Addendum to the note for guidance on evaluation of medicinal products indicated for treatment of bacterial infections (CPMP/EWP/558/95 REV 2) to address indication-specific clinical data

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Addendum to the note for guidance on evaluation of medicinal products indicated for the treatment of bacterial infections (CPMP/EWP/558/95 Rev 2) to address indication-specific clinical data.
EXECUTIVE SUMMARY

During the revision of the Guidance on evaluation of medicinal products indicated for treatment of bacterial infections (CPMP/EWP/558/95 Rev 2) consideration was given to the need to provide recommendations for the design of clinical studies intended to support the approval of specific indications for use. During the consultation period and at a Workshop held before finalisation of the revised Guidance the CHMP was requested to provide detailed advice on several issues including patient selection criteria, primary endpoints, indications for which superiority or non-inferiority study designs would be expected and suggestions for non-inferiority margins. In addition, the CHMP was asked to suggest possible clinical development programmes for new antibacterial agents with very narrow spectra of antibacterial activity and/or with activity against multi-resistant pathogens for which there are very limited treatment options.

This addendum reiterates that the primary assessment of efficacy should usually occur at a test of cure visit that takes place within the same post-randomisation window in each treatment group and is timed to occur when a minimum numbers of days have elapsed from the last possible dose of protocol-defined treatment. With a few exceptions, it is not required that the primary assessment of efficacy should be confined to patients with a confirmed pathogen relevant to the type of infection under study.

Detailed guidance is provided for studies in five types of infection in which it is accepted that indications for use can be supported by a demonstration of non-inferiority of the test agent to an appropriate comparative regimen. Some suggestions for acceptable non-inferiority margins are provided. There is a lack of reliable evidence relevant to current clinical management practices to gauge the likely spontaneous resolution rates in the types of infection under consideration. The suggested non-inferiority margins have been selected on the basis that they are very likely to be sufficient to differentiate the treatment effect of the test agent from no antibacterial therapy and reflect a clinically acceptable difference to an appropriate active comparative regimen.

In indications for which a demonstration of superiority over placebo or an active comparative regimen could be required some suggestions are made for exploring appropriate patient populations and endpoints in the light of the current lack of data to support definitive recommendations for study design. In the specific case of acute otitis media recognition is given to accepting evidence of efficacy from non-inferiority studies subject to restriction of the study population and conduct of appropriate analyses.

There are several situations in which only limited evidence of clinical efficacy can be generated. Suggestions are made for possible approaches to establishing the efficacy of a test antibacterial agent in patients with severe infections for which there are limited treatment options. The development of new agents to treat multi-resistant Gram-negative aerobes/facultative anaerobes is used as an example. One possible approach could include an extensive non-clinical evaluation, robust pharmacokinetic/pharmacodynamic (PK/PD) analyses and at least one non-inferiority study in a major indication to support an indication for use against specific multi-resistant pathogen(s) even if very few such organisms had actually been treated. Additional consideration is given to clinical development programmes for new agents with very limited antibacterial spectra that may preclude their use as monotherapy for some types of infection.

Limited guidance is provided regarding the clinical assessment of treatment modalities intended to exert a local antibacterial effect as a result of direct administration to the site of infection. The specific examples covered are the topical treatment of superficial skin infections, inhalational...
therapy (excluding patients with cystic fibrosis) and oral administration of agents intended to exert an action within the gut.

Finally, consideration is given to the assessment of efficacy to support use of an antibacterial agent for treatment of some other types of infections. These include some infections for which there are special issues to consider regarding study designs and interpretation of results.

1. Introduction

It is essential that this addendum is read in conjunction with CPMP/EWP/558/95 Rev 2 in which broadly applicable general guidance is provided for the development of antibacterial agents.

CPMP/EWP/558/95 Rev 2 covers the general approach to the development of antibacterial agents. In particular, it covers matters such as microbiological investigations, study designs in treatment and prophylaxis, selection of active comparative regimens, general patient characteristics, diagnostic methods, analysis populations, primary endpoints, timing of assessment of outcomes, data analyses, studies in children and the evaluation of safety. It also addresses the development of fixed drug combinations, including antibacterial agents administered with compounds intended to inhibit a bacterial mechanism of resistance (e.g. beta-lactam agents with beta-lactamase inhibitors).

This addendum provides additional guidance on studies and clinical development programmes intended to support specific indications for use. It includes a consideration of the possible content of feasible clinical development programmes for antibacterial agents whose properties preclude their clinical evaluation along well-established lines and/or with potential for clinical activity against specific multi-resistant pathogens.

2. Scope

The addendum provides guidance on clinical data requirements to support:

- **Indications for which non-inferiority study designs are acceptable**
  This section considers five commonly sought indications that are supported by studies that demonstrate non-inferiority of the test regimen to an appropriate reference regimen.

- **Indications for which superiority study designs could be required**
  This section considers indications for which demonstration of superiority over placebo or over an active intervention is required for a pre-specified clinically relevant parameter(s). It also considers possible exceptions within these indications (e.g. in terms of patient and infection characteristics) for which non-inferiority study designs might be acceptable.

- **Circumstances in which only limited clinical data can be generated**
  This section considers the evaluation of efficacy of a test agent against uncommon or rarely encountered infections and pathogens. As an example, suggestions are made for collecting a body of evidence to support likely clinical efficacy against organisms that express specific types of resistance or patterns of multi-resistance that are currently uncommon or rare. Consideration is also given to the development of agents with a very narrow antibacterial spectrum of activity, including circumstances in which it will not be possible to evaluate these agents as monotherapy unless the pathogen can be determined before commencing treatment.
Other indications for use that could be sought

This section includes examples of indications for which some special considerations and/or problems apply to the design of clinical studies and the interpretation of data.

