuki et al[9][8]</td>
<td>● ● ● ●</td>
</tr>
<tr>
<td>Uzun et al[9][3]</td>
<td>● ● ● ●</td>
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<tr>
<td>Simpson et al[9][4][6]</td>
<td>● ● ● ●</td>
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<tr>
<td>Berkhof et al[9][6]</td>
<td>● ● ● ●</td>
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<tr>
<td>Donath et al[9][2][3]</td>
<td>● ● ● ●</td>
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<tr>
<td>Herath and Poole[9][3][6]</td>
<td>● ● ● ●</td>
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<tr>
<td>Simoons et al[9][2][7]</td>
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<tr>
<td>Yao et al[9][2][8]</td>
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<tr>
<td>Lee et al[9][2][9]</td>
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<tr>
<td>Ni et al[9][0][0]</td>
<td>● ● ● ●</td>
</tr>
<tr>
<td>Wedzicha et al[9][2]</td>
<td>● ● ● ●</td>
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</table>

investigated, intervention (type, dose and duration of macro-

Cohort studies
Several retrospective cohort studies were identified, with the largest two (Pomares et al and Yamaya et al) including a total number of 326 patients.88 89 The quality of evidence in reporting exacerbations and QOL was deemed poor due to the risk of selection and attrition bias.

Included studies
Based on the aforementioned appraisal of evidence, the included studies for recommendations in this guideline were focused on RCTs and systematic reviews as demonstrated in the evidence matrix (Table 8).

Other recent guidance
GOLD 2019
In patients with COPD optimised with inhaled therapy who are still experiencing exacerbations, the best available evidence exists for the use of azithromycin, especially in those who are not current smokers. Consideration to the development of resistant organisms should be factored into decision-making.90

NICE 2018
Recommends after non-pharmacological and pharmacological optimisation to consider azithromycin (usually 250 mg three times a week) for people with COPD more prone to daily sputum production if they have stopped smoking and have more than four exacerbations per year, prolonged exacerbations or exacerbations resulting in hospital admission.
ERS/ATS recommendation
Patients with COPD with moderate to very severe airflow obstruction (post-bronchodilator FEV1/FVC <0.70 and an FEV1% predicted of <80%) and exacerbations despite optimal inhaled therapy, treatment with a macrolide antibiotic to prevent future exacerbations is suggested as a conditional recommendation.9

Cochrane systematic reviews
Prophylactic antibiotics for patients with COPD that included macrolide therapy found for every eight participants treated, one person would be prevented from suffering an exacerbation. Not all the antibiotic regimens had the same impact on exacerbations, with results suggesting that antibiotics Prescribing off-label given at least three times per week may be more effective than pulsed antibiotic regimens.8

Symptoms and QOL
Health status measurement is a standardised and objective means of quantifying the impact of disease on patients’ daily life, health and well-being. Health status questionnaires usually address emotional and psychological effects of the illness as well as the physical. The St George’s Respiratory Questionnaire (SGRQ) is validated to measure health impairment in patients with COPD. It is in two parts; Part 1 produces the Symptoms score, and Part 2 the Activity and Impacts scores, which results in a total score.9 The COLUMBUS trial83 enrolled patients with at least three exacerbations in the year prior to entering the trial, the severity of COPD and the definition of the severity of an acute exacerbation.8

Exacerbations
Nine RCTs were identified that reported the effect of long-term macrolide antibiotics on AECOPD. The study entry criteria varied in terms of the number of exacerbations in patients in the year prior to the trial, the severity of COPD and the definition of the severity of an acute exacerbation.

Variations
The COLUMBUS trial83 enrolled patients with at least three exacerbations in the year prior to entering the trial, Blasi et al98 reported the number of exacerbations in patients the year before entering the trial as 3.1 (±0.9) and 3.0 (±1.1) in the control and intervention arm, respectively. The largest RCT (Albert et al 2011) included in the entry criteria at least one acute exacerbation in the year prior to trial entry.9 The remainder of studies did not define number of exacerbations prior to entering the trial in the inclusion criteria.

There was significant variation in severity of disease of patients enrolled in trials, including reporting of severity of disease and airflow limitation measurements reported as the

### Table 9 Summary of COPD trials

<table>
<thead>
<tr>
<th>Country conducted</th>
<th>Population</th>
<th>Mean age: intervention/control</th>
<th>COPD intervention/control</th>
<th>Required an exacerbation in year before</th>
<th>Intervention</th>
<th>Follow-up on treatment</th>
<th>Patients</th>
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<tbody>
<tr>
<td>Albert et al83 USA</td>
<td>Mean age: 65/66 COPD FEV1: %: 39/40</td>
<td>Azithromycin 250 mg a day</td>
<td>12 months</td>
<td>558/559</td>
<td>Time to first exacerbation</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>He et al87 China</td>
<td>Mean age: 68/69 COPD FEV1: %: 44.3/42.1</td>
<td>Erythromycin 150 mg three times a day</td>
<td>6 months</td>
<td>18/18</td>
<td>Exacerbations airway inflammation</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Seemungal et al80 UK</td>
<td>Mean age: 66/67 COPD FEV1: %: 49.5/50.6</td>
<td>Erythromycin (stearate tablets) 250 mg bd</td>
<td>12 months</td>
<td>53/56</td>
<td>Exacerbations airway inflammation</td>
<td></td>
<td></td>
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<tr>
<td>Suzuki et al82 Japan</td>
<td>Mean age: 69/71 COPD FEV1: %: 47/61</td>
<td>Erythromycin (ethylsuccinate tablets) 200–400 mg a day</td>
<td>12 months</td>
<td>55/54</td>
<td>Frequency of common cold and exacerbations</td>
<td></td>
<td></td>
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<td>Uzun et al81 Netherlands</td>
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<td>Azithromycin 500 mg three times a week</td>
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ERS/ATS recommendation
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<td>Seemungal et al80 UK</td>
<td>Mean age: 66/67 COPD FEV1: %: 49.5/50.6</td>
<td>Erythromycin (stearate tablets) 250 mg bd</td>
<td>12 months</td>
<td>53/56</td>
<td>Exacerbations airway inflammation</td>
<td></td>
<td></td>
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<tr>
<td>Suzuki et al82 Japan</td>
<td>Mean age: 69/71 COPD FEV1: %: 47/61</td>
<td>Erythromycin (ethylsuccinate tablets) 200–400 mg a day</td>
<td>12 months</td>
<td>55/54</td>
<td>Frequency of common cold and exacerbations</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Uzun et al81 Netherlands</td>
<td>Mean age: 64/76/64 COPD FEV1: %: 44/24/5</td>
<td>Azithromycin 500 mg three times a week</td>
<td>12 months</td>
<td>47/45</td>
<td>Rate of exacerbations</td>
<td></td>
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mean FEV₁ measured in litres or as percentage predicted. Albert et al described severity (GOLD criteria) of patients in the intervention/control arm as follows: GOLD 2 (26%/26%), GOLD 3 (40%/40%) and GOLD 4 (34%/33%). Uzun et al included patients as follows: GOLD 1 (47%), GOLD 2 (30/27%), GOLD 3 (38/40%) and GOLD 4 (28/22%).

The long-term macrolide therapy was more effective than placebo (RR=0.46, 95% CI 0.18 to 1.18, p=0.11). In contrast, studies as reported by Berkhof et al demonstrated a similar effect (RR=0.53, 95% CI 0.43 to 0.83, p=0.01). Studies using azithromycin in 1200 patients showed a significant reduction (RR=0.59, 95% CI 0.37 to 0.93, p=0.02) and with erythromycin (254 patients) for 6–12 months a similar effect (RR=0.53, 95% CI 0.43 to 0.83, p=0.01).

