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Standardising neonatal and paediatric antibiotic clinical trial design and conduct: the PENTA-ID network view

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ABSTRACT

Antimicrobial development for children remains challenging due to multiple barriers to conducting randomised clinical trials (CTs). There is currently considerable heterogeneity in the design and conduct of paediatric antibiotic studies, hampering comparison and meta-analytic approaches. The board of the European networks for paediatric research at the European Medicines Agency (EMA), in collaboration with the Paediatric European Network for Treatments of AIDS—Infectious Diseases network (www.penta-id.org), recently developed a Working Group on paediatric antibiotic CT design, involving academic, regulatory and industry representatives. The evidence base for any specific criteria for the design and conduct of efficacy and safety antibiotic trials for children is very limited and will evolve over time as further studies are conducted. The suggestions being put forward here are based on the adult EMA guidance, adapted for neonates and children. In particular, this document provides suggested guidance on the general principles of harmonisation between regulatory and strategic trials, including (1) standardised key inclusion/exclusion criteria and widely applicable outcome measures for specific clinical infectious syndromes (CIS) to be used in CTs on efficacy of antibiotic in children; (2) key components of safety that should be reported in paediatric antibiotic CTs; (3) standardised sample sizes for safety studies. Summarising views from a range of key stakeholders, specific criteria for the design and conduct of efficacy and safety antibiotic trials in specific CIS for children have been suggested. The recommended criteria are intended to be applicable to both regulatory and clinical investigator-led strategic trials and could be the basis for harmonisation in the design and conduct of CTs on antibiotics in children. The next step is further discussion internationally with investigators, paediatric CTs networks and regulators.

WHAT IS THE PROBLEM?

Antimicrobial resistance (AMR) is a rapidly emerging problem, causing morbidity and mortality especially in vulnerable populations. Mortality attributable to AMR may be associated with discordant therapy, which is particularly challenging in neonates and children due to the limited number of approved effective antimicrobials, the inadequate pipeline for novel antibiotics and the long delays noted in many documents between the adult and paediatric licensing of novel antibiotics.1,2 There is no evidence that the significant delays in paediatric licensing of new antibiotics is improving. Antimicrobial development for children remains challenging due to multiple barriers to conducting clinical trials (CTs), with nearly half of paediatric medicines in Europe prescribed off-label, without evidence on the optimal dosage or safety data.3 The Clinical Trials Transformation Initiative, aiming to develop and drive adoption of practices that will increase the quality and efficiency of CTs, recently organised a Multi-Stakeholder Expert Meeting with the aim to identify and address barriers in conducting antibacterial CTs in neonates and children.4 We have previously reported on the marked heterogeneity in the design and conduct of paediatric antibiotic trials, with a lack of standardisation of the key inclusion/exclusion criteria and endpoints for specific clinical infectious syndromes (CIS) hampering comparison between studies and meta-analytic approaches.4

Among the initiatives put in place to improve the efficiency and feasibility of paediatric CTs was the publication of a Paediatric Addendum to the Guideline on the evaluation of medicinal products indicated for treatment of bacterial infections5 by the Committee for Medicinal Products for Human Use and the Infectious Diseases Working Party at the European Medicines Agency (EMA). The purpose of this publication was to provide some general consideration on the paediatric clinical development programmes required to support the authorisation of antibacterials for treatment of infectious diseases in children AND in the optimal design and conduct of clinical investigator-led strategic trials.5 The board of the
European networks for paediatric research at the EMA, in collaboration with the Paediatric European Network for Treatments of AIDS—Infectious Diseases network, therefore developed a Working Group (WG) on paediatric antibiotic CT design, involving academic, regulatory and industry representatives from both the USA and Europe. This group aimed to provide practical guidance on the design and conduct of neonatal and paediatric antibiotic CTs in order to improve international harmonisation in this important area. Currently, the EMA recommends the conduct of single-dose/multi-dose pharmacokinetic (PK) studies to support the approval of an antibacterial agent to treat infectious diseases in paediatric patients.3