This addendum does not address treatment modalities that do not exert a direct antibacterial effect. For example, agents intended to modify the course of an infectious process wholly or partly via mechanisms other than inhibition of bacterial replication.

3. Main guideline text

3.1 Introduction

The sections that follow are intended to be as broadly applicable as possible. Individual clinical development programmes may need to be tailored to fit specific circumstances.

3.2 Indications for which non-inferiority study designs are acceptable

This section considers five commonly sought indications that are supported by demonstrating non-inferiority of the test regimen to an appropriate reference regimen.¹ The following observations are relevant in each example:

a) Non-inferiority margins

There is a lack of very reliable evidence relevant to current clinical management practices to gauge the likely spontaneous resolution rates (i.e. without specific antibacterial therapy) in the types of infection under consideration. In the examples that follow, the suggestions for appropriate non-inferiority margins are considered very likely to be sufficient to differentiate the effect of the test agent from no antibacterial treatment and take into account clinically acceptable differences for a test agent compared to an appropriate active comparative regimen. Sponsors should note that the suggested non-inferiority margins are applicable whether two pivotal studies are conducted or a single pivotal study is proposed.

If a single study is proposed the sponsor should give consideration to pre-defining a smaller level of significance than is usual in such studies (e.g. 0.01 rather than 0.05).

Sponsors may wish to propose alternative non-inferiority margins to those suggested (e.g. based on emerging methods for estimating the placebo effect). These proposals will be given due consideration according to the strength of the supportive evidence.

b) Route of administration

Patients with any of the five types of infection considered below usually require initial parenteral treatment, with or without a switch to oral therapy. For studies in patients with community acquired pneumonia or urinary tract infections using only oral treatment the inclusion criteria would require adjustment but the suggestions for the primary analysis are still applicable.

¹ The suggested patient characteristics in sections 3.2.1, 3.2.4 and 3.2.5 generally equate with selection of cases previously referred to as complicated infections.

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c) Pre-study antibacterial treatment

In general, up to 24 hours of prior therapy within 72 hours of enrolment may be acceptable. The protocol should specify limits for the most likely agents that would be used depending on the type of infection under study. For example, in community-acquired pneumonia (CAP) and urinary tract infection (UTI) studies the limit may be a single dose of an agent usually given once daily and 2-3 doses of agents that are routinely administered more than once a day. In intra-abdominal infections (IAI) it may be appropriate to limit prophylaxis to one pre-operative and one further dose administered during or at the conclusion of surgery. An exploratory analysis of outcomes in subgroups of patients that did and did not receive prior therapy within 72 hours for the infection under study is recommended in all studies.

Pre-study antibacterial treatment up to the time of enrolment is acceptable in a patient who has clearly failed to respond to a suitable course of antibacterial treatment (in terms of dose and duration along with documented susceptible pathogen). The protocol should specify whether prior failure includes failure to improve as well as worsening on pre-study treatment.

3.2.1 Skin and soft tissue infections

- Patient selection criteria

Acceptable types of infection for study include cellulitis, erysipelas, wound infections (traumatic or post-surgical) and major abscesses. The extent of the infection should be documented, taking into account that the acute infection may surround a chronic lesion (e.g. a varicose ulcer) that will likely remain unchanged by systemic antibacterial therapy. A minimum area affected (e.g. area of erythema, wound dimensions) or estimated size of abscess should be stated in the protocol. The proportion of patients enrolled with abscess should be limited (e.g. up to approximately 30% of total patients) and the protocol should specify a time window within which drainage should occur.

Patients should demonstrate a protocol-defined minimum number of signs and symptoms associated with an ongoing acute infectious process.

If patients with infected burns are to be enrolled the maximum extent and thickness should be specified in the inclusion criteria and the protocol should set a limit on the proportion of patients with burns that are enrolled. It is preferred that efficacy in patients with diabetic foot infections is evaluated in separate dedicated studies.

Patients with suspected or confirmed osteomyelitis or septic arthritis and those with severe necrotising infections that require specific surgical and pharmacological management should be excluded.

- Primary analysis

Clinical outcome documented at a test of cure (TOC) visit timed from randomisation so that it occurs within a window of approximately 7-14 days after the last day of treatment would be an acceptable primary endpoint. The suggested non-inferiority margin is -10%.

3.2.2 Community-acquired pneumonia

- Patient selection criteria

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All patients must have a good quality chest radiograph obtained within 48 hours prior to enrolment that shows new infiltrates in a lobar or multilobar distribution. Patients should demonstrate a protocol-defined minimum number (e.g. at least 3-4) of new onset cough, purulent sputum, fever, dyspnoea, tachyypnoea and pleuritic chest pain as well as at least one characteristic finding on percussion and/or auscultation associated with consolidation.

Sufficient data should be collected and recorded before enrolment to assign patients within the Patient Outcomes Research Team (PORT) classification system for the purposes of stratification.

- When treatment is to be initiated by the intravenous route eligible patients should have a minimum PORT score of III and at least 25% (and preferably ~50%) should have a score of IV-V. It may be appropriate to exclude patients with a score of V who require immediate ICU admission.

- In studies that involve only treatment by the oral route patients should have PORT scores of II or III at the time of randomisation and at least 50% should have a score of III.