The recent published ERS/ATS guideline pooled data for studies that included treatment for 12 months and excluded the Suzuki trial (due to risk of bias, non-blinded study as described above) and demonstrated a reduced rate of AECOPD (RR=0.76, 95% CI 0.68 to 0.86).

Data analysis demonstrated no significant difference in exacerbation rates at 3 months; it is recognised that these studies were likely underpowered to demonstrate effect size. Banerjee et al reported the effects of clarithromycin for 3 months in 67 patients (RR=3.27, 95% CI 0.53 to 20.18, p=0.2) and Simpson et al on 30 patients with azithromycin for 12 weeks (RR=0.38, 95% CI 0.14 to 1.05, p=0.06). The number of exacerbations at 3 months of treatment did not demonstrate significant variation, as reported by Berkhoff et al, with azithromycin in 84 patients (RR=0.46, 95% CI 0.18 to 1.18, p=0.11). In contrast, studies reporting exacerbation rates at 6–12 months demonstrated a similar effect (RR=0.53, 95% CI 0.43 to 0.83, p=0.01).

Onset of first exacerbation
Macrolide therapy significantly delayed onset of first exacerbation. Pooled data of patients on macrolide antibiotics for 12 months demonstrates a mean 81-day delay in first AECOPD (95% CI 53.3 to 109.8 days longer) and the study by Seemungal et al in 109 patients with erythromycin for 12 months demonstrated a mean 182 days delay in first AECOPD (p=0.02).

Daily use of azithromycin long term (12 months) was demonstrated in Albert et al (1117 patients) to significantly reduce exacerbation rate (RR=0.83, 95% CI 0.72 to 0.95, p=0.01). Two studies adopted intermittent dosing regimens (see table 9). The Suzuki et al rather than 6 months, as reported by Uzun et al, appeared to be more effective to reduce exacerbation rate (RR=0.58, 95% CI 0.42 to 0.80, p<0.01).

Subgroups
Han et al published subgroup analysis of the largest RCT (1117 patients) using azithromycin 250 mg a day for 12 months. There was no convincing evidence that treatment efficacy was affected by sex, presence of chronic bronchitis, use of oxygen therapy or concomitant COPD therapy. Long-term azithromycin therapy was found to be more effective in older patients (>65 years, relative hazard 0.59, 95% CI 0.57 to 0.74, p<0.01) and in patients who stopped smoking (ex-smoker relative hazard 0.65, 95% CI 0.55 to 0.77; smokers relative hazard 0.99, 95% CI 0.71 to 1.38; p value for interaction=0.03).

Severity of exacerbations on macrolide therapy
The impact of long-term macrolide therapy on the severity of exacerbations was not reported in many studies. Seemungal et al showed that macrolide use was associated with a lower median number of exacerbation days. The macrolide group reported median exacerbation days of 9 with IQR 6–13 days, and the placebo group had median of 13 exacerbation days with IQR 6–24 days (p=0.036). There is insufficient evidence to conclude what the impact on severity of AECOPD is in patients on long-term macrolide therapy.

Hospitalisation for AECOPD
Five studies involving 1424 patients reported on the number of hospitalisations due to acute exacerbations of COPD as secondary outcomes. Pooled data, as reported by Ni et al, did not demonstrate a significant reduction in hospital admissions in patients with AECOPD on macrolide therapy in comparison with the control group (RR 0.89, 95% CI 0.64 to 1.24, p=0.50). Albert et al found a non-significant trend towards reduction in the rate of hospital admissions using macrolides, whereas Uzun and colleagues found no significant delay in time to first hospitalisation in patients on long-term macrolide therapy.

The published evidence does not support the use of long-term macrolides in patients with COPD to reduce severity of exacerbations or requirement for hospitalisation due to exacerbation. The studies were underpowered and definition of severity of exacerbations not comparable with pool data.

Lung function tests and disease progression
The natural history of COPD is characterised by progressive airflow limitation and historically, FEV₁ decline has been considered the single most important marker of disease progression. However, data obtained in two different retrospective cohorts have shown that FEV₁ decline is not invariably progressive and FEV₁ is only weakly correlated with patient-related outcomes.

BTS Guideline
Exacerbation rate and number of exacerbations
Seven randomised control studies involving 1614 patients reported the number of acute exacerbations during the period of intervention. Long-term macrolide therapy was shown in a meta-analysis to significantly reduce acute exacerbations in the intervention group when compared with the comparison group (RR=0.7, 95% CI 0.56 to 0.87, p<0.01, I²=66.43%). The Suzuki et al 2001 trial was non-blinded and in appraisal of the study methodology, this raised concerns for bias. Analysis of the remaining six studies continued to show benefit for macrolide therapy over placebo, with a 20% relative risk reduction of AECOPD (RR=0.8, 95% CI 0.72 to 0.88, p<0.01).

Eight RCTs reported the rate of exacerbation per patient per year in a total of 1582 patients. The long-term macrolide group had a significant reduction in the rate of exacerbations (RR=0.58, 95% CI 0.43 to 0.78, p<0.01, I²=67.8%). The recent published ERS/ATS guideline pooled data for studies that included treatment for 12 months and excluded the Suzuki trial (due to risk of bias, non-blinded study as described above) and demonstrated a reduced rate of AECOPD (RR=0.76, 95% CI 0.68 to 0.86).

Randomised trials assessing the impact of long-term macrolide treatment in COPD have not reported on disease progression as a secondary outcome. Rate of decline in FEV₁, as single measure is not accurate in the measurement of disease progression, or in assessing the effect of therapeutic interventions in altering the progression of the disease process.107

Lung function was reported in four studies. Pooled analysis was not possible due to the varied ways in which results were reported. Banerjee et al reported no difference in the intervention and placebo groups after 12 weeks of clarithromycin in spirometry measurements or shuttle walk distance.83 Simpson et al reported no difference in FEV₁, reported as percentage predicted with 12 weeks of azithromycin 250 mg a day (30 patients, 15 in each arm).84 No difference in FEV₁ after 12 months’ treatment with erythromycin-study was withdrawn due to 250 mg or placebo was demonstrated by Seemungal and colleagues in patients with COPD.108 No significant changes were shown between groups in post-bronchodilator FVC, FEV₁ or 6 min walk test in patients on azithromycin 500 mg three times a week after 12 months.83

There is no evidence that long-term macrolide therapy results in improvements, or lessened decline, of lung function parameters in patients with COPD.

Safety and adverse events

Long-term treatment with macrolides has been associated with several adverse effects.108 The most common side effects relate to the gastrointestinal tract by stimulating gastrointestinal motility through motilin-like activity. Symptoms including anorexia, nausea, vomiting, diarrhoea and abdominal pain were reported in patients with COPD in the studies reviewed. Cessation of treatment due to gastrointestinal side effects were reported in the studies published by Banerjee et al, Suzuki et al, Seemungal et al, He et al and Blasi et al.96–98 101 102 Uzun published side-effect profiles, indicating that diarrhoea was the only event that was higher statistically significantly more frequent in the azithromycin group compared with placebo.83

Albert et al reported hearing impairment in 142 (25%) patients with COPD treated with azithromycin 250 mg a day and in 110 (20%) patients receiving placebo (p=0.04).91 A small, but significant difference was seen at 3-month review in audiometry measurements (in all four sound frequencies) in the treatment arm.93 Repeat audiology testing demonstrated improvement regardless if azithromycin was stopped or not. The authors postulate too stringent criteria for measurable deficit and over reporting of hearing impairment as explanation. None of the other studies reported on hearing impairment as an outcome. Erythromycin and clarithromycin are both associated with otoxicology that includes vertigo, hearing loss, deafness and tinnitus.109 A single patient receiving long-term erythromycin in the Seemungal et al study was withdrawn due to significant tinnitus.101

The recognised side-effect profile associated with macrolide therapy includes allergic reactions, skin eruptions, hepatotoxicity, cardiac arrhythmias and QTc prolongation.110–112 These side effects were not reported as adverse events in the studies reviewed based on the trial protocols and exclusion criteria (designed to exclude high-risk patients specifically with regards to cardiac toxicity). Three studies including 212 patients reported four cardiovascular events in the treatment arm and two in the control arm (p=0.43).106 97 107 Albert and colleagues reported the death rate due to cardiovascular events in both the azithromycin and placebo arm as 0.2%.