The evidence base for any specific criteria for the design and conduct of efficacy and safety antibiotic trials for children is very limited and will evolve over time as further studies are conducted. The suggestions being put forward here are based on the adult EMA guidance, adapted for neonates and children.4-6 In particular, this document provides suggested guidance on the general principles of harmonisation between regulatory and strategic trials, including:

1. Standardised key inclusion/exclusion criteria and widely applicable outcome measures for specific CIS to be used in CTs on efficacy of antibiotic in children.
2. Key components of safety that should be reported in paediatric antibiotic CTs.
3. Standardised sample sizes for safety studies.

**GENERAL PRINCIPLES**

There are clear differences between regulatory trials being conducted to obtain a marketing authorisation for a new molecular entity and strategic trials usually sponsored by academic institutions. However, where possible, similar standards should apply across both types of studies. This distinction is becoming less wide as collaboration between clinical academic CT networks and pharma to drive efficiency increases, with both groups committed to the more rapid delivery of high quality trials.7 Recent data has noted that there has been in general inadequate reporting of safety in investigator-led paediatric antibiotic CTs and marked heterogeneity of the key trial elements, for example, inclusion/exclusion criteria and definition and timing of end points.5 4

There is increasing recognition that for the great majority of paediatric regulatory antibiotic trials, for well-established classes, both efficacy and safety can be bridged from adult studies. Single-dose PK studies are difficult to perform and there needs to be a clear focus on reducing barriers to recruitment. In our view, single-dose PK studies do not need to be conducted only in the CIS where there is an adult licence but will recruit more efficiently as an ‘all comers’ study where the child may be in hospital with any CIS. We can see no scientific rationale why the PK for the great majority of antibiotics (eg, a beta lactam/beta lactam inhibitor combination—BL/BLI) will be different if the child is stable and completing a course of antibiotics for complicated urinary tract infection, or a complicated intra-abdominal infection. The child should be clinically stable, on either intravenous or oral antibiotics, being given for treatment or prophylaxis. If the antibiotic has very predictable linear PK and a well described safety profile, then the cohorts across all ages, including neonates, should be opened at the same time. Wherever possible, the neonatal single-dose PK cohort should be included in the same protocol as the older age cohorts, as separate protocols may lead to significant delays in determining the neonatal dose.

The scientific rationale for a multi-dose study needs to be determined on a case by case basis. As can be seen from the sample sizes given below, multi-dose studies of less than around 100 children for well-established classes of antibiotics will not be adequately powered to determine any new safety signal that is not entirely predictable from that drug class. Some antibiotics that, for example, require a loading dose calculated from the single-dose PK study, will need a multi-dose validation PK study. There may be a rationale for certain novel antibiotics to gain experience of routine clinical use from an open label, all comers, multi-dose treatment study, while recognising that the study is not required for PK and is not powered for either safety or efficacy. There does not appear to be any clear scientific rationale for the great majority of well-established classes of antibiotics for randomisation between the novel agent and a standard of care (SOC) arm. Even if SOC can be controlled to a limited number of regimens (which is often difficult in studies requiring multiple sites to achieve the recruitment targets), as seen from the sample sizes given below, the trial would need to be recruiting a very substantial number of children to detect any novel safety signal that was not entirely predictable from the drug class. It should be emphasised that these comments only apply to well-established drug classes (eg, BL/BLIs, aminoglycosides, etc).

The optimal study design for neonates is evolving and requires further international consensus. There is an urgent global unmet clinical need for novel antibacterial agents to treat neonates, both term and preterm, with multidrug resistant bacterial infection causing neonatal sepsis. Equally, not every new antibiotic under development needs evaluation in neonates, where, for example, there are already other treatment options. For many new antibiotics single-dose PK studies would be all that would reasonably be required.