Protocols may also capture sufficient data to determine CURB-65 scores (i.e. a scoring system based on confusion, urea, respiratory rate and blood pressure) as part of the documentation of the baseline condition of patients.

Consideration should be given to stratification of enrolment according to age < 65 years and ≥ 65 years and no upper age limit should be set.

The sponsor may include strategies to try to enrich or to minimise the study population infected with specific pathogens, such as the use of urinary antigen tests for *S. pneumoniae* or *L. pneumophila*.

Patients suspected of having pneumonia that is secondary to aspiration or a specific obstruction (e.g. malignancy and inhaled foreign body) and those with cystic fibrosis should not be enrolled.

- **Primary analysis**

Clinical outcome (based on pre-defined resolution of signs and symptoms) documented at a test of cure (TOC) visit timed from randomisation so that it occurs within a window of approximately 5-10 days after the last day of treatment would be an acceptable primary endpoint. The suggested non-inferiority margin for each study is -10%. In studies that enrol a large proportion of patients with PORT scores of IV-V, in whom the spontaneous resolution rate is expected to be lower, a wider non-inferiority margin could be acceptable.

### 3.2.3 Hospital-acquired pneumonia and ventilator-associated pneumonia

- **Patient selection criteria**

Studies may be confined to either hospital-acquired pneumonia (HAP) or ventilator-associated pneumonia (VAP). A convincing demonstration of efficacy in VAP could support an indication that includes HAP but not *vice versa*. Studies that include patients with either HAP or VAP should employ stratification to ensure that representative samples of patients in each category are enrolled (e.g. it is suggested that at least 30% should have VAP).

Patients with HAP should have been hospitalised for at least 48 hours before onset of the first signs or symptoms or these should occur within 7 days of hospital discharge. Patients should present with a minimum number of clinical features (as suggested for CAP but not including the signs on...
examination and auscultation, which may be absent) plus a new infiltrate on chest radiograph. Patients who have only been assessed in an emergency care setting should be excluded in order to enhance the likelihood that the infection is due to a pathogen highly characteristic of nosocomial infections that are commonly acquired in acute care hospitals.

In addition to clinical and radiographic features, patients with VAP should have received mechanical ventilation via an endotracheal or nasotracheal tube for at least 48 hours (i.e. the VAP population should not include patients receiving only positive pressure ventilation without intubation). Additional inclusion criteria to assist the selection of ventilated patients with an acute onset pneumonia may include documentation of the Clinical Pulmonary Infection Score (e.g. a minimum CPIS of 6), partial pressure of oxygen < 60 mm Hg in arterial blood (on room air), oxygen saturation < 90% (on room air) and worsening of the PaO2/FiO2 ratio.

Protocols may employ other scoring systems to select for a patient population that is severely ill (e.g. in whom the mortality rate is likely to exceed 10-20%). For example, the sequential organ failure assessment (SOFA) score, the multiple organ dysfunction score (MODS) and the acute physiology and chronic health evaluation score (APACHE II).

Sponsors may include pre-enrolment rapid tests that attempt to enrich or exclude patients infected with or colonised by certain species. If sponsors choose to include specifications for respiratory secretion specimens and minimum bacterial loads (in colony forming units/mL) for classifying organisms as pathogens it is imperative that the protocol also plans for analyses in which outcomes are assessed in all patients with any positive culture of a relevant pathogen from any pre-treatment respiratory tract specimen.

Primary analysis

Clinical outcome documented at a TOC visit timed from randomisation so that it occurs within a window of approximately 7-14 days after the last possible day of treatment would be an acceptable primary endpoint. The secondary endpoints should include all-cause mortality (e.g. deaths that occur up to day 28 post-randomisation) and the proportions of patients that are discharged from hospital within a pre-specified post-randomisation follow-up period.

The suggested non-inferiority margin should not exceed -12.5% in studies confined to VAP or HAP or including both HAP and VAP patients.

3.2.4 Intra-abdominal infections

Patient selection criteria

Patients should have a diagnosis of intra-abdominal infection (IAI) established during procedures such as laparotomy, laparoscopy or percutaneous drainage. Suitable diagnoses include (but are not limited to) perforations of the gall bladder, a diverticulum or the appendix, established peritonitis secondary to trauma and abscesses associated with any of these conditions. It is recommended that the proportion of patients with infections originating in the appendix should not exceed approximately 30% and that patients should be stratified at enrolment according to infection type (e.g. appendicitis-associated IAI vs. IAI secondary to other primary lesions). Patients with perforations of the stomach and small intestine should not usually be enrolled unless there is clear evidence of an established secondary infectious process within the abdominal cavity.
Primary analysis

Clinical outcome documented at a TOC visit timed from randomisation so that it occurs within a
window of approximately 7-14 days after the last possible day of treatment would be an acceptable
primary endpoint.

A non-inferiority margin of -12.5% is suggested.

3.2.5 Urinary tract infections

Patient selection criteria

Patients should have at least one of indwelling urethral (i.e. not percutaneous) catheter, urinary
retention, urinary obstruction or neurogenic bladder. Patients with ileal loops or vesico-ureteric
reflux should not be enrolled. As far as is possible, patients with signs and symptoms suggesting
prostatitis should not be enrolled.

Patients with acute pyelonephritis do not always require parenteral treatment and it is preferred
that efficacy in acute pyelonephritis is studied separately. If a study is planned to enrol patients
with any of the above conditions or acute pyelonephritis in patients considered unable to
commence oral therapy there should be stratification at enrolment according to these diagnoses
and it is recommended that the proportion with pyelonephritis should be limited.