Pooled analysis of all nine RCTs reviewed demonstrated higher adverse events reported in the intervention group, with gastrointestinal side effects the most commonly reported (OR 1.55, 95% CI 1.00 to 2.39, p=0.05).84 93 94 96–98 101 102 Albert and colleagues reported only 11/558 patients in the treatment arm on azithromycin that had to stop the treatment due to GI side effects and 6/559 patients in the control group stopped placebo for the same reason.93

Antimicrobial resistance

Macrolide antibiotics have direct antibacterial activity against Gram-positive, Gram-negative and atypical bacteria such as Legionella spp, Mycoplasma and Chlamydia spp. Macrolides inhibit RNA-dependent protein synthesis by reversibly binding to the P-site on the 50S subunit of bacterial ribosomes and inhibit transpeptidation or translocation of nascent peptides.81

The severity and rate of AECOPD have been linked to both C. pneumoniae and Haemophilus influenzae.113–117 The antimicrobial activity of macrolide antibiotics with possible reduction in bacterial load may explain in part the effect on exacerbation rate seen in clinical trials in patients with COPD. The acquisition of macrolide resistance with long-term low-dose use of macrolide antibiotics remain uncertain in patients with COPD.

Seven studies monitored and reported on macrolide antibiotic susceptibility changes during treatment period.83 93 94 96–98 101 102

Albert et al found only 15% of patients could expectorate sputum after a 3-month period on macrolide therapy.93 Investigators used sputum in patients who could expectorate and nasopharyngeal swabs to report on colonisation and resistance. This study subsequently reported Staphylococcus aureus as the most common organism isolated (treatments group 60 (10.7%) vs placebo group 71 (12.7%)) likely due to nasopharyngeal sampling. Moraxella spp and S. pneumoniae were both isolated and did not show significant difference between treatment and placebo arms in the study. During the 12-month study period, significantly more patients in the placebo group became colonised (172 patients) versus 66 patients in the treatment arm on azithromycin 250 mg a day (p<0.001). Newly colonised patients in the treatment group demonstrated significantly higher resistance to macrolide antibiotics 81% compared with 41% in the placebo group (p<0.001).

Seemungal et al reported in 109 patients receiving erythromycin 250 mg twice a day over a 12-month period that only one patient developed resistance to S. pneumoniae. H. influenzae was isolated in 22 of the 109 patients; all of the isolates were found to be resistant to erythromycin.101

At baseline, Uzun et al obtained 22 sputum samples in the treatment group and 20 sputum samples in the placebo group. After a 12-month study period, 51 sputum samples in the treatment group and 57 sputum samples in the placebo group were analysed. This study also reported fewer new colonised patients in the treatment arm, 4 patients versus 12 patients (p=0.044). Macrolide resistance was reported in 6% (3 patients) receiving azithromycin 500 mg three times a week and in 24% (11 patients) receiving placebo for the 12-month period (p=0.036).83

Banerjee et al and He et al did not observe significant differences in colonisation rate or resistance to macrolide antibiotics, but these studies were shorter in duration, 3 and 6 months, respectively.96 97

In a single-centre, single-blind randomised placebo controlled trial, Brill et al demonstrated, after 3 months of azithromycin 250 mg three times a week, no difference in airway inflammation or bacterial load (0.42log₁₀ cfu/mL) in sputum when compared with placebo (0.08 CI 0.38 to 0.54).118
Mortality
In studies where mortality was specifically reported, no significant difference was observed between the control and macrolide groups (RR=0.9, 95% CI 0.48 to 1.69). There is therefore insufficient evidence to indicate a reduction in mortality with the use of long-term macrolides, but no evidence of increased mortality in relation to treatment.

Outcome weighting
The guideline group identified rate of AECOPD, time to first exacerbation, hospitalisation, serious adverse events and mortality as critical outcomes. Important outcomes identified include QOL, disease progression, lung function tests, airway inflammation and acquisition of macrolide resistance.

Evidence summary
Long-term macrolide treatment showed statistically significant improvement in QOL in patients with COPD as measured by SGRQ, but this does not reach the MCID—4 unit change. (Moderate)

Long-term macrolide antibiotics are effective in reducing the acute exacerbation rate in patients with COPD with high exacerbation rates (ie, more than three exacerbations per year, prolonged exacerbations or exacerbations resulting in hospitalisation). (Moderate)

Treatment courses of 12 months demonstrated the biggest effect size in reduction of exacerbation rate. (Low)

The number of hospitalisations was not significantly reduced in patients receiving long-term macrolide therapy. (Moderate)

No significant mortality benefit has been shown in 12-month follow-up studies using macrolide therapy, and there is insufficient evidence to demonstrate the effect of long-term macrolide therapy on the mortality in patients with COPD. (Low)

There is no evidence to suggest that long-term macrolide therapy impacts on disease progression or spirometry measurements and exercise capacity measurements. (Low)

Insufficient evidence is available on airway colonisation and acquisition of macrolide resistance in patients with COPD receiving long-term, low-dose macrolide therapy. (Moderate)

Recommendations

- Long-term macrolide therapy could be considered for patients with COPD with more than three acute exacerbations requiring steroid therapy and at least one exacerbation requiring hospital admission per year to reduce exacerbation rate. (Conditional)
- Long-term macrolide therapy could be considered for a minimum of 6 months and up to 12 months to assess the impact on exacerbation rate. (Conditional)

Good practice points

- Non-pharmacological and pharmacological therapies should be optimised prior to considering long-term macrolide therapy. This includes smoking cessation, optimised inhaler technique, optimised self-management care plan, airway clearance techniques and attendance at pulmonary rehabilitation courses.

- Macrolides should only be started following discussion and shared decision-making between the patient and a respiratory specialist.
- For safety purposes, an ECG should be performed prior to initiation of macrolide therapy to assess QTc interval. If QTc is >450 ms for men and >470 ms for women, this is considered a contraindication to initiating macrolide therapy. Baseline liver function tests should also be measured.
- Patients should be counselled about potential adverse effects before starting therapy including gastrointestinal upset, hearing and balance disturbance, cardiac effects and microbiological resistance.
- Microbiological assessment of sputum should be performed before therapy, including investigation for NTM. Macrolide monotherapy should be avoided if an NTM is identified. Repeat assessments are recommended with clinical decline or during exacerbations to monitor resistance patterns.
- Accurate assessment of baseline exacerbation rate should be determined before starting long-term macrolides for patients with COPD and a CT scan should be considered to exclude a possible diagnosis of bronchiectasis.
- A risk:benefit profile should be considered in each individual if significant side effects from oral macrolide therapy develop. If gastrointestinal side effects occur at the higher dose of azithromycin (500 mg thrice weekly), a dose reduction to azithromycin 250 mg thrice weekly could be considered if macrolide therapy has been of clinical benefit.
- Liver function tests should be checked 1 month after starting treatment and then every 6 months. An ECG should be performed 1 month after starting treatment to check for new QTc prolongation. If present, treatment should be stopped.
- Subsequent follow-up at 6 and 12 months should determine whether benefit is being derived from therapy by using objective measures such as the exacerbation rate, CAT score or QOL as measured by a validated assessment tool such as SGRQ. If there is no benefit, treatment should be stopped.
- It is not necessary to stop prophylactic azithromycin during an acute exacerbation of COPD unless another antibiotic with potential to affect the QT interval has also been prescribed.

See quick reference guide in online supplementary file 1.