The next step would be a prioritisation of the novel antibiotics that are a high priority for multi-dose safety and efficacy trials in neonates. This could be based on the WHO Priority Pathogen List, focussing on the most critical pathogens, specifically those agents active against carbapenem resistant organisms.9 For these relatively few antibiotics, PK, safety and efficacy data are required in the indication of neonatal serious/severe bacterial infection (SBI, sepsis). Evaluation of penetration of the drug into the central nervous system for these antibiotics is also required. With the majority of neonatal sepsis caused by
multidrug resistant Gram-negative pathogens, much of which is healthcare-associated, there is no obvious scientific basis to divide neonatal sepsis into early and late onset sepsis and the general term neonatal serious bacterial infection is the most suitable term (as used by the WHO). These studies will need to recruit babies across all stages of prematurity and postnatal age. Given the challenges of recruitment into such a population and the need for such trials to recruit globally, active consideration should be given for establishing close collaboration between Pharma, the WHO, global paediatric infectious diseases CTs networks and other major stakeholders, similar to the structures that were developed for paediatric HIV infection. Novel trial designs need to be urgently considered allowing the inclusion of multiple agents within protocols, focussing on obtaining both regulatory and public health outcomes within single trials. We urgently recommend the WHO to convene a consensus meeting focused specifically on neonatal sepsis to drive forward the global collaboration required.

The reporting of pharmacovigilance data on antibiotics in neonates and children is currently limited. At the moment, a standardised method of conducting antibiotic pharmacovigilance in children and neonates has not been developed, particularly for drugs that are used off-label. This is an increasingly important area for all medicines as key regulatory trials have smaller sample sizes related to cost considerations. The establishment of a network of different stakeholders (academics, physicians, regulators and industry) involving centres in all regions across the world would allow the conduct of prospective cohort studies using electronic data records as part of post-marketing surveillance (as has already been set up with paediatric HIV registries). Such approach could potentially allow data to be collected and easily pooled out at a relatively low cost.

Suggested key clinical and laboratory components of inclusion/exclusion criteria and endpoints for CIS in paediatric antibiotic CTs

In the absence of any clearly accepted criteria, while recognising the very limited evidence base but given the wide variation seen in reported CTs, the WG has developed suggestions for paediatric inclusion/exclusion criteria and endpoints for the most common CIS.

Based on the results of a recently conducted systematic review, the most frequently reported CIS-specific clinical and laboratory criteria for the enrolment and evaluation of children in antibiotic CTs were collected. These criteria were then compared with the EMA Guideline on the evaluation of medicinal products indicated for treatment of bacterial infections, revised according to the expert opinion of the WG members, and summarised in table 1. The WG decided to adapt the adult EMA criteria for children and neonates in all those CIS in which a similar pathophysiology and a similar spectrum of pathogens across the target age groups could be anticipated. This has been also the principle that has been adopted in the Paediatric

Addendum for the extrapolation of efficacy against an infectious disease from adults to paediatric patients. This situation applies to the majority of infectious diseases that occur both in adults and in one or more paediatric age subgroups. However, there are some cases in which the pathophysiology and the spectrum of pathogens differ substantially between children/neonates and adults. As discussed above, for example, this is the case of neonatal sepsis (neonatal SBI). In this case, age-specific criteria have been adopted specifically designed for the neonatal age.

Key components of safety in paediatric antibiotic CTs

Proper reporting of safety data when publishing clinical studies would increase translation of results into clinical practice. We have previously published a systematic review of safety data reported in CTs of anti-bacterial agents in children and neonates to determine if age-specific adverse events (AEs) could be identified for different antibiotic classes. The quality of reporting AEs was suboptimal in the great majority of CTs, due to the lack of detailed definitions of expected/unexpected AEs (with respect to the AEs that have been reported in adults and/or the mechanism of action of the study drug), grading, reference for Coding System, and age stratification of the results. To improve the quality of safety reporting we recommend that there should be a specific section on safety in every paediatric antibiotic CT. To allow an appropriate comparison between CTs, studies should provide:

- Justification of the sample size for safety, and definition of the safety population in studies having safety as a primary endpoint.
- Definition for:
  - How harms-related information was collected (mode of data collection, timing, attribution methods, harms-related monitoring and stopping rules).
  - Predefinition of each specific clinical/laboratory/imaging addressed AE.
  - Grading (mild, moderate, severe).
  - Relationship with the study drug (expected vs unexpected).
  - Reference for Coding System (taking into account that most groups are now using the Division of AIDS grading system).
- Overall (all age groups together) analysis presented first, followed by stratification of safety assessments and results by different age groups.
- Data on any modification to randomised treatment OR withdrawals because of AEs.
- All the denominators and all absolute risks per arms and per AE type, grade, seriousness and severity.

Standardising sample sizes for paediatric antibiotic CTs

Data obtained from underpowered studies limits the implementation of the result itself, wastes resources, and undermines the ethics of patients’ involvement. However,
### Table 1 The key components of inclusion/exclusion criteria and endpoints for infectious CIS in paediatric AB CTs

#### Community-acquired pneumonia (CAP)

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
<th>Efficacy endpoints</th>
</tr>
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<tbody>
<tr>
<td>▶ Chest X-ray with new infiltrates in a lobar or multilobar distribution. AND ▶ A minimum number (at least 3–4) of new onset: Cough, Fever, Dyspnoea, Tachypnoea. AND ▶ Pleuritic chest pain.</td>
<td>▶ Chronic/underlying conditions (eg, moderate or severe asthma, cystic fibrosis, immunodeficiency, malignancy). ▶ Pneumonia secondary to aspiration or a specific obstruction (eg, malignancy and inhaled foreign body).</td>
<td>▶ Timing for evaluation: ▶ End of treatment (EOT). ▶ Test of cure (TOC) 5–10 days after the EOT. ▶ Follow-up 28 days after randomisation. ▶ Clinical cure: ▶ Resolution or significant improvement of the baseline signs and symptoms documented at TOC visit AND discontinuation of antibiotics. OR ▶ Treatment failure: ▶ Any change, modification, or discontinuation of allocated AB therapy because of deterioration in patient’s condition, development of serious intercurrent illness or complications. ▶ Persistence of signs and symptoms present at the enrolment. ▶ Relapse of the hypoxemic pneumonia during the following 2 weeks. ▶ Death (up to 28 days post-randomisation).</td>
</tr>
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</table>

#### Ventilator-associated pneumonia (VAP) and hospital-acquired Pneumonia (HAP)

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
<th>Efficacy endpoints</th>
</tr>
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<tbody>
<tr>
<td>HAP: ▶ Hospitalised for at least 48 hours before onset or onset within 7 days of hospital discharge. AND ▶ Chest X-ray with new infiltrates in a lobar or multilobar distribution. AND ▶ Minimum number of clinical features (as suggested for CAP but not including the signs on examination and auscultation).</td>
<td>▶ VAP population: patients receiving only non-invasive positive pressure ventilation.</td>
<td>▶ Timing for evaluation: ▶ End of treatment (EOT). ▶ Test of cure (TOC) 7-14 days after the EOT. ▶ Follow-up 28 days after randomisation. ▶ Clinical cure: ▶ Resolution or significant improvement of the baseline signs and symptoms documented at EOT visit AND discontinuation of antibiotics. OR ▶ Treatment failure: ▶ Any change, modification, or discontinuation of allocated AB therapy because of deterioration in patient’s condition, development of serious intercurrent illness or complications. ▶ Persistence of signs and symptoms present at the enrolment. ▶ Relapse of the pneumonia during the following 2 weeks. ▶ Death (up to 28 days post-randomisation).</td>
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Combined HAP/VAP studies: enrol representative samples in each category (at least 25%-30% VAP).
Complicated urinary tract infections (UTI)\(^9\) (including pyelonephritis, renal abscess, catheter-related UTI, bacteraemia from urinary tract without specification)