The clinical picture should be consistent with an ongoing acute infectious process likely to have a
primary focus within the urinary tract. For example, protocols may require that patients have a
minimum number of signs of systemic upset accompanied by one or more of flank or pelvic pain,
tenderness in the costo-vertebral area, fever, dysuria, frequency or urgency.

Patients may be enrolled before microbiological culture results are available on the basis of
documented pyuria (≥ 10 WBCs/mm3) in suitable fresh urine samples, noting that specimens from
urine collection bags are not acceptable. If a mid-stream or clean catch specimen is not possible it
is preferred that patients with indwelling catheters have the catheter replaced before the sample is
obtained.

It is essential that the culture methods allow for an estimation of the bacterial load (expressed in
colony forming units [CFU]) in urine. Based on experience and consensus it would be acceptable
that patients deemed to have an infection should have > 1 x 10^5 CFU/mL. Some samples may not
meet this cut-off with a single colony type but may have at least this number of colonies in a mixed
culture based on visual inspection of morphology on an appropriate selective medium. It is
recommended that the microbiologically evaluable population should be confined to those who
have only a single colony type. Speciation is expected in clinical studies rather than reporting only
enterobacteria or other general descriptive terms.

Primary analysis

Microbiological success should be defined as < 1 x 10^3 CFU/mL. The microbiological success rate,
documented at a TOC visit timed from randomisation so that it occurs approximately 7 days after
the last possible day of treatment, would be an acceptable primary endpoint. It is expected that a
reduction of the bacterial load in urine to < 1 x 10^3 CFU/mL would usually be accompanied by
resolution of the clinical signs and symptoms suggesting infection within the urinary tract. Patients
who meet the criterion for microbiological success without clinical resolution should be fully
described and investigated.
The suggested non-inferiority margin is -10%.

### 3.3 Indications for which superiority study designs could be required

In some types of infection and/or in subsets of patients with specific conditions that may be ascribed to bacterial infection the use of active antibacterial treatment has not been established to be superior to no treatment. The reasons include, among others, high spontaneous resolution rates in certain types of infection, or at least in subsets of patients with such infections, and/or low likelihood that the clinical picture is due to a bacterial infection. These infections include (among others) acute bacterial maxillary sinusitis (ABS), acute bacterial exacerbations of chronic bronchitis (ABECB), acute otitis media (AOM) and superficial skin infections (such as impetigo and minor wounds). Another example is the use of inhaled antibacterial agents to prevent infective exacerbations in patients with chronic airways obstruction or bronchiectasis or as add-on therapy to systemic antibacterial regimens for the treatment of exacerbations or acute bacterial pneumonias.

In these instances the clinical benefit of a test agent cannot be assessed with confidence in a non-inferiority study vs. an antibacterial agent that has been approved in the past for the type(s) of infection under consideration. Therefore, efficacy should be evaluated in studies that are designed to demonstrate superiority of the test agent compared to placebo or, possibly, compared to active comparative therapy for a pre-specified clinically important endpoint. It is not possible to provide definitive recommendations for clinical development programmes in these circumstances but some suggestions are provided for consideration.

#### 3.3.1 Study designs

In several types of infection discussed in the following sections, demonstrating superiority of the test agent over placebo or over an active comparator based on clinical cure rates at a TOC visit is unlikely to be a feasible objective. To assist in selecting appropriate patient populations for study and endpoints for evaluation it is suggested that at least one exploratory study is conducted before proceeding to pivotal studies with pre-defined objectives. These exploratory studies could serve to identify potentially clinically important endpoints for which there is some likelihood that the test agent would demonstrate superiority in an adequately powered study in a carefully selected patient population. Before embarking on pivotal studies it is recommended that study designs and efficacy endpoints are discussed with EU Regulators.

For example, in studies in which patients are randomised to commence either the test agent or placebo from the outset it may be that a benefit for active treatment is demonstrated only during and/or at end of treatment i.e. active treatment speeds up resolution of the infection but it does not significantly affect cure rates assessed at a post-therapy TOC visit. An effect of active treatment on time to resolution of an infection might be regarded as clinically important if it is of sufficient magnitude. This situation is especially likely to be encountered in studies involving topical treatments for impetigo or superficial wounds. It may also apply in subsets of patients with AOM, ABS and ABECB.

One possible alternative to a study against placebo is to randomise patients either to a full course of the test agent that is commenced at study entry or to commence with placebo for a specified number of days (e.g. 48-72 hours) followed by a full course of an appropriate licensed agent. If the test agent has a safety profile that allows for a wide range of doses and if PK/PD suggests the strong possibility of a clear dose-response relationship these features could allow for a further
alternative study design that avoids a placebo group. Thus, all patients could be randomised to one
of several dose regimens of the test agent starting from the minimum that might be clinically
active at least against some potential pathogens based on PK/PD considerations.

In each of these examples the final wording of the indication would reflect the clinical benefit that
was actually demonstrated.

### 3.3.2 Acute otitis media

It is considered that published data support a specific exception to the general requirement for a
superiority study against placebo in AOM. Based on the findings reported by Tähtinen et al. (2011)
and Hoberman et al. (2011) a placebo-controlled study is not required in adequately diagnosed
AOM in children aged from 6 months up to 3 years. Nevertheless, the available data do not provide
an unequivocal indication of the primary endpoint and non-inferiority margin to apply.