Research recommendations

Long-term (>12 months) follow-up trials to evaluate impact of long-term macrolide therapy on mortality, antimicrobial resistance, long-term potential cardiac toxicity and disease progression.

Studies to evaluate the impact of short-term breaks in chronic therapy with long-term macrolide antibiotics are needed.

Studies phenotyping COPD in large trials where subgroup analysis can potentially identify groups of patients with COPD who will benefit most from long-term macrolide therapy.

Trials investigating head to head the benefits and adverse effects of oral agents that reduce acute exacerbations in patients with COPD (long-term, low-dose macrolides, carbo-cysteine, rolumlustil).

SECTION 7: BRONchiOLITIS OBLITERANS (INClUding POST TRANSPLANTATION)

Introduction
Bronchiolitis obliterans is a condition characterised by subepithelial inflammatory and fibrotic narrowing of small airways
within the lung. It occurs following a variety of insults to the lung such as infections or inhaled toxins. It also occurs in the context of systemic diseases such as rheumatoid arthritis and other connective tissue diseases. It can occur as a complication of transplantation of lung or haematopoietic stem cells (HSCT). The histological correlate is obliterator bronchiolitis or constrictive bronchiolitis.

In the context of lung transplantation, bronchiolitis obliterans causing lung function impairment without histological confirmation has been called bronchiolitis obliterans syndrome (BOS). A standard definition of BOS was first developed in 1993 based on changes in FEV, following transplantation. The definition has undergone subsequent revisions in 2002 and in 2014 when the over-arching term of chronic lung allograft dysfunction (CLAD) with subdivisions of restrictive CLAD (restrictive allograft syndrome, RAS) and obstructive CLAD (BOS) were proposed. BOS is common after lung transplantation occurring in over 50% of those surviving more than 5 years. It is less common after HSCT, between 2% and 14% in the first 5 years post transplant. Long-term macrolides have been used both to prevent BOS and to treat BOS when it occurs. Several observational studies have noted a positive response to azithromycin in a proportion of patients with BOS following lung transplantation, especially those with neutrophilic bronchoalveolar lavage results. This has been labelled neutrophilic reversible allograft dysfunction or azithromycin responsive allograft dysfunction distinguishing it from the non-reversible CLAD.

Evidence base
There is a significant body of literature relating to bronchiolitis obliterans occurring in the context of lung or HSCT. In the lung transplantation field, the majority of published studies are case series. There is marked variation between studies in patient groups, concomitant immunosuppressive regimens, choice of macrolide and dosing regimens, and choice of outcome measures making meaningful systematic analysis of the data impossible. In patients with bronchiolitis obliterans complicating HSCT, there are fewer studies, predominantly case series, also with a high degree of variation in populations, background immunosuppressive regimens, macrolide interventions (often in conjunction with other novel therapeutic agents) and outcome measures precluding meaningful systematic analysis.

Macrolides in BOS/CLAD following lung transplantation
An International Society for Heart and Lung Transplantation/American Thoracic Society (ATS)/European Respiratory Society (ERS) clinical practice guideline on the diagnosis and management of BOS was published in 2014 following the GRADE approach to develop specific clinical recommendations including the use of azithromycin. This guideline identified 10 studies (one observational study and nine case series) that described the effects of azithromycin on the lung function of lung transplant patients with BOS. It also identified two observational studies that described the effect of azithromycin on mortality in lung transplant patients with BOS. The guideline recommended a trial of azithromycin (250 mg orally daily for 5 days and then thrice weekly for a minimum of 3 months) for lung transplant recipients who developed a decline in FEV, consistent with the onset of BOS. The recommendation was conditional.

There have been two RCTs investigating the role of macrolides in BOS following lung transplantation. Vos and colleagues examined the effect of azithromycin versus placebo for the prevention of BOS following lung transplantation. Corris et al studied the effect of azithromycin versus placebo in the treatment of patients who developed BOS following lung transplantation. In the Vos study, patients who had undergone lung transplantation were randomised to receive either azithromycin, 250 mg three times a week (40 patients) or placebo (43 patients) for 2 years in a fully blinded study. Primary end points were freedom from BOS and survival at 2 years. If patients developed BOS on placebo, they were switched to open-label ‘rescue’ treatment with azithromycin. Patients receiving prophylactic azithromycin were less likely to develop BOS than those on placebo (12.5% vs 44.2%, p=0.0017). Overall survival was comparable between groups. This group of investigators have subsequently published a post hoc analysis of the same patients who were treated for a further year applying the newer CLAD classification system over an extended 7-year period of observation. The treated patients were less likely to develop CLAD at any stage compared with the placebo patients (28% vs 51%, p=0.043). There were also benefits in CLAD-free survival, long-term pulmonary function and functional exercise capacity, but graft loss (re-transplantation and mortality) was similar in both groups (53% vs 40%, p=0.27). In the Corris et al paper, 46 patients with BOS following lung transplantation were randomised to receive either azithromycin, 250 mg on alternate days (23 patients) or placebo (23 patients) over a 12-week study period in double-blind fashion. The primary end point was change in FEV, at 12 weeks. Patients who had a rapid and severe deterioration in lung function were withdrawn from the study and received open-labelled azithromycin. On an intention-to-treat (ITT) analysis, there was no significant difference between treatments in the primary outcome (0.035 L, p=0.6). For study completers (16 azithromycin and 17 placebo), there was a significant difference of 0.278 L (p=0.001). In addition, 9/23 ITT patients in the azithromycin group had a ≥10% gain in FEV, from baseline while no patient in the placebo group had ≥10% gain in FEV, while on placebo (p=0.002). This latter observation, previously noted in other studies, suggests a subgroup of BOS patients who exhibit a response to azithromycin and is one of the drivers for recommending the re-classification of CLAD following lung transplantation noted above.

Evidence summary
The evidence for macrolides in BOS following lung transplantation covers both prophylactic use to prevent BOS and treatment following the occurrence of BOS. In both areas, there are single RCTs and a number of case series. The overall quality of the evidence is at best modest; however, BOS is a devastating complication of lung transplantation, so any intervention that offers the chance of prevention, reversal or stabilisation is welcome. Long-term macrolide use is a low-risk intervention. On this basis, we can make two recommendations. (Low)

Recommendations
- Low-dose, long-term azithromycin (250 mg thrice weekly) could be considered to prevent the occurrence of BOS post lung transplantation. (Conditional)
- Low-dose azithromycin (250 mg alternate days for a trial period of 3 months) could be considered to treat BOS occurring in lung transplant recipients. (Conditional)

Macrolides in BOS following HSCT
There are a number of case series and one RCT examining the role of macrolides (sometimes in combination with other emerging therapies) in BOS following HSCT. Khalid et al reported prospectively from Saudi Arabia on a series of eight
HSCT patients with BOS diagnosed on the basis of lung function and high-resolution CT scanning who were treated with azithromycin for 12 weeks. All patients showed improvement in FEV₁ and all but one in FVC following treatment. The changes were statistically (FEV₁, p=0.0052; FVC, p=0.0067) and clinically (FEV₁, 21.6%; FVC 20.6%) significant. Norman et al described a retrospective, single-centre series of eight HSCT patients in North America with newly diagnosed BOS who were treated with a combination of fluticasone (inhaled), azithromycin and montelukast (the FAM regimen) in addition to a more rapid taper in systemic corticosteroids than was normal practice. The intention was to reduce systemic corticosteroid exposure in these patients. Comparison with historical controls (HSCT patients with BOS treated with much higher overall doses of systemic steroids) suggested equivalent efficacy in maintaining lung function. Jo et al from South Korea, in a retrospective study, described the effect of prophylactic azithromycin in reducing the incidence of BOS in a series of 100 HSCT patients compared with over 1000 similar patients who did not receive prophylactic azithromycin. The incidence of BOS in those who received azithromycin was 12% compared with 6.4% in those who did not. Subsequent multivariate analysis did not suggest that prophylactic azithromycin was associated with the development of BOS. In a multicentre open-labelled prospective study, Williams et al examined the effect of the FAM regimen on non-BSCT patients. The primary end point was treatment failure by 3 months defined as an absolute decline in % predicted FEV₁ of > or =10%. Thirty-six patients from 10 centres were studied. Two patients (6%) had treatment failure compared with 40% of historical controls. The single-centre RCT from Lam et al in Hong Kong compared the effect of 12 weeks’ treatment with daily azithromycin, 250 mg, compared with placebo in 22 HSCT patients with BOS. Primary outcomes were changes in QOL (SGRQ) and lung function (FEV₁ and FEF 25–75) measured monthly and 1 month after completion of treatment. The authors reported no change in lung function or SGRQ although the QOL data were analysed in an unusual fashion.