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
<th>Efficacy endpoints</th>
</tr>
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<tbody>
<tr>
<td>► Abnormal urinary dipstick test (leucocyte esterase &gt;1+, or nitrite positive) OR ► Urinalysis ≥5 white cell counts per high-power field in centrifuged urine or ≥10 white cell counts per mm(^3) in uncentrifuged urine and bacteriuria with any bacteria per high-power field. AND ► At least two of the following clinical or biological signs: – Fever with temperature of 38°C or higher. – General, non-specific signs (for children &lt;2 years irritability, vomiting, diarrhea, feeding problems; for children &gt;2 years abdominal or flank pain, urgency, frequency, dysuria, or suprapubic tenderness). – O-reactive protein or procalcitonin concentrations elevated according to the local laboratory. AND ► Positive urine culture result with no more than two species of micro-organisms OR ► Spontaneously voided urine with ≥10(^5) micro-organisms per mL of urine OR ► Suprapubic aspirate or urinary catheter with ≥10(^4) micro-organisms per mL of urine OR ► Positive blood culture result and no other recognised cause.</td>
<td>► Chronic/underlying conditions (eg, known UT abnormalities, malignancy, immunodeficiency, shock).</td>
<td>Timing for evaluation: ► End of treatment (EOT). ► Test of cure (TOC) 5–7 days after the EOT. ► Follow-up 28 days after randomisation. Treatment success: Concomitant clinical and microbiological evaluation for the TOC ► Clinical cure defined as defervescence. ► Microbiological cure defined as urine sterilisation.</td>
</tr>
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Suspected or proven neonatal severe bacterial infection

<table>
<thead>
<tr>
<th>Inclusion criteria(^a)</th>
<th>Exclusion criteria</th>
<th>Efficacy endpoints</th>
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<tr>
<td>At least two clinical symptoms and at least two laboratory signs in presence of or as a result of suspected or proven infection. Clinical criteria: ► Hyperthermia or hypothermia and/or temperature instability. ► Reduced urinary output or hypotension or mottled skin or impaired peripheral perfusion. ► Apnoea or increased oxygen requirement or requirement for ventilation support. ► Bradycardia spells or tachycardia and/or rhythm instability. ► Feeding intolerance or abdominal distension. ► Lethargy or hypotonia or irritability. ► Skin and subcutaneous lesions, such as petechial rash. Laboratory criteria: ► WBC count &lt;4000 or &gt;20,000 cells/mm(^3). ► Platelet count &lt;100,000/mm(^3). ► CRP &gt;1.5 mg/dL or procalcitonin ≥2 ng/mL. ► Glucose intolerance as expressed by blood glucose values &gt;180 mg/dL confirmed at least twice or hypoglycaemia &lt;40 mg/dL (2.5 mmol/L). ► Acidosis as characterised by base excess &lt;−10 mmol/L or lactate with value above 2 mmol/L.</td>
<td>► Major underlying conditions or major congenital malformations. ► Deep seated localised infection (including osteomyelitis, endocarditis and meningitis).</td>
<td>Timing for evaluation: ► End of Treatment (EOT). ► Test of Cure (TOC) 7–10 days after the EOT. ► Long-term follow-up 28 days after randomisation. Clinical cure: ► Resolution or significant improvement of the baseline clinical signs and symptoms of infection AND microbiological eradication or presumed eradication (in case of positive blood culture) AND no new symptoms suggestive to neonatal sepsis. OR Treatment failure, any of the following: ► Death up to 28 days post-randomisation. ► Persistence of signs and symptoms present at the enrolment. ► Clinical deterioration: emergence of any sign of critical illness at any time or re-emergence of a sign of clinical severe infection after initial disappearance. ► Any change, modification, or discontinuation of allocated AB therapy because of deterioration in patient’s condition, development of serious intercurrent illness or complications.</td>
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Table 1  Continued