An acceptable non-inferiority study in AOM must employ strict inclusion criteria. It is recommended
that all eligible children should present with acute onset (within 48 hours) otalgia and a bulging
tympanic membrane on otoscopy as a minimum. AOM may be unilateral or bilateral and
stratification is suggested. All signs and symptoms compatible with an ongoing acute infection
should be documented and the use of a scoring system is recommended. Based on the two
published studies the comparative regimen should be oral amoxicillin-clavulanate administered at
the highest dose that is approved for treatment of AOM in this age group across the study sites and
for at least 7 days.

Clinical success should require resolution of abnormalities on repeat otoscopy (in both ears if AOM
was bilateral) and resolution of otalgia. There should also be resolution of signs and symptoms of
an ongoing acute infectious process that were present at baseline. A demonstration of non-
inferiority could be based on comparison of clinical success rates at a visit timed from
randomisation to occur at 1-2 days post-therapy. It is suggested that the pre-defined non-
inferiority margin should be less than -10%. There should also be a comparison of sustained
success rates at approximately 14-21 days post-randomisation, depending on the length of
treatment and timing of the TOC visit.

At the current time an approval for treatment of AOM in other age groups and in populations that
do not meet these diagnostic criteria is not possible based solely on non-inferiority studies.

### 3.3.3 Acute bacterial sinusitis

An approval based solely on non-inferiority studies is not currently acceptable. There is a need for
further clinical data in adequately diagnosed and well-characterised patient populations before
definitive suggestions for clinical studies that could support approval for use in ABS can be made.

Meanwhile, if this indication is pursued it is recommended that the study population should consist
of patients with evidence compatible with an acute bacterial infection of the maxillary sinuses. In
addition to clinical symptoms such as facial pain and headache, diagnostic imaging should be
compatible with an ongoing infection within one or both maxillary sinuses. Establishing that the
clinical picture is due to a bacterial infection remains problematical. Maxillary drainage is currently
the only definitive method for establishing the aetiology.
3.3.4 Acute bacterial exacerbations of chronic bronchitis

An approval for the treatment of infective exacerbations of chronic bronchitis based solely on non-inferiority studies is not currently acceptable. Studies are hampered by a lack of consensus on the criteria that constitute an exacerbation and the criteria that should determine the need for specific antibacterial therapy in addition to other treatment modalities. Nevertheless, if sponsors wish to conduct studies in such patients it could be acceptable to use criteria to identify exacerbations that might benefit from antibacterial therapy suggested by at least one professional body including experts in the field.

The judgment of clinical success is also not straightforward when a return to pre-exacerbation status is likely the best that can be achieved and when each exacerbation may result in some further deterioration. All of these issues underline the need for high quality placebo-controlled studies in well-defined patient populations.

3.3.5 Inhalational antibacterial regimens in non cystic fibrosis patients

Sponsors may wish to assess the potential for an inhaled antibacterial regimen to prevent infective exacerbations of underlying conditions such as chronic bronchitis or bronchiectasis and/or to assess inhalational treatment of acute bacterial pneumonia or acute exacerbations in addition to a systemic regimen. Currently the efficacy of these possible uses of inhalational antibacterial therapy has not been established and a demonstration of superiority for the test regimen over placebo is required. In addition, since the relationship between demonstrating an effect on bacterial loads in respiratory secretions and a documented clinical benefit has not been established in any of these conditions the primary analysis must be based on an appropriate clinical endpoint.

In the case of treatment or prophylactic regimens in patients with chronic bronchitis or bronchiectasis it is essential that there are adequate pre-study investigations to fully document the presence and severity of the underlying lung condition. A major issue for the conduct and interpretation of these studies is the lack of consensus regarding the definition of an acute bacterial exacerbation. Rational criteria for the definition need to be proposed (e.g. taking into account definitions proposed by professional associations of pulmonologists) and justified in protocols.

In studies that assess the effect of single or multiple courses of an inhaled antibacterial agent on preventing bacterial exacerbations an appropriate primary endpoint could be time to exacerbation assessed over 12 months after completion of an initial or first course of the test agent (depending on the regimen under evaluation).

In the most likely scenario, studies of the treatment of acute bacterial exacerbations of underlying conditions or of acute pneumonias will involve addition of the test and placebo inhaled regimens to a standard systemic antibacterial regimen. In such cases it could be acceptable that the study demonstrates superiority for the test inhaled regimen over inhalation of a placebo based on one or more pre-specified clinical criteria (e.g. time to resolution of clinical signs and symptoms, return to baseline status).

In the case of treatment of pneumonia, subsequent to compelling results from adequate exploratory studies, sponsors may wish to demonstrate non-inferiority of an inhalational therapy alone compared to an appropriate systemic antibacterial treatment in terms of cure rates. In this instance the suggestions made in sections 3.2.2 and 3.2.3 would apply.
3.3.6 Superficial skin infections

An approval based solely on non-inferiority studies is not currently acceptable. Placebo-controlled studies in patients with impetigo, superficial wound infections and some types of secondary infected dermatoses are feasible. These should be studied separately and with appropriate limitations placed on the use of adjunctive therapies, including the use of antiseptics.

It would be acceptable if the test agent was shown to be superior to placebo based on time to resolution of the infection, which could be assessed at end of treatment. Clinical resolution should also be assessed at post-therapy visits to document relapse rates. Organisms within the two major pathogenic species (S. aureus and S. pyogenes) may manufacture a range of toxins, some of which could have a negative impact on the success of oral or topical antibacterial treatment. It is recommended that pathogens recovered from infections that have not resolved by end of treatment or which relapse should be investigated for production of toxins.