### Evidence summary

The literature for macrolides in BOS following HSCT is limited. There is only one RCT which has significant methodological limitations. The majority of case series are small and half are retrospective. There are differing diagnostic criteria for BOS and in some studies multiple drugs including azithromycin are used. Several authors comment on the difficulty of performing studies in this patient group where the condition is a rare complication of HSCT (making it difficult to gather high patient numbers) with devastating consequences (making it difficult to justify and recruit to a placebo limb). This situation is discussed in detail in an editorial. There is insufficient evidence for us to make a recommendation.

### Macrolides for other causes of bronchiolitis obliterans

There are no published studies looking at macrolide treatment for other causes of bronchiolitis obliterans. There is a single case report of bronchiolitis obliterans following dust exposure on 9/11 with response to azithromycin and one recent review article suggested a possible role for macrolides in the treatment of bronchiolitis obliterans associated with rheumatoid arthritis.
Hodgson et al demonstrated no significant difference between placebo and intervention in LCQ, nor for any secondary outcome measurements. A subgroup analysis was conducted (post hoc), identifying a subgroup of responders to azithromycin therapy (i.e., those with a concomitant diagnosis of asthma, mean LCQ improvement compared with placebo 5.77; 95% CI 2.75 to 8.79). However, given the small size of this group (18 patients in total) and post hoc nature of the subgroup, it is highly likely that this analysis suffers with bias.146

Side effects and adverse events
Side-effect profiles were published in both trials; Yousaf et al reported two patient withdrawals on erythromycin 250 mg per day for 12 weeks, one of whom reported dizziness which resolved.147 Hodgson et al reported one withdrawal on azithromycin due to GI side effects. In the study as a whole, there were no statistically significant differences in side effects between intervention and control group (eight patients with GI side effects in the intervention group vs five in the placebo group).148

Evidence statements
Long-term macrolide antibiotics are ineffective in improving any outcomes in chronic cough, accepting that only two small RCTs have been conducted (totalling only 72 patients). These outcomes include those proven to be of importance in chronic cough such as the LCQ, number of coughs and 100 mmVAS for cough. (Low)

Recommendations
- Long-term macrolide antibiotics should not be used to manage patients with unexplained chronic cough. (Conditional)

Research recommendation
High-quality, adequately powered randomised trials are needed to assess the effect of macrolides on outcomes of importance in chronic cough.

Organising pneumonia
Organising pneumonia (OP) is a clinicopathological syndrome associated with characteristic patterns on lung imaging and biopsy which represents an aberrant healing response to injury within the small airways and alveolar spaces. CT scanning typically shows patchy consolidation with air bronchograms, typically in subpleural locations, which may appear migratory on serial imaging. Ground glass change, reversed halo (atoll) sign, reticulation or nodules may also be seen. Histological features of OP include fibroblasts and inflammatory cells embedded in extracellular matrix within the small airways and alveoli, often forming pseudocysts masses known as Masson bodies. OP can occur in a variety of conditions including infection, connective tissue disease, malignancy, following radiotherapy, drug reactions and immunodeficiency. If no cause can be found on exhaustive testing the term cryptogenic organising pneumonia (COP) (previously called bronchiolitis obliterans organising pneumonia) is used. COP is a rare disease (annual incidence ~1:100 000) that presents with breathlessness and cough over weeks or months, often accompanied by fever, myalgia and elevated blood inflammatory markers. Lung function tests typically show restriction and impaired gas transfer.

The prognosis of COP is good, particularly when consolidation is the primary pattern on CT. The natural history is spontaneous remission in many untreated cases.149 In non-remitting or progressive COP, oral corticosteroids have been used often in reported case series.150 There have been no placebo-controlled trials of steroid therapy, but is believed that there is an autoimmune aetiology in a proportion of cases and that steroids speed resolution. In steroid-responsive cases, relapse may occur after stopping therapy.

Evidence summary
The evidence base for macrolide therapy in OP consists of case reports and small case series totalling fewer than 50 patients.151–161 Macrolides were used either first-line in untreated patients or less commonly, as an add-on therapy with oral steroids. When used first-line, no studies described an initial observation period to identify self-resolving disease. Clinical outcome reporting is variable, but overall the published studies report clinical improvement in response to macrolide therapy.162 The risk of publication bias is high. Because the natural history of untreated COP is often self-resolution, in the absence of placebo-controlled trials, no conclusions can be made about the efficacy of macrolide therapy.

There is insufficient evidence to make a recommendation.

Diffuse panbronchiolitis
Diffuse panbronchiolitis (DPB) was defined as a specific clinicopathological entity in the 1960s, being characterised histologically by multiple micronodular lesions consisting of chronic inflammatory cells infiltrating bronchiolar walls.163 164 It has almost exclusively been reported in East Asian countries, notably Japan, but also others such as China and South Korea.163 Relatively few cases have been diagnosed in Caucasian and other populations.163 165 166

DPB was one of the first respiratory conditions for which it was identified that a regular macrolide, erythromycin, at a lower dose than usually used in therapy, could be associated with notably improved outcomes for sufferers. This was discovered in the early 1980s, although there has been a paucity of published rigorous and well-designed placebo-controlled trials subsequently to test the hypothesis, with further supporting evidence largely in the form of (usually small) case series and observational studies.161 167 The largest published study was retrospective, involving nearly 500 Japanese patients with DPB.168 These patients were divided into three cohorts on the basis of likely date of diagnosis, which were then compared with respect to outcome. There was a statistically significant survival advantage (p<0.0001) to being in the third cohort and thereby diagnosed between 1985 and 1990. This was ascribed as being directly related to when the beneficial role of low-dose erythromycin in DPB had first been reported.168 To provide support for this, the records of this third cohort of 87 patients were analysed in more detail—and there was a significantly higher survival rate at 5 years of those who were known to have received erythromycin compared with those who had not (p=0.01).168 Moreover, those who had not received erythromycin had a similar 5-year survival rate to those in the first cohort, who had been diagnosed between 1970 and 1979, and therefore before the first reports of benefit with such a strategy.163 168 DPB case series studies have also shown the effect of low-dose macrolide therapy in controlling clinical symptoms, such as by reducing sputum production, and objectively improving lung function.165

The benefits of using low-dose macrolide therapy in the setting of DPB have been endorsed by clinical guidelines developed in the Far East.161 However, due to the current rarity of this condition in the UK, the GDG felt that it was not warranted for BTS to conduct a comprehensive evidence-based review and
SECTION 9: SAFETY ISSUES

Gastrointestinal effects

Gastrointestinal (GI) side effects of macrolides have been recognised since the introduction of erythromycin in 1952. Although initially thought to be due to alterations in gut flora, it is now clear that macrolides act through interactions with motilin receptors in the gut. In humans, macrolids potentiate gastric and small bowel motility, increase lower oesophageal sphincter pressure, and influence colonic transit and gall bladder function.