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
<th>Efficacy endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complicated intra-abdominal infections (cIAI) (including localised infections (e.g., biliary and non-biliary organ-specific infections, abscesses, postsurgical secondary infections) and diffuse peritonitis. 10% of patients should have a diagnosis other than complicated appendicitis)</td>
<td>Diagnosis of cIAI established during procedures such as laparotomy, laparoscopy or percutaneous drainage</td>
<td>Chronic/underlying conditions affecting surgical decision making or that would limit recovery (e.g., haemophilia, severe cardiac or respiratory comorbidities).</td>
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<tr>
<td>► Patient has organisms cultured from purulent material from intra-abdominal space obtained during a surgical operation for the current infection or needle aspiration OR ► Organisms cultured from drainage from surgically placed drain (e.g., closed suction drainage system, open drain, T-tube drain) OR ► Organisms cultured from blood and radiographic evidence of infection (e.g., abnormal findings on ultrasound, CT scan, MRI, or radiolabel scans (gallium, technetium, etc) or on abdominal X-ray). AND ► At least two of the following signs or symptoms with no other recognised cause: – Fever (&gt;38.5°C). – Nausea. – Vomiting. – Abdominal pain. – Jaundice.</td>
<td>Timing for evaluation: ► End of treatment (EOT). ► Test of cure (TOC) 7–10 days after the EOT. ► Long-term follow-up 28 days after randomisation. Clinical cure: ► Clearance of infection with resolution of the baseline signs and symptoms such that no additional antibacterial therapy or surgical or percutaneous intervention is required at EOT AND eradication or presumed eradication of micro-organisms. OR Treatment failure: ► Any change, modification, or discontinuation of allocated AB therapy because of deterioration in patient’s condition, development of serious intercurrent illness or complications (e.g., abscess, wound infection, prolonged fever &gt;3 days, prolonged bacteraemia), need for additional antibiotics, need for operative or percutaneous intervention.</td>
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</table>

| Acute bacterial skin and skin structure infections (ABSSSI) (including surgical wound infections, deep abscesses, cellulitis and erysipelas) | Diagnosis of ABSSSI requiring systemic antibiotic treatment, with at least one of the following: – Drainage/discharge. – Erythema. – Fluctuance. – Heat/localised warmth. – Pain/tenderness to palpation. – Swelling/induration. AND at least two of the following: – Fever or hypothermia. ► Leucocytosis or leucopenia or a left shift of band neutrophils. – Tachycardia (98th percentile for age).21 – Tachypnoea (2 SD of normal for age).21 | Chronic/underlying conditions that would limit recovery (e.g., neutropenia, diabetes, other immunoendocrinopathies). ► Patients with suspected and/or confirmed osteomyelitis or septic arthritis. ► Patients with mild infections that do not need systemic antibiotics. |
| Timing for evaluation: ► End of Treatment (EOT). ► Test of cure (TOC) 7–10 days after the EOT. ► Long-term follow-up (LFU) 28 days after randomisation. Clinical cure: ► Resolution or significant improvement of the baseline clinical signs and symptoms of infection at EOT AND ► No need for additional antibiotics. OR Treatment failure: ► Persistence or progression of signs and symptoms or development of new lesions at a different site. ► Any change, modification, or discontinuation of allocated AB therapy because of deterioration in patient’s condition, development of serious intercurrent illness or complications. |

AB, antibiotics; CIS, clinical infectious syndromes; CT, clinical trial.
Table 2 Sample size for single-arm interventional paediatric antibiotic CTs having safety as a primary endpoint, according to the rates of adverse events (AEs) per single drug class reported in the systematic review