In studies in impetigo the number of lesions should be counted and an estimate made of the total body surface affected. Protocols may set limitations on numbers and/or surface area, especially if treatment is topical. The protocol may designate treatment of only the single largest lesion, a specific number of lesions or all lesions present to be treated. Depending on the strategy adopted, pre-defined additional analyses may be needed according to lesion numbers or area since untreated neighbouring lesions can affect the likelihood of clinical success at treated lesions.

The design of studies in secondary infected dermatoses should take into account the possibility of stratifying according to the underlying diagnosis, the need for ongoing topical steroid treatment and the use of occlusion.

3.4 Circumstances in which only limited clinical data can be generated

3.4.1 Introduction

This situation includes, among others, the evaluation of treatments for infections due to organisms that demonstrate specific types and/or patterns of multi-resistance that are currently uncommon or rare. No or very few patients who are infected with such organisms are likely to be enrolled in pivotal efficacy studies in commonly sought indications. Thus, alternative approaches are needed to accumulate sufficient overall evidence to support a specific endorsement for treatment of these organisms.

Additional issues arise regarding the generation of clinical efficacy data for new agents with a very narrow antibacterial spectrum of activity but a potential to be active against multi-resistant organisms.

In light of the paucity of new antibacterial agents in development and, in particular, the lack of new agents likely to be active against multi-resistant Gram-negative aerobes/facultative anaerobes, this section considers possible development programmes for such agents as an example. The approaches suggested could be applied (with modifications) to other situations in which few efficacy data can be obtained. Additional modifications of the following suggestions and tailoring of the clinical programme could be considered in certain scenarios (e.g. if an established antibacterial agent were to be co-administered with a new beta-lactamase inhibitor).
3.4.2 General considerations

The minimum level of evidence required for approval of a specific claim must be judged on a case by case basis that takes into consideration the characteristics of agent, the target population and the perceived unmet clinical need.

3.4.3 Evaluation of clinical efficacy against uncommon or rare multi-resistant pathogens

Building on the general guidance provided in CHMP/EWP/558/95 Rev 2, some possibilities for demonstrating efficacy and accumulating adequate safety data to support claims for use against multi-resistant organisms could include (but are not limited to) development programmes along the lines suggested below. Alternative approaches could be considered acceptable according to the various scenarios that can be envisaged. As one example, the total evidence for safety and efficacy that is required for approval of a fixed drug combination product in which one active substance is new and the other is already approved for use alone in certain indications (e.g. combining a licensed beta-lactam agent with a new inhibitor of beta-lactamase) would take into account relevant prior data for the known active substance.

i) In all cases it is essential to accumulate evidence to support a strong prediction of efficacy in the intended use(s) from PK/PD analyses that are founded on a thorough documentation of in-vitro activity, non-clinical evidence of efficacy and relevant human PK data. These data should address the likelihood that the test agent will be clinically active against organisms that are resistant to many or all of the licensed treatments. Since several different mechanisms of resistance could co-exist in these organisms and any one new agent may not be active in all cases it is essential that these issues are fully explored. For example, a new beta-lactamase inhibitor may prevent hydrolysis of a partner beta-lactam agent by extended spectrum beta-lactamases (ESBLs) and serine-based carbapenemases but the in-vitro activity of the combination may be considerably reduced (and it may not be clinically active) if enzyme production is accompanied by impermeability of the outer membrane or an efficient efflux pump. If these mechanisms often co-exist, then the actual efficacy of the combination may be considerably less than expected based only on enzyme inhibition data.

ii) If the antibacterial spectrum and pharmacokinetics of the test agent permit, the preferred approach would be to obtain clinical data from at least one randomised and active-controlled study in a specific type of infection. For example, if the test agent is expected to be active against multi-resistant Gram-negative aerobes/facultative anaerobes it could be studied for efficacy in HAP/VAP or IAI since many of the patients will be infected by organisms of relevant genera/species. An alternative for some new agents could be a study in UTI but this could limit extrapolation of the data due to pharmacokinetic considerations (see below). These studies are not expected to enrol sufficient numbers of patients infected with multi-resistant organisms to allow for an assessment of efficacy, although any cases that are enrolled should be carefully scrutinised for outcomes.

Patients infected with multi-resistant Gram-negative organisms may have received several prior courses of antibacterial agents and may have been hospitalised for some time. They may be debilitated and have a range of underlying chronic conditions. It is essential that the study population shares these features and includes at least a subset of patients that can be considered to be severely ill. There should be adequate PK sampling to detect any
possible effects of severe systemic upset on plasma concentrations and, as may be needed, additional PK/PD analyses.

Provided that non-inferiority is convincingly demonstrated for the test product compared to the active comparator the evidence accumulated as recommended in i) could then be used to support a claim for efficacy against specific multi-resistant organisms in this indication, assuming that the safety data collected would also support a conclusion of a favourable benefit-risk relationship. In addition, depending on non-clinical data and detailed knowledge of the PK of the test agent, consideration could be given to allowing an indication for use in patients infected with specific multi-resistant organisms when causing other types of infection under specified circumstances, as discussed in section 3.4.4.

iii) In addition to i) and ii), it is highly desirable that some pre-approval evidence is provided to support a claim for clinical efficacy against target multi-resistant pathogens, even if is based only on well-documented cases collected from a prospective non-randomised study that enrols patients regardless of the site of the infection. For example, this might be achievable if the target multi-resistant pathogens are known to be especially problematic in certain countries or specific institutions where data on clinical experience can be amassed.

iv) Additional difficulties apply to the clinical evaluation of antibacterial agents that have a very limited spectrum of activity (e.g. confined to a single genus or species). Evaluating such agents for use as monotherapy compared to an appropriate comparator is desirable since this provides a clear picture of safety. However, this is feasible only in types of infection that are commonly due to a single species and it would require availability of rapid diagnostic tests (that would need to be commercially available or developed in parallel with the antibacterial agent) to detect the presence of the target pathogen(s). If the only feasible monotherapy study were to be in patients with UTI and the pharmacokinetic data showed that very high concentrations of the test agent were achieved within the urinary tract further cautionary wording might be needed regarding a claim for treating the same multi-resistant pathogens when causing other types of infection, as discussed in section 3.4.4.