The prokinetic effects of macrolides on the GI tract has led to their therapeutic use in conditions characterised by reduced GI motility including diabetic gastroparesis, anorexia nervosa, colonic pseudo-obstruction, postoperative ileus and in critical care patients.

GI symptoms are the most commonly reported side effects of macrolides occurring in up to 70% of patients taking erythromycin, less commonly with clarithromycin and azithromycin. Symptoms include nausea, vomiting, abdominal pain, diarrhoea and anorexia. Commenting on 11 studies of long-term, low-dose macrolide treatment in various chronic respiratory diseases in nearly 1000 children and adults, Altenburg et al found that mild to moderate GI complaints had been reported which hardly ever caused study drug discontinuation.

Thus, although GI symptoms are common and predictable, they are rarely sufficiently troublesome to necessitate discontinuation of therapy. Dose reduction may improve tolerability.

Good practice points

✓ Prior to initiating low-dose macrolide therapy, patients should be warned of the possibility of GI side effects.
✓ GI side effects may be ameliorated by dose reduction although this may also reduce clinical efficacy.
✓ Clinicians should carefully consider the risk-to-benefit balance when considering therapy for those with pre-existing GI symptomatology.

Cardiac effects

Macrolide antibiotics, along with a wide range of other pharmacological agents, have the potential to interfere with conduction in cardiac tissue and hence to cause dangerous cardiac arrhythmias and sudden cardiac death. Macrolides can both directly prolong the QT interval and also inhibit the metabolism of other proarrhythmogenic drugs by acting on cytochrome P450 in the liver.

In recent years, there have been reports of an increased risk of cardiovascular death in patients treated with short, oral antibacterial courses of azithromycin and of no increased risk of death in a younger group of similarly treated patients. Discussion of these seemingly contradictory findings commented that the group of patients with increased risk exhibited a high level of baseline risk and comorbidity. Factors increasing the risk of malignant arrhythmias with macrolides are well recognised and include age >80, female gender, heart disease, use of other QT prolonging medication, reduced drug elimination, bradycardia, prolonged QT interval before therapy and genetic predisposition. These events are rare; 85 deaths for each 1 million courses prescribed in the increased risk group described above, so it is not surprising that such events do not appear in the literature of long-term, low-dose macrolide use.

A helpful review article for the American COPD Clinical Research Network describes a pragmatic approach to risk stratification when considering long-term, low-dose macrolide therapy. The authors combined history taking targeting evidence of heart failure, episodes of hypokalaemia, a family history of long QT syndrome, or use of other medications known to prolong the QT interval together with a pre-treatment ECG and an ECG 1 month after starting therapy. With this approach in a COPD population entering a macrolide trial, 7.1% were excluded on history and 1.6% because of a prolonged baseline QTc. A further 0.8% were withdrawn after developing QTc prolongation with treatment or on placebo. Using such a strategy, exclusions in asthma and bronchiectasis populations are likely to be fewer.

Good practice points

✓ Prior to initiating low-dose macrolide therapy, patients should be asked if they have a history of heart disease, previous low serum potassium measurements, a slow pulse rate, a family history of sudden death or known prolonged QT interval. Patients with such a history should not receive low-dose macrolide therapy without careful consideration and counselling of the increased risk of adverse cardiac effects.
✓ Prior to initiating low-dose macrolide therapy, a drug history looking for agents that might prolong the QTc interval should be sought (see online supplementary appendices 3 and 4). Patients taking such agents should not receive low-dose macrolide therapy.
✓ Prior to initiating low-dose macrolide therapy, an ECG should be performed to exclude a prolonged QTc interval defined as >450 ms for men and >470 ms for women (see methodology in online supplementary appendices 3 and 4). Patients with a prolonged QTc interval should not receive low-dose macrolides.
✓ One month after initiating low-dose macrolide therapy, a second ECG should be performed to exclude the development of a prolonged QTc interval. Patients who develop a prolonged QTc interval on low-dose macrolides should stop the macrolide.
✓ If any new drug that could potentially prolong QTc time is started or if dose increases are made repeat ECG assessment.

Otoxicity

Macrolide ototoxicity is well recognised usually causing a dose-dependent, reversible, sensorineural hearing loss (SNHL). The only systematic review of this topic was published recently, including data from three prospective studies. In these studies including 93 patients, 14 (15%) developed SNHL confirmed by audiometry following exposure to macrolide antibiotics. All but one recovered on cessation of the drug. Earlier studies of long-term macrolide use for the treatment of mycobacterial infections have also reported reversibility of ototoxicity following either withdrawal or dose reduction of the macrolide. Albert reported hearing decrements in 25% of a COPD population (vs 20% in the placebo group) with normal hearing at the outset of 1 year of low-dose azithromycin. Hearing subsequently improved in 34% who discontinued treatment, but also in 32% who continued due to protocol errors. Similar proportions of placebo-associated hearing loss also improved with or without drug withdrawal. The authors of this study felt in discussion that their criteria for defining hearing loss were too stringent and that the incidence of hearing decrements were overestimated.
Overall, the literature suggests that for low-dose, long-term macrolide use the incidence of ototoxicity is rare and almost always reversible. It would seem sensible to assess potential recipients for hearing impairment or difficulties with balance prior to initiating therapy.

**Good practice point**
- Prior to initiating low-dose macrolide therapy patients should be asked if they have a history of hearing or balance difficulties. Such patients should be made aware of the potential for a further, almost always reversible, deterioration in hearing or balance with macrolide therapy. Patients with pre-existing hearing or balance difficulties who wish to proceed with treatment should be asked to report any change in hearing or balance promptly.

**Other side effects**
A low rate (1%–5%) of asymptomatic elevation of serum aminotransaminase levels is known to occur with any of the four orally absorbed macrolide antibiotics. The elevation is generally mild to moderate in degree and rarely requires dose modification or discontinuation. More seriously, a cholestatic picture can occur which carries a higher risk of permanent liver damage.\(^\text{194,195}\)

Monitoring liver function at the start of therapy seems a sensible precaution.

**Good practice points**
- Prior to initiating low-dose macrolide therapy, baseline liver function tests (LFTs) should be checked.
- LFTs should be checked after 1 month of treatment and then every 6 months thereafter for the duration of therapy.

**Prescribing off-label**
It is important to be aware that none of the macrolide antibiotics have a product license in the UK for the indication of long-term, low-dose usage as immunomodulatory agents. Their prescription in this context is thus off label. Clinicians should make patients aware of this and follow GMC guidance in this area which states that “decisions should be made in collaboration with the patient by discussing the options with them and ensuring that they have sufficient information about the medicine to allow them to make an informed decision”. An example of an information leaflet used to support such discussions is shown in online appendix 4.

**SECTION 10: ANTIMICROBIAL RESISTANCE**
Antimicrobial agents are different to other medications, as they can be associated with undesirable consequences not only in the patient receiving the treatment, but also for other individuals - both at the time and potentially at much later dates in the future. Exposure of bacteria to an antibiotic may be associated with subsequent alterations in those bacteria such that they become resistant to its effect. This means that the antibiotic loses its therapeutic value. If the antibiotic exposure, that is, with its associated selection pressure, is stopped such resistant bacteria do not necessarily revert back to being susceptible in future. The encoding mechanism of the bacterial resistance to an antibiotic’s effect may be located on mobile genetic elements, such as plasmids or transposons, which will then be inherited by the future progeny of that bacterium, and may also be transferred to other strains of the same species, and indeed to other species. Such mobile genetic elements may carry the gene(s) encoding resistance to the specific antimicrobial agent, or its class, being used; and also resistance mechanisms to other, completely unrelated, antimicrobial agents/classes—that is, conferring resistance to multiple different agents. These resistant bacterial strains can then be transmitted to other individuals or establish persisting reservoirs in the environment. The resistance mechanism does not need to evolve in the pathogenic organism(s) primarily being targeted by the antimicrobial treatment. Other bacterial species, of low or no virulence, may develop such mechanisms to survive in their antibiotic-exposed milieu, such as the GI or upper respiratory tracts, but may then transmit the means to other, more virulent, species. The rise in antimicrobial resistance is of national and international concern.\(^\text{196,197}\) Not all bacteria are able to develop resistance to all antimicrobials to which they are naturally susceptible. However, this can then lead to the disappearance of some non-pathogenic commensals, ‘good’, bacterial species in any exposed microbiome; and their replacement by other organisms which are resistant, either intrinsically or by acquisition, which in turn may lead to subsequent infections. Such a shift in pathogens could include metacinillin-resistant *Staphylococcus aureus*, Clostridium difficile, Gram-negative bacilli and yeasts. These ecologically adverse effects have been termed the ‘collateral damage’ associated with antibiotic use.\(^\text{198}\)

