Assessment of flomoxef combined with amikacin in a hollow-fibre infection model for the treatment of neonatal sepsis in low- and middle-income healthcare settings

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Background: Annual mortality from neonatal sepsis is an estimated 430 000–680 000 infants globally, most of which occur in low- and middle-income countries (LMICs). The WHO currently recommends a narrow-spectrum β-lactam (e.g. ampicillin) and gentamicin as first-line empirical therapy. However, available epidemiological data demonstrate high rates of resistance to both agents. Alternative empirical regimens are needed. Flomoxef and amikacin are two off-patent antibiotics with potential for use in this setting.

Objectives: To assess the pharmacodynamics of flomoxef and amikacin in combination.

Methods: The pharmacodynamic interaction of flomoxef and amikacin was assessed in chequerboard assays and a 16-arm dose-ranged hollow-fibre infection model (HFIM) experiment. The combination was further assessed in HFIM experiments mimicking neonatal plasma exposures of clinically relevant doses of both drugs against five Enterobacterales isolates with a range of flomoxef/amikacin MICs.

Results: Flomoxef and amikacin in combination were synergistic in bacterial killing in both assays and prevention of emergence of amikacin resistance in the HFIM. In the HFIM assessing neonatal-like drug exposures, the combination killed 3/5 strains to sterility, (including 2/5 that monotherapy with either drug failed to kill) and failed to kill the 2/5 strains with flomoxef MICs of 32 mg/L.

Conclusions: We conclude that the combination of flomoxef and amikacin is synergistic and is a potentially clinically effective regimen for the empirical treatment of neonatal sepsis in LMIC settings and is therefore suitable for further assessment in a clinical trial.

Introduction

Despite improvement in neonatal mortality in recent decades,1 neonatal sepsis continues to contribute significant global mortality, with an estimated 430 000–680 000 deaths per annum, with a majority of deaths occurring in low- and middle-income countries (LMICs).2–4 The WHO currently recommends a narrow-spectrum β-lactam (e.g. benzylpenicillin or ampicillin) in combination with gentamicin for the first-line empirical treatment of neonatal sepsis, with a third-generation cephalosporin (e.g. cefotaxime, ceftiraxone) recommended as second line.5,6 A recent prospective observational neonatal sepsis study in South Asia and sub-Saharan Africa demonstrated resistance rates in Gram-negative bacteria of 95%, 83% and 60% to ampicillin, cefotaxime and gentamicin, respectively.7 Another prospective study in New Delhi demonstrated 56% of Gram-negative bacteria being resistant to ≥3 classes of broad-spectrum antibiotics (i.e. extended-spectrum cephalosporins, piperacillin/tazobactam, fluoroquinolones, aminoglycosides and
carbapenems) and 38% methicillin resistance rates in Staphylococcus aureus. A similar pattern of high resistance rates is seen in other retrospective studies. Alternative antimicrobial regimens for empirical treatment of neonatal sepsis in LMIC settings are urgently required.

Alternative antimicrobial agents for empirical treatment of neonatal sepsis in LMICs should meet the following criteria: (1) be effective against relevant pathogens with resistance mechanisms complicating the current WHO-recommended regimen; (2) be safe and well tolerated; (3) licensed for use in neonates (or with experience of extensive usage); and (4) be affordable.

Amikacin is an aminoglycoside with molecular modifications that render it stable to aminoglycoside-modifying enzymes that inactivate gentamicin. Flomoxef is an oxacephem β-lactam with stability to degradation by non-AmpC ESBLs. Both agents fulfil the above criteria, and each have been demonstrated to have in vitro synergy in combination with fosfomycin. We therefore studied the potential utility of these agents in combination by assessing their in vitro activity, the presence and magnitude of any pharmacodynamic (PD) interaction using dynamic in vitro models, and assessed the potential utility of this candidate combination regimen using clinically relevant drug exposures.

Materials and methods

The methodology is similar to that of previous experimental work assessing the pharmacodynamics of antibiotics in neonates in a pre-clinical setting.

Antimicrobial agents

Pure compounds of flomoxef (Shionogi, Osaka, Japan) and amikacin (Sigma-Aldrich, St Louis, USA) were obtained for all in vitro experiments. Both agents were stored at 2°C–8°C in anhydrous form and were prepared in sterile distilled water prior to any experiment.