If an evaluation of monotherapy is not possible (e.g. the PK of the agent precludes a study in UTI and the spectrum does not allow for a study of monotherapy in another indication) a possible approach would be to compare addition of the test agent to one or more other agents that do not cover the same genus/species vs. standard of care in at least one type of infection. As above, patient selection should include the use of rapid diagnostic tests for the pathogen(s) of interest.

If the total data, including evidence amassed as suggested in i) and iii), were to be strongly supportive of possible clinical efficacy consideration could be given to allowing an indication for use in patients infected with specific multi-resistant organisms when causing other types of infection under specified circumstances, as discussed in section 3.4.4.

3.4.4 Reflecting the evidence in the Summary of Product Characteristics (SmPC)

There are several possible options regarding reflection of the evidence for efficacy in the SmPC and the final wording can only be decided after a full review of the data. The following proposals should be viewed as preliminary.
A test agent expected or shown to be clinically active against multi-resistant Gram-negative pathogens could be indicated for use in the types of infections that have actually been studied in the usual way and without qualification by pathogen. In this case the details of the actual organisms treated would be reflected in 5.1 along with mention of the evidence supporting activity also in the case of specific multi-resistant organisms.

In addition, consideration could be given to allowing use in types of infection that have not been studied if they are known or highly suspected to be due to specific multi-resistant pathogens. Thus, a pathogen-specific indication is a possibility. Depending on the level of evidence, the PK profile and the safety profile, such an indication might be further qualified by a restriction to use when other commonly used agents are not suitable for the individual patient.

3.5 Other indications for use that could be sought

3.5.1 Bacteraemia

Non-pathogen-specific

It may be possible to accumulate sufficient clinical data to support an indication for use of an antibacterial agent in the treatment of bacteraemia that is associated with specific types of infection, with or without restriction to certain pathogens. For example, in the case of agents that have been in use for many years and are indicated for use in a broad range of infections the total evidence may be considered sufficient for an indication that reads Treatment of patients with bacteraemia that occurs in association with, or is suspected to be associated with, any of the infections listed above (i.e. referring to the list of indications approved).

It is likely that at the time of first approval there will be very little clinical experience with an antibacterial agent in the treatment of bacteraemic patients. If no concern arises from review of the subset with accompanying bacteraemia then no statement is made about use in such patients in the SmPC except to mention the limited experience. If the antibacterial agent has been evaluated in several indications and the total number of bacteraemic patients treated across these indications is deemed sufficient (e.g. ~50 or more) to support a conclusion that efficacy is comparable to that in other patients or, at least, comparable to that of other treatments, then the addition of the sentence above could be considered appropriate.

Pathogen-specific

Studies that enroll patients with bacteraemia due to a specific pathogen but regardless of the underlying infection are not usually considered sufficient to support a pathogen-specific indication without additional qualification because this would imply that the test agent could be used to treat such cases regardless of the location of the primary focus/foci of infection (which will anyway be unknown in a proportion of cases).

An exception to this approach could apply to agents that are expected to be clinically active against uncommon or rare pathogens and/or multi-resistant pathogens for which there are few treatment options. In such cases, depending on the level of evidence that can be provided, an indication that includes bacteraemic patients regardless of the focus of infection might be considered possible with an adequate qualification of the circumstances of use.
3.5.2 Treatment of acute bacterial infections in neutropenic patients

The institution of an antibacterial agent prior to or at the time of onset of expected neutropenia is now a common practise in some patient populations and centres so that rates of breakthrough infections may be comparatively low compared to other patient groups. The study population actually enrolled with acute bacterial infections during neutropenia will comprise some ratio of patients with breakthrough infections despite prophylaxis and patients who have not received routine prophylaxis. The two sub-groups may be substantially different in terms of their underlying conditions and are likely to be enrolled at different centres with variable routine management protocols. On this basis stratification according to prior or no prophylaxis may be appropriate. The protocol should provide clear criteria to be met in terms of neutropenia (cut-off and expected duration). The definition of fever will also require alignment across sites.

If the test agent must be co-administered due to its spectrum of activity then the additional agent(s) should be specified, including dose regimen and any dose adjustments. If possible the range of agents allowed should be standardised. The protocol should include clear criteria for stopping therapy in terms of susceptibility data, clinical progress, culture results and recovery of the granulocyte count. It is critical that the criteria for failure are very carefully specified (e.g. persistence of the baseline pathogen beyond ~48 hours of treatment).

The most objective basis for the assessment of efficacy would be the comparison of bacterial eradication rates in the subset of patients with a positive blood culture pre-treatment between the test and comparative regimens. Patients with an obvious primary focus should also have a resolution of infection.