The macrolides, such as erythromycin, clarithromycin and azithromycin, inhibit RNA-dependent protein synthesis by bacteria by binding to the 50S ribosomal subunit.\(^\text{199}\) Various mechanisms by which bacteria may resist their action have been elucidated, with resistance often being cross-class.\(^\text{199}\) Many Gram-negative bacteria, such as Enterobacteriaceae (‘coli-forms’), *Pseudomonas* and *Acinetobacter* species, show reduced permeability of their outer cell envelope to macrolides. Azithromycin is more active in vitro against a range of Gram-negative bacteria than erythromycin. This is thought most likely due to the molecule being able to penetrate such bacterial cells more effectively.\(^\text{200}\) Organisms, such as *S. aureus* and coagulase-negative staphylococci, may have plasmid-mediated macrolide resistance by increased efflux.\(^\text{201}\) The 50S target site may be altered, reducing the binding affinity.\(^\text{202}\) This has been reported in a wide range of potential pathogens, including *Streptococcus pneumoniae*, *Streptococcus pyogenes*, *S. aureus*, *Helicobacter pylori* and *Mycobacterium avium* complex.\(^\text{203,204}\) The genetic mutation is mediated by *erm* genes, located either on transposons in the bacterial chromosome or on plasmids. Such resistance may also be associated with resistance to lincosamides such as clindamycin. Resistance may also be conferred by enzymatic inactivation, such as by phosphotransferases in *S. aureus*, *Escherichia coli* and *Nocardia* species.\(^\text{199}\)

Rising bacterial resistance rates to macrolides have been reported globally. For *S. pneumoniae* isolates, it can vary markedly between countries, ranging from less than 10% to over 90%.\(^\text{203}\) Extensive macrolide use provides a strong selective pressure for the spread of macrolide resistance in pneumococci.\(^\text{203,204}\) There is a notable trend for *S. pneumoniae* isolates, which are resistant to penicillin, also to be resistant to macrolides—in the USA, there are reports of 30% of *S. pneumoniae* isolates overall being erythromycin resistant, but virtually 70% of high-level penicillin-resistant isolates also exhibiting erythromycin resistance.\(^\text{199,205}\) However, it should be recognised that the increasing use of pneumococcal conjugate vaccines, which include serotypes commonly associated with macrolide resistance such as serotype 14 in the 7-valent and also serotype 19A in the 13-valent, can then be followed by marked shifts in the prevalence of macrolide-resistant *S. pneumoniae* isolates.\(^\text{203,206}\) *S. pyogenes* isolates showing resistance to erythromycin were first reported in the UK in 1955, with resistance rates rising in the 1980s–1990s.\(^\text{199,207}\) However, reduction in resistance rates with reduced macrolide usage has also been found. For instance,
Finland likewise experienced a rise in macrolide resistance in the 1980s, which was temporally associated with a rise in macrolide use; however, after nationwide policies were introduced to restrict their use in treating respiratory and soft-tissue infections after 1991, the resistance rate fell in clinical isolates of *S. pyogenes* from 16.5% in 1992 to 8.6% in 1996. Similar to *S. pneumoniae*, erythromycin resistance rates are much higher in *S. aureus* isolates that are meticillin resistant compared with those that are meticillin susceptible. The emergence of erythromycin resistance in *S. aureus*, isolated from individual patients on macrolide treatment, has been known since the discovery of the antibiotic.

As to be expected, increased macrolide resistance has been found in studies evaluating the impact of long-term macrolide use in the setting of chronic lung diseases. A recent meta-analysis of azithromycin use found that, while overall colonisation rates by potential pathogens, such as *S. pneumoniae*, *Haemophilus influenzae* and *Moraxella catarrhalis*, decreased, the risk of macrolide resistance, macrolide resistance among such pathogens rose 2.7-fold in patients receiving azithromycin when compared with placebo. However, it should also be acknowledged that for any individual patient, the actual clinical impact of the presence of a macrolide-resistant respiratory pathogen, such as *S. pneumoniae*, at least in the short term, is not clear.

Organisms for which the macrolides are a key pillar of treatment, but in which macrolide resistance can emerge, notably if any macrolide treatment is not accompanied by two or three other active agents in combination, is of particular concern in the use of long-term (ie, months to years) courses of macrolides. Such pathogens include many non-tuberculous mycobacterial (NTM) species, notably the slower growing such as *M. avium* complex. Such organisms may develop resistance by single point mutations, for example, in the 23S rRNA gene. It is well recognised that the long-term success rate of medically treating macrolide-resistant NTM disease is very poor, and much worse than for treating a susceptible organism with a combination including an effective macrolide. Current NTM infection should be managed with reference to the BTS Guidelines for the management of non-tuberculous mycobacterial pulmonary disease.

The macrolides, notably azithromycin, also have significant applications outside the respiratory setting—including in regards to some sexually transmitted diseases (STDs), GI infections and zoonoses. One needs to be mindful that their value in such indications is affected by macrolide resistance and also by the rising bacterial resistance rates to other antibiotic classes. Circulating strains of *Neisseria gonorrhoeae* now show resistance to a range of antibiotic classes, to which this organism was initially susceptible. In consequence, the recent standard first-line regimen for suspected gonorrhoea has been ceftriaxone and azithromycin; however clinical isolates of *N. gonorrhoeae* also resistant to both these agents have now been described. The latest UK guidelines, published January 2019, now recommend therapy with ceftriaxone alone as first choice for cases where organism susceptibilities are not known, partly on the grounds of concern of rising macrolide resistance, both among *N. gonorrhoeae* but also in other organisms causing STDs.

Azithromycin is also a recommended first-line therapy for other STDs such as chlamydia (*Chlamydia trachomatis*), chancroid (*Haemophilus ducreyi*) and donovanosis (*Klebsiella granulomatis*). Rising fluoroquinolone resistance rates means that macrolides are the empirical treatment of choice for diarrhoea due to *Campylobacter*; and, in the form of azithromycin, a first-line option for enteric fever acquired in Asia.

Any indications for the macrolide class of antibiotics have been classified, in the main, in the ‘Watch’ group of antibiotics by WHO; along with six other classes. This categorisation consists of three groups: ‘Access’, ‘Watch’ and ‘Restrict’, the AWaRe classification. The uses of antibiotics in the Watch group should be carefully monitored to ensure they are in accordance with recommended indications. It should be recognised that the available evidence for the potential benefits and harms of long-term macrolide therapy will not capture all the associated risks of increasing antimicrobial resistance. Such studies are not designed to do so, and are too short in timescale and too restricted in patients, and organisms, being studied.

Virtually any antibiotic may be associated with diarrhoea; notably if by precipitating *C. difficile* disease. Although the macrolides appear relatively rare among antibiotic classes as being directly associated with *C. difficile*; clearly if their prior use has led to a pathogen shift, then use of an alternative antibiotic may be required which is more often associated with subsequent *C. difficile* disease.