Media

CAMHB (Sigma-Aldrich, St Louis, USA) was used as the primary medium in all experiments. Mueller–Hinton agar (MHA) was used in all agar plates. Commercially pre-prepared 20 mL round MHA plates (Fisher Scientific, Waltham, USA) or self-prepared 50 mL square MHA plates (MHA from Sigma-Aldrich; square plates from VWR, Radnor, USA) were used in all experiments. For drug-containing plates, MHA was supplemented with antibiotic and prepared within each antibiotic’s stability limits and stored at 2°C–8°C (1 week for both agents). Drug concentrations in agar were four times the MIC of the specific bacterial strain used in a given experiment.

Bacterial isolates

Isolates were supplied by JMI Laboratories, IHMA, PHE, LGC standards, University of Birmingham, University of Oxford and Royal Liverpool University Hospital. For the initial non-dynamic in vitro experiments, a collection of strains was collated representing a range of possible neonatal sepsis bacterial pathogens and resistance mechanisms in an antimicrobial resistance (AMR)-prevalent environment. In total, this included 10 strains of each of the following: Streptococcus agalactiae, MRSA, Escherichia coli and Klebsiella pneumoniae. All Enterobacteriales were ESBL (nine E. coli and nine K. pneumoniae strains) or carbapenemase producers (one E. coli and one K. pneumoniae strain). Some of these strains were used also in the hollow-fibre infection model (HFIM) based on their MICs. Full details of the isolates are detailed in Table S1, available as Supplementary data at JAC Online, with isolates used in the HFIM detailed in Table 1. All isolates were stored in glycerol at −80°C and subcultured onto two MHA plates for 18–24 h at 37°C prior to each experiment. In each experiment, colonies were suspended in PBS to MacFarland standard 0.5 (1 × 10⁶ cfu/mL) and diluted to the target concentration.

Antimicrobial susceptibility testing

Amikacin and flomoxef MICs for a panel of representative neonatal sepsis bacterial pathogens were determined using the EUCAST broth microdilution methodology. E. coli ATCC 25922 or S. aureus ATCC 29213 were used as quality control isolates in all experiments, interpreted using QC values from EUCAST and the Japanese Society of Chemotherapy. The antibiotic gradient strip assay method was used for amikacin MIC determination from isolates from the hollow-fibre experiment. Briefly, an inoculum of the isolate was made using a suspension of a sweep of colonies into PBS to a McFarland standard of 0.5. A lawn of the inoculum was plated onto an MHA plate and an antibiotic gradient strip (Etest, bioMérieux, Marcy-l’Étoile, France) placed on the plate, which was subsequently incubated for 18–24 h at 37°C before reading.

In vitro PD assays

Chequeboard assays were used on selected strains to assess the PD interaction of the flomoxef/amikacin combination, using a similar method as described previously. Strains were selected based on having MICs ≤32 mg/L and >0.0625 mg/L to both amikacin and flomoxef. A total of 100 μL of antimicrobials in sterile distilled water was added to an 8 x 8 grid on a 96-well plate, with concentration gradients created with 1:2 serial dilutions along each axis, with the final row/column having 0 mg/L of the appropriate drug. Each plate was assembled bespoke to each strain, with the maximum concentration of antimicrobial being 4 × the MIC for that strain. One hundred microtitre of a 1 × 10⁶ cfu/mL inoculum was added to each well of the prepared chequeboard. The well containing 0 mg/mL of each drug acted as the positive control; an additional row of blank MHB on the plate acted as negative control. Plates were incubated for 18–24 h at 37°C before being read by an optical densitometer (Varioskan, Thermo Fisher) at 600 nm. Plates were considered valid if the MIC on the monotherapy rows of the chequeboard were within one dilution of previously determined MICs, the negative controls had no growth, and quantification of the inoculum was within 6 × 10⁵–14 × 10⁶ cfu/mL.