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Overall percentage experiencing AEs*</th>
<th>Sample size to provide &gt;0.80 probability that final 95% CI around estimated AE rate is no more than 10% above this</th>
<th>Upper 97.5% confidence limit around an observation of 0/N (%)</th>
<th>Sample size to provide &gt;0.90 probability that final 95% CI around estimated AE rate is no more than 10% above this</th>
<th>Upper 97.5% confidence limit around an observation of 0/N (%)</th>
<th>Sample size to provide &gt;0.95 probability that final 95% CI around estimated AE rate is no more than 10% above this</th>
<th>Upper 97.5% confidence limit around an observation of 0/N (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penicillins</td>
<td>13</td>
<td>106</td>
<td>3.4</td>
<td>139</td>
<td>2.6</td>
<td>172</td>
<td>2.1</td>
</tr>
<tr>
<td>Aminoglycosides</td>
<td>3</td>
<td>51</td>
<td>7.0</td>
<td>70</td>
<td>5.1</td>
<td>79</td>
<td>4.6</td>
</tr>
<tr>
<td>Cephalosporins</td>
<td>16</td>
<td>114</td>
<td>3.2</td>
<td>152</td>
<td>2.4</td>
<td>190</td>
<td>1.9</td>
</tr>
<tr>
<td>Macrolides</td>
<td>22</td>
<td>135</td>
<td>2.7</td>
<td>180</td>
<td>2.0</td>
<td>229</td>
<td>1.6</td>
</tr>
<tr>
<td>Penicillins+BLI</td>
<td>46</td>
<td>165</td>
<td>2.2</td>
<td>226</td>
<td>1.6</td>
<td>283</td>
<td>1.3</td>
</tr>
<tr>
<td>Fluoroquinolones</td>
<td>36</td>
<td>161</td>
<td>2.3</td>
<td>225</td>
<td>1.6</td>
<td>277</td>
<td>1.3</td>
</tr>
<tr>
<td>Carbapenems</td>
<td>33</td>
<td>158</td>
<td>2.3</td>
<td>214</td>
<td>1.7</td>
<td>270</td>
<td>1.4</td>
</tr>
<tr>
<td>Linezolid</td>
<td>61</td>
<td>153</td>
<td>2.4</td>
<td>205</td>
<td>1.8</td>
<td>258</td>
<td>1.4</td>
</tr>
<tr>
<td>Glycopeptides</td>
<td>75</td>
<td>117</td>
<td>3.1</td>
<td>153</td>
<td>2.4</td>
<td>185</td>
<td>2.0</td>
</tr>
<tr>
<td>Sulfonamides+trimethoprim</td>
<td>5</td>
<td>59</td>
<td>6.1</td>
<td>85</td>
<td>4.2</td>
<td>102</td>
<td>3.6</td>
</tr>
<tr>
<td>Amphenicols</td>
<td>4</td>
<td>55</td>
<td>6.5</td>
<td>73</td>
<td>4.9</td>
<td>91</td>
<td>4.0</td>
</tr>
</tbody>
</table>

The third, fifth and seventh columns represent the sample size that would provide a >0.80, >0.90 and >0.95 probability, respectively, that the final 95% CI around the estimated percentage experiencing AEs in the new trial was no more than 10% higher than the average rate provided in the second column. The fourth, sixth and eighth columns provide the upper 97.5% confidence limit around an observation of zero AEs of a particular type from this number of children.

*Data are expressed as median proportion of overall AEs among the studies included in the systematic review by Pansa et al.3 rounded to the nearest percentage point.

BLI, beta lactamase inhibitor; CT, clinical trial.
in the safety review, only two trials provided the justification for the sample size specifically for the safety population, including those designed with safety as the primary endpoint.

In an attempt to provide a standardised sample size to be used in single-arm interventional paediatric antibiotic CTs having safety as a primary endpoint, based on the rates of AEs per single drug class reported in the systematic review, the WG considered some key underpinning concepts. First, the rates of AEs and serious AEs (SAEs) in children are generally low, often lower than in adults, and usually predictable by class; AEs/SAEs specific to children occur extremely rarely, but are important to detect; and blinded (placebo-controlled) or unblinded comparative trials aim to estimate the difference between AE rates with the new antibiotic versus a comparator, with sample sizes typically large if designed to exclude differences outside a non-inferiority margin, or powered only to detect very large reductions in AEs which may not be realistic.