Antibiotics have been shown to reduce the microbial diversity and alter composition of microbiomes in murine and human studies. Such effects may be greater, and more prolonged, with macrolides than for some other antibiotics classes, such as penicillins. In children at least, receipt of macrolides, and the associated changes in microbial commensal flora, have been linked with other health outcomes, such as asthma and being overweight. This illustrates that if there are identified benefit(s) to offering certain patients with specific chronic respiratory conditions, a macrolide as long-term supportive therapy; the known, and as yet unknown, risks of associated collateral damage should also be carefully considered, by both macrolide prescriber and macrolide consumer.

**Good practice points**

- The risks associated with increasing antimicrobial resistance should be discussed with patients prior to starting low dose macrolide therapy. Patients should understand the risk that there may not be an effective antibiotic for them, or someone else, when needed in the future.
- Prior to initiating low-dose macrolide monotherapy, patients should be asked if they have a history of previous or current *C. difficile* infection or disease. Current *C. difficile* infection should be managed with reference to BTS guidance and precludes low-dose macrolide monotherapy. Successfully treated *C. difficile* infection should not preclude low-dose macrolide monotherapy.
- If there is any clinical suspicion of possible *NTM* disease, patients should be screened via examination of sputum samples prior to starting therapy. If positive for recognised potential pathogenic species, low-dose macrolide prophylaxis is contraindicated.

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**Disclaimer** Healthcare providers need to use clinical judgement, knowledge and expertise when deciding whether it is appropriate to apply recommendations for the management of patients. The recommendations cited here are a guide and may not be appropriate for use in all situations. The guidance provided does not override the responsibility of healthcare professionals to make decisions appropriate to the circumstances of each patient, in consultation with the patient and/or their guardian or carer.

Competing interests BTS Declarations of Interest forms have been completed by all members for each year they partook in the GDG. Details of these forms can be obtained from BTS Head Office.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

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Supplementary 1: Quick reference guide for azithromycin

<table>
<thead>
<tr>
<th>Asthma</th>
<th>COPD</th>
<th>Bronchiectasis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Identify if suitable for Azithromycin therapy</strong></td>
<td><strong>Identify Contra-indications to macrolide therapy</strong></td>
<td><strong>Perform safety checks before starting therapy</strong></td>
</tr>
<tr>
<td>Confirmed diagnosis of asthma</td>
<td>Confirmed diagnosis of COPD</td>
<td>Confirmed diagnosis of bronchiectasis</td>
</tr>
<tr>
<td>Symptomatic despite &gt;800mcg/BED</td>
<td>3 or more exacerbations in previous 12 months OR</td>
<td>3 or more exacerbations in previous 12 months OR</td>
</tr>
<tr>
<td>At least 1 exacerbation in previous 12 months</td>
<td>1 or more severe exacerbation with hospitalisation/morbidity</td>
<td>1 or more severe exacerbation with hospitalisation/morbidity</td>
</tr>
<tr>
<td>Inhaled therapies optimised including inhaler technique and adherence review</td>
<td>Inhaled therapies optimised including inhaler technique and adherence review, smoking cessation and pulmonary rehabilitation completed</td>
<td>Optimisation of other interventions such as airway clearance and pulmonary rehabilitation</td>
</tr>
<tr>
<td><strong>Baseline ECG</strong></td>
<td><strong>Standard sputum for baseline culture if able to expectorate</strong></td>
<td><strong>Review concomitant medications for potential interactions</strong></td>
</tr>
<tr>
<td>If QTc prolonged (&gt;450msec for men, &gt;470msec for women) do not give macrolide</td>
<td>If bronchiectatic or clinical concern of NTM infection investigate to exclude (following BTS guideline on NTM disease)</td>
<td>Review concomitant medications for potential interactions</td>
</tr>
<tr>
<td><strong>Start Azithromycin therapy</strong></td>
<td><strong>Monitoring during therapy</strong></td>
<td><strong>Review therapy at 6-12 months</strong></td>
</tr>
<tr>
<td>Azithromycin (250mg/500mg) thrice weekly</td>
<td>Liver function tests at 1 month and every 6 months</td>
<td>Objective evidence of improvement:</td>
</tr>
<tr>
<td>Warn of potential side effects</td>
<td>Repeat ECG at 1 month—if QTc prolonged (&gt;450msec for men, &gt;470msec for women) stop macrolide</td>
<td>Reduction in exacerbation rate</td>
</tr>
<tr>
<td><strong>Liver function tests at 1 month and every 6 months</strong></td>
<td><strong>Enquire about side effects, especially GI upset and hearing and balance problems</strong></td>
<td>Improvement in symptoms, QoL or CAT score</td>
</tr>
<tr>
<td><strong>Review therapy at 6-12 months</strong></td>
<td><strong>Standard sputum for culture at review if able to expectorate</strong></td>
<td>Change in sputum microbiology including NTM growth</td>
</tr>
<tr>
<td><strong>Perform individual risk/benefit analysis</strong></td>
<td><strong>Medication review for potential drug interactions and QT prolongation</strong></td>
<td>Medication review for potential interactions</td>
</tr>
<tr>
<td><strong>Consider treatment break for 3-6 months each year to reduce treatment burden (and possibly reduce microbiological resistance)</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Appendix 1 - Guideline Group Members

Dr Ingrid Du Rand (Co-Chair)
Dr David Smith (Co-Chair)
Dr Charlotte Addy
Dr Tim Collyns (microbiologist)
Dr Simon Hart
Dr Phil Mitchelmore
Professor Najib Rahman
Ms Ravijyot Saggu (pharmacist)
Mrs Joan McCarthy was the lay representative (October 2016-December 2017)
Appendix 2 - Macrolide Guideline Stakeholder Organisations

Royal College of Physicians
Royal College of Physicians of Edinburgh
Royal College of Physicians and Surgeons of Glasgow
Royal College of General Practitioners
Royal College of Pathologists
Association of Respiratory Nurse Specialists
Royal College of Nursing
British Society for Haematology
British Lung Transplantation Society
UK Clinical Pharmacy Association Respiratory Group
Association for Palliative Medicine
Primary Care Respiratory Society
Association of Chartered Physiotherapists in Respiratory Care
Chartered Society of Physiotherapy
Association for Respiratory Technology and Physiology
Appendix 3 - Drugs that prolong the QT interval

**Antimicrobials**
- Moxifloxacin
- Fluconazole
- Ketoconazole

**Antiarrhythmics**
- Dronedarone
- Sotalol
- Quinidine
- Amiodarone
- Flecaïnide

**Antipsychotics**
- Risperidone
- Fluphenazine
- Haloperidol
- Pimozide
- Chlorpromazine
- Quetiapine
- Clonazepam

**Antidepressants**
- Citalopram/escitalopram
- Amitryptaline
- Clomipramine
- Dosulepin
- Doxepin
- Imipramine
- Lofepramine

**Antiemetics**
- Domperidone
- Droperidol
- Odansetron/Granesitron

**Others**
- Methadone
- Some antimalarials
- Some antiretrovirals

This list is not exhaustive. Readers are guided to reference texts such as the BNF for complete listings of individual agents.
Appendix 4

Measuring the QT interval

The QT interval is measured from the start of the Q wave to the end of the T wave. The QT interval varies inversely with heart rate, increasing as the rate slows and decreasing as the rate increases. The corrected QT interval (QTc) estimates the value of QT at a standard rate of 60 beats per minute. There are a variety of methodologies for arriving at a value for QTc from an ECG at a different heart rate.

There are free smartphone applications (such as MedCalX) and websites (such as mdcalc.com) that will generate a value for QTc from QT interval and heartrate (RR interval)

QTc is prolonged if > 450ms in men or >470ms in women

Graphic from “Life in the fast lane“ blog (litfl.com)

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https://litfl.com/qt-interval-ecg-library/

Dr Ed Burns

Accessed 26/11/2018