Raw OD readings were normalized to that of the positive control. The readouts were then modelled using Greco’s model of drug synergy using ADAPT 5, 26, 27 The model produced an interaction parameter, α, with 95% CI characterizing the PD interaction; which was interpreted as follows: a lower bound of the CI > 0 indicates synergy; an upper bound of the CI < 0 indicates antagonism; a CI containing 0 indicates additivity only. Meta-analysis was performed on the output of the model between individual strains using the R package ‘Metafor’. 28

Hollow-fibre infection model

The HFIM is a well-established dynamic model simulating the PD effect of antimicrobials with physiological dynamic concentrations. The HFIM method was used largely as described previously. Briefly, each arm in the HFIM is set up as demonstrated in Figure S1; monotherapy arms omitted the supplementary compartments. CAMHB was pumped into the central compartment at a rate set to simulate a physiological clearance rate (80°C and subcultured onto two MHA plates for 18-24 h at 37°C prior to each experiment. In each experiment, colonies were suspended in PBS to MacFarland standard 0.5 (1 × 10⁶ cfu/mL) and diluted to the target concentration.

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Table 1. Details of strains used in the HFIM experiments

| Strain ID | Species                  | Resistance mechanisms                                      | Flomoxef MIC (mg/L) | Amikacin MIC (mg/L) |
|----------|--------------------------|------------------------------------------------------------|---------------------|---------------------|
| ST195    | *E. coli*                | CTX-M-14, aph(3'), aac(3), TEM-OSBL, mdf(A)                | 1                   | 4                   |
| SPT 731  | *E. coli*                | CTX-M-3, TEM, ST131, O25b                                   | 0.125               | 16                  |
| I1025    | *E. coli*                | mdf(A), ampC promoter mutation                              | 8                   | 4                   |
| 1203217  | *K. pneumoniae*          | SHV-12, CTX-M-9, OXA-48                                    | 0.5                 | 1                   |
| 1280740  | *K. pneumoniae*          | SHV-OSBL, TEM-OSBL, CTX-M-15, DHA-1                        | 32                  | 4                   |
| 1256506  | *K. pneumoniae*          | SHV-OSBL; TEM-OSBL; CTX-M-2; CMY-2                         | 32                  | 2                   |

To account for the difference in clearance between flomoxef and amikacin, supplementary arms were incorporated according to the principles laid out by Blaser.32 Protein binding is negligible for both drugs,36,37 so no adjustment in the administered dosage to account for this was necessary. Biological and technical replicates were not performed due to cost, as is the standard in other published HFIM experiments.36

Preliminary monotherapy experiments were performed with the ESBL-producing ST195 *E. coli* strain (flomoxef MIC 0.125 mg/L; amikacin MIC 4 mg/L; supplied by the University of Birmingham).38 PK and PD outputs of these experiments were modelled using PMetrics38 and parameters simulated using ADAPT27 to determine the flomoxef and amikacin doses required to achieve EC20, EC50 and EC80 in terms of bactericidal effect within the HFIM. A 16-arm HFIM experiment was performed using a 4×4 dosing matrix using these three doses and no dose for both antibiotics in combination. The experiment was run over 96 h, with a target initial inoculum of 1×10⁶ cfu/mL inoculated into the hollow-fibre cartridges. Doses of flomoxef were administered every 12 h to the primary central compartment only; amikacin doses were administered to the primary and supplementary central compartments every 24 h.

PK samples were taken for bioanalysis at four timepoints [pre-dose and 1, 4 and 12 h (for flomoxef)/24 h (for amikacin) post-dose] in dosing windows on Days 1 and 3 of the experiment, with drug concentrations determined via an LC-MS/MS bioanalysis methodology (see Text S1 in Supplementary data for full details). Inoculum samples were taken from each hollow-fibre cartridge at four timepoints (pre-first dose and 2, 4 and 6 h post-first dose) during the first 24 h, then once daily before administration of any doses until the 96 h timepoint. Inoculum concentrations were determined using the track dilution method39 plated onto three MHA plates: one drug-free and two containing either flomoxef or amikacin. An additional 100 μL of the original inoculum was plated onto a drug-free MHA plate to lower the limit of detection for total bacterial quantification (i.e. to 10 cfu/mL). Plates were then incubated at 37°C for 18–24 h for drug-free plates, and 42–48 h for drug-containing plates. After incubation, colonies were counted for at least two dilutions and the cfu/mL of the original inoculum calculated. MICs from any viable colonies from each arm on the final timepoint were determined via antibiotic gradient strip assay for amikacin or broth microdilution MIC methodology (for flomoxef).

Suspected sterility of individual HFIM arms was confirmed at the end of the experiment by centrifugation of contents of the hollow-fibre cartridge at 3000 rpm for 10 min. The supernatant was discarded, and remainder of the sample resuspended in 1 mL MHB before being plated onto MHA plates and incubated for 24 h at 37°C.

Modelling
Population PK models were constructed using the PK and PD outputs of the 16-arm HFIM experiment using the population PK program PMetrics using a non-parametric adaptive grid (NPAG) estimation routine38 and a structural model was based on Greco’s models of pharmacological synergy39 (see Text S2 for details).

Results
In vitro susceptibility testing
The flomoxef and amikacin MICs were determined for a panel of strains representative of pathogens with identified resistance mechanisms pertinent to LMICs (Table 2). The modal flomoxef MIC was 0.25 mg/L. Of the six Enterobacteriales strains with flomoxef MICs ≥8 mg/L, 3/6 carried a plasmid-borne gene encoding an AmpC enzyme (e.g. CMY-II); 1/6 was a carbapenemase (KPC3) producer; 1/6 carried an AmpC promoter mutation; and 1/6 had no identified relevant resistance mechanism. The flomoxef MICs were ≤0.5 mg/L for all Streptococcus agalactiae, and ≤4 mg/L for 9/10 MRSA strains. The modal MIC for amikacin was 4 mg/L (excluding the intrinsically amikacin-resistant *S. agalactiae*, which had a modal amikacin MIC of >32 mg/L).

In vitro drug–drug interaction modelling
Static chequerboard assays were performed on strains with both amikacin and flomoxef MICs >0.0625 mg/L and ≤32 mg/L (n = 16). A mathematical model of drug interaction originally described by Greco39 was fitted to the observed data to estimate...
Darlow et al.

The combination demonstrated evidence of synergy for 8/16 strains (i.e. $\alpha$ and the 95% CI were $>0$). For the remaining 8/16 strains, the combination was additive (i.e. the 95% CI of $\alpha$ included 0). A meta-analysis of the individual strains demonstrated low heterogeneity between strains and species ($I^2 = 0\%$) with a summary value of $\alpha = 0.2013$ (95% CI 0.0575–0.3451), indicating an overall synergistic interaction.

**PD interaction of flomoxef and amikacin**

In preliminary HFIM experiments, the drug exposures of both agents that produced the EC$_{20}$, EC$_{50}$, and EC$_{80}$ (quantified in terms of maximal bacterial kill) against the CTX-M-14-producing *E. coli* ST195 (flomoxef MIC 0.125 mg/L, amikacin MIC 4 mg/L) were determined as fAUC$_{0–24}$ of 15, 65 and 120 mg·h/L for flomoxef (with $C_{\text{max}}$ values of 2, 8.5 and 15 mg/L) and 45, 190 and 375 mg·h/L for amikacin (with $C_{\text{max}}$ values of 5, 20 and 40 mg/L).

These EC$_{20}$, EC$_{50}$, and EC$_{80}$ exposures for both drugs were used in a 16-arm 4×4 matrix, representing all possible monotherapy and combination regimens to explore the PD interaction between the two agents (Figure 2). Increasing exposures of flomoxef (as monotherapy) resulted in rapid bacterial killing in the first 24 h. However, none of the arms became sterile for the duration of the experiment [Figure 2(b–d)], although emergence of flomoxef resistance was not observed in the arm despite this. Increasing amikacin monotherapy saw an exposure-dependent decline of bacterial growth and progressive exposure-dependent emergence of amikacin resistance [Figure 2(e, i and m)]. All combination arms [Figure 2(f–h; j–l; n–p)] resulted in a greater magnitude of bacterial kill compared with comparable monotherapy doses without emergence of resistance to either amikacin or flomoxef.

A PD interaction model based on the model described by Greco$^{26}$ was fitted to the data. The mean and median parameter estimates, and their dispersions, are summarized in Table 3. The coefficient of determination values for observed-versus-individual predicted values (using mean parameter estimates) were 0.985 (free flomoxef concentrations), 0.981 (free amikacin concentrations), 0.858 (total bacterial count), 0.875 (flomoxef-resistant bacterial count) and 0.923 (amikacin-resistant bacterial count). The mean $\alpha$ interaction parameter values were 54.96 (95% credibility interval 47.69–74.50) for bacterial kill and 35.66 (95% credibility interval 11.05–

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**Table 2. Flomoxef (top) and amikacin (bottom) MIC distributions for panel of 40 representative bacterial strains**

| Bacterial species | Flomoxef MIC (mg/L) | Amikacin MIC (mg/L) |
|-------------------|---------------------|---------------------|
|                   | ≤0.0625 | 0.125 | 0.25 | 0.5 | 1 | 2 | 4 | 8 | 16 | 32 | >32 |
| *E. coli*         | 1       | 4     | 2    | 1   | — | — | — | 1  | —  | —  | 1   |
| *K. pneumoniae*   | —       | 1     | 4    | 1   | — | — | — | 1  | —  | —  | 2   |
| MRSA              | —       | —     | —    | 1   | 1 | 1 | 5 | —  | 1  | —  | —   |
| *S. agalactiae*   | 1       | 1     | 7    | 1   | — | — | — | —  | —  | —  | —   |

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**Figure 1.** Fitted $\alpha$ values from the Greco model using chequerboard assays outputs for 16 strains. A total summary statistic using a meta-analysis of the 16 strains is demonstrated in the final raw (‘Total’) with numerical values given above the figure.
53.78) for prevention of amikacin resistance emergence. Modelled experimental time–concentration and PD profiles can be seen in Figures S2 and S3. An interpretable value of α could not be estimated for prevention of flomoxef resistance given the lack of flomoxef resistance emergence in flomoxef-containing arms.

**Assessment of flomoxef and amikacin using neonatal regimens**

The pharmacodynamics of the flomoxef/amikacin combination replicating exposures of candidate neonatal regimens (i.e. IV 15 mg/kg q24h for amikacin and IV 20 mg/kg q12 h for flomoxef)11,32 were studied using five Enterobacterales strains with a range of flomoxef and amikacin MICs (Table 1).

The final end-experiment PD outcomes from each arm are shown in Figure 3 (modelled time–concentration profiles are shown in Figure S4 with full PD outputs shown in Figures S5–S9). Flomoxef monotherapy resulted in sterilization of *E. coli* strain SPT 731 (flomoxef MIC 0.125 mg/L) but not for all other strains. Amikacin monotherapy failed to sterilize any strain. The combination regimen sterilized 2/4 strains not killed by either monotherapy. Neither of the two strains with a flomoxef MIC value of 32 mg/L were sterilized by the combination.

Colonies retrieved from end-experiment non-sterile arms following use of a flomoxef- or amikacin-containing regimen (i.e. either monotherapy or combination) had an increase in MIC by ≥4-fold to the respective agent.

**Discussion**

In our experimental work, we have demonstrated that flomoxef and amikacin in combination provide a synergistic bactericidal effect (Figures 2 and S5–S9) and extends the spectrum of activity to strains that would otherwise not be successfully killed by monotherapy with either agent (Figures S6 and S7). The combination also synergistically prevents emergence of amikacin resistance, although we did not determine the effect of amikacin on protecting flomoxef resistance.

The pharmacodynamics of amikacin monotherapy in the HFIM is similar to previous work.21 Its relatively poor performance when administered as monotherapy may result from the emergence of small-colony variants at a greater rate in vitro than in vivo.41 These experimental data also support the recent downward revision of amikacin breakpoints by EUCAST and the new recommendation to avoid use of amikacin (and other aminoglycosides) as monotherapy in systemic infections.42

As monotherapy, flomoxef was not able to cause any overall bacterial kill in strains with significant production of AmpC. Significant kill and sterilization occurred only in strains with flomoxef MICs ≤0.5 mg/L. Combined with results from similar

![Figure 2](image_url)
In the so-called ESCPM organisms (Enterobacter spp., Serratia spp., Citrobacter freundii, Providencia spp. and Morganella spp.) these chromosomal genes are quickly derepressed to produce clinically significant levels of AmpC. In other Enterobacteriales, chromosomal ampC is either absent (e.g. Klebsiella spp.), or poorly expressed (e.g. E. coli). The latter group can acquire mutations in ampC promotors leading to clinically significant AmpC production. Additionally, all Enterobacteriales can acquire plasmid-borne ampC (e.g. blaCMY or blaOXA), which causes clinically significant production of AmpC.

Flomoxef resistance is primarily driven by the production of AmpC enzymes and carbapenemases in Enterobacteriales. Resistance to flomoxef also results from one or more additional mechanisms including porin loss and, most likely, efflux, as with other β-lactams. However, the specific characterization of these mechanisms for flomoxef is limited and their contribution to resistance requires further study. Where characterizations are made (e.g. loss of OmpF or OmpK35 porins), the effect on flomoxef MIC is relatively minor. Nevertheless, it remains likely the overall efficacy of flomoxef monotherapy will be directly related to the epidemiology of AmpC and carbapenemases.

Many Gram-negative bacteria contain chromosomal ampC genes. In the so-called ESCPM organisms (Enterobacter spp., Klebsiella spp., and Serratia spp.), these chromosomal genes are quickly derepressed to produce clinically significant levels of AmpC. In other Enterobacteriales, chromosomal ampC is either absent (e.g. Klebsiella spp.), or poorly expressed (e.g. E. coli). The latter group can acquire mutations in ampC promotors leading to clinically significant AmpC production. Additionally, all Enterobacteriales can acquire plasmid-borne ampC (e.g. blaCMY or blaOXA), which causes clinically significant production of AmpC.

The recent BARNARDS study showed ESCPM organisms accounting for 247/1038 (23.8%) of Gram-negative bacteria and 247/2483 (9.9%) of total isolated bacteria from LMIC neonatal sepsis patients. These rates are likely overestimates of the true prevalence due to outbreaks of clonal Serratia spp. at one of the study sites. Plasmid-mediated ampC genes and carbapenemase genes were present in a further 17/258 (6.6%) and 69/258 (26.7%) of total isolated bacteria from LMIC neonatal sepsis patients. These rates are likely overestimates of the true prevalence due to outbreaks of clonal Serratia spp. at one of the study sites. WGS data from the NeoOBS study suggest plasmid-borne ampC carriage rates of 7% and 2.5% in isolated K. pneumoniae and E. coli isolates, respectively. WGS data from the NeoOBS study suggest plasmid-borne ampC carriage rates of 7% and 2.5% in isolated K. pneumoniae and E. coli isolates, respectively. Whilst there is regional variation, these point estimates suggest that the prevalence of AmpC-producing strains is high enough in these LMIC settings that a combination of flomoxef with amikacin would be beneficial compared with flomoxef monotherapy for the empirical treatment of neonatal sepsis in these settings.

There are several considerations for using this combination regimen. As monotherapy, both flomoxef and amikacin are safe and well tolerated. Whilst it is likely that combination therapy will be safe (given the experience of both agents as monotherapy, and the experience of ampicillin and gentamicin in combination), it is possible there may be unanticipated drug-
drug interactions and potentiated toxicities when the drugs are used in combination. Additionally, the CSF penetration is relatively low for both agents, with neonatal CSF partition coefficients of 0.05 and 0.1 estimated for flomoxef and amikacin, respectively.\textsuperscript{50,51} These numbers compare favourably with the WHO regimen of ampicillin and gentamicin (CSF partition coefficients of \(\sim 0\) for gentamicin and 0.05–0.1 for ampicillin).\textsuperscript{52–54} but our work did not model the pharmacokinetics within CSF, nor the pharmacodynamics within the CNS, and we therefore cannot comment on the efficacy of this regimen for the treatment of CNS infections.

There are potential limitations of this work. First, the HFIM does not contain or replicate immunological effectors. Secondly, the inoculum that was used in the HFIM is higher than that found in neonatal sepsis (\(10^5\)–\(10^6\) cfu/mL compared with the estimated bacterial density in neonatal sepsis of \(10^5\)–\(10^3\) cfu/mL).\textsuperscript{55,56} Both limitations are likely to underestimate the predicted in vivo effect of the combination regimen. A smaller bacterial inoculum and present immune effectors (even if immature) in individuals with neonatal sepsis would likely lead to a greater overall effect of the regimen than seen in vitro here. The conclusions drawn from these experiments are therefore conservative and the HFIM likely represents a worst-case scenario.

Overall, we conclude that this experimental study has demonstrated this combination regimen is synergistic, in terms of both bactericidal effect and protection against amikacin resistance, and potentially efficacious in the empirical treatment of neonatal sepsis caused by ESBL-producing Enterobacteriales species. Consequently, we believe this regimen should be considered for clinical assessment as empirical treatment of MDR neonatal sepsis in LMIC settings, where the current standard of care has become decidedly suboptimal.\textsuperscript{21,30}

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### Transparency declarations

Shionogi provided the flomoxef material free of charge, but provided no additional funding. W.H. holds or has recently held research grants with UKRI, National Institutes of Health, National Institute of Health Research, F2G, Spero Therapeutics, Antabio, Pfizer, Bugworks