Due to the complex nature of these patients, difficulties in ascertaining the range of co-existing pathogens and lack of clear distinction between the treatment and prophylactic role of antibacterial agents (even in the subset with a documented bacterial pathogen) the resulting indication would likely reflect the utility of the agent in the overall management of such patients rather than specifying use in the treatment of bacterial infections.

3.5.3 Eradication of carriage

Sponsors may wish to pursue studies that have the primary aim of demonstrating an effect of test agents on carriage of specific bacterial species.

Indications that relate to the reduction or eradication of a pathogen from a specified body site are not acceptable unless the microbiological findings have been shown to result in a measurable clinical benefit. In most examples that could be envisaged the provision of published data alone to support a link between an effect on carriage and a clinical benefit would not be acceptable. In these cases the clinical benefit associated with the effect on carriage should be assessed in a placebo-controlled study. Demonstration of non-inferiority versus an active regimen would only be acceptable if current clinical opinion rules out the possibility of using a placebo.

Possible exceptions could include the use of oral treatment regimens to eradicate carriage of meningococci from the nasopharyngeal area of contacts of cases and the eradication of *S. pyogenes* in order to reduce the risk of post-streptococcal syndromes (e.g. rheumatic fever and glomerulonephritis). In these examples a study of the test agent against placebo/vehicle is not feasible. Pivotal studies would have to demonstrate non-inferiority for the test agent regimens against recommended regimens based on microbiological eradication rates (see below).
In addition, sponsors may be able to justify that eradication of *S. aureus* carriage at some body sites prior to specific types of surgical procedures can be expected to reduce the rate of post-operative infections. It is most likely that such studies will involve direct application of the test agent to the anterior nares. It is expected that pivotal studies to support this use will aim to demonstrate superiority of the test agent compared to placebo/vehicle in terms of microbiological eradication rates (see below) at least until such time as clinical practise would make this study design no longer feasible.

Microbiological culture techniques cannot demonstrate absolute eradication since there will always be a minimum number of organisms that cannot be detected. Therefore only a reduction in *numbers* (within a range that can be differentiated by culture) or apparent eradication (i.e. negative cultures) can be demonstrated. In cases that involve topical applications there is also the issue of a carry over effect from residual active agent at the sampling site influencing the numbers of organisms cultured, which may give a falsely optimistic view of the real effect. For all these reasons it is essential that there is an extensive documentation of the detection limits of the sampling and culture methodologies applied in pivotal studies. Other detection methods, such as PCR, cannot differentiate live from dead organisms and data obtained from these methods should not be used for the primary assessment of efficacy.

Pivotal studies should be conducted in the patient population and setting(s) in which the product is proposed for routine use. In this way some assessment of the treatment duration required to achieve the required effect and of the risk of and time to re-colonisation would be facilitated. This requires that there are adequate means available for differentiating re-growth of initial strains from new colonisation events. Organisms recovered from patients who fail to achieve apparent eradication or who show a very slow response to treatment, rapid re-growth or re-colonisation should be fully characterised in terms of susceptibility, mechanisms of resistance and, as may be appropriate to the species, other features such as sub-type and toxin encoding genes/toxin production.

### 3.5.4 Oral treatment intended to exert an action within the gut

Currently, antibacterial regimens intended to exert an action within the gut (some of which are and some not absorbed systemically to any potentially clinically useful extent) have been approved for the treatment of *C. difficile* infections producing diarrhoea and for the treatment of travellers’ diarrhoea (with variably specified usages according to genera).

The systemic absorption of agents intended for these uses should be adequately characterised and an appropriate range of pharmacokinetic studies should be conducted accordingly. The implications of any systemic absorption for selection of drug-resistant organisms colonising body sites other than the gut should be discussed.

In these types of indications PK/PD analyses do not assist in predicting an effective dose and adequate dose-finding studies are needed.

For treatment of *C. difficile* associated diarrhoea a demonstration of non-inferiority of the test agent compared to a licensed agent would be acceptable. The patient population should have carefully documented changes in bowel habit within a pre-defined pre-study period accompanied by detection of toxin (A or B) in stools. An established *C. difficile* infection (CDI) severity index should be applied within the inclusion criteria. The primary efficacy endpoint should be the cure rate using a definition of cure that encompasses resolution of symptoms and no requirement for...
further antibacterial treatment. The suggested non-inferiority margin is 10%. There should be sufficient follow-up to document relapse rates.

In the case of travellers’ diarrhoea the rate and rapidity of spontaneous resolution varies according to the pathogen. In a population presenting with recent onset travellers’ diarrhoea that is not associated with any features suggestive of the presence of an invasive pathogen it is expected that the test agent is shown to be superior to placebo. A third treatment arm in which subjects receive an antibacterial agent approved for use in this setting could be included for assay sensitivity purposes. Protocols should make adequate provision for subject management when a pathogen that requires specific treatment is detected after enrolment and/or there is rapid worsening (e.g. onset of blood in stool) during the study period.

Eligible subjects should have an acute onset of diarrhoea within a defined number of days before enrolment that is characterised by a minimum number of unformed stools per day. The recommended primary endpoint is time to last unformed stool (TLUS).

Suitable test agents should at least demonstrate in-vitro activity against *E. coli*. The risk of encountering organisms of this and other species that are unlikely to be susceptible to the test agent at concentrations expected within the gut should be taken into account in the study design and may influence the geographical location of study sites. It is particularly important that the identity and in-vitro susceptibility of pathogens recovered from subjects who do not respond to the test agent are fully documented since the clinical effect of test agents within the gut may differ from expectations based solely on in-vitro and PK data.