Given this, a reasonable approach would be to ensure sufficient children receive a novel antibiotic to enable (1) a high probability of determining that the overall AE/SAE rate is estimated reasonably precisely and (2) a reasonable probability of observing an AE which occurs in 1/20 children, or equivalently, that an observation of zero events has an upper 97.5% CI which lies below 5%. This could be done within a single-arm interventional trial with a standard proportion test (as, eg, in Flahault et al). Given an expected proportion of children experiencing one or more AEs, and a maximum acceptable value for this proportion, the sample sizes in table 2 provide the 0.95, 0.90 and 0.80 probability that the upper 95% CI around the proportion of children experiencing one or more AEs in the new trial is below the maximum acceptable value. An observation of no AEs of a particular type from this number of children has an upper 97.5% CI limit which is approximately 3/N (as a proportion). For example, for 0 events observed from 60 children, the approximation is 3/n=3/60=1/20=0.05 (compared with the actual exact upper limit, which is 0.06).

A further analysis then considered the potential class-specific sample sizes using data from the safety systematic review discussed above. In table 2, the third, fifth and seventh columns represent the sample size that would provide a >0.80, >0.90 and >0.95 probability, respectively, that the final 95% CI around the estimated percentage experiencing AEs in the new trial was no more than 10% higher than the average rate provided in the second column. The fourth, sixth and eighth columns provide the upper 97.5% confidence limit around an observation of zero AEs of a particular type from this number of children (ie, the degree of certainty that an AE that was not observed in the trial genuinely had a low frequency).

These sample sizes are intended to inform investigators of the number of children to be enrolled to adequately power single-arm studies on these antibiotic classes having safety as a primary endpoint.

**WHAT NEXT?**

The WG has discussed general principles for the design of studies and put forward practical suggestions on clinical inclusion/exclusion criteria for children for antibiotic trials, where none previously existed. We have also put forward suggestions on how to improve safety reporting.

The collaboration between clinical academic CT networks and pharma is improving. However, one of the barriers in conducting CTs in children is the complexity of the inclusion/exclusion criteria which can be a barrier to recruitment. The group therefore attempted to draft criteria for each CIS that would be as simple and inclusive as possible, to try and encourage as wide an adoption by both clinicians and industry, relying on the fact that investigators are keen to use widely recognised criteria when available. In this process, the regulatory agency will have the responsibility for the approval of CTs designed for obtaining a marketing authorisation for a new molecular entity; on the other side, the sponsor will have the responsibility to ensure that the protocol is designed to be as efficient as possible and reflects the relevant current guidance documents for that specific clinical infection.

Considering the limited data currently available on paediatric pharmacology, it is clear that robust evidence of efficacy and safety of different drugs in children can only be gained if CTs are properly conducted and reported. This issue was raised by Saint-Raymond et al who suggested additional reporting requirements to the 2010 Consolidated Standards of Reporting Trials Statement specifically for trials in children. Data on neonates are even more limited. To improve the quality of reporting and strengthen research in this age group, an extension of the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement for neonatal infection research has been published recently—STROBE-Neonatal Infection. This would help the process of harmonisation in data collection and reporting, therefore increasing translation of results into clinical practice.

In summary, this document is intended to be complimentary to the draft EMA ‘Addendum to the guideline on the evaluation of medicinal products indicated for treatment of bacterial infections to address paediatric-specific clinical data requirements’. The WG focused on those aspects not specifically addressed in the draft Addendum, gathering evidence from both published literature and experience from the networks and members involved. The next step is further discussion internationally with investigators, paediatric CTs networks and regulators, and to work towards a wider harmonisation of trial design and conduct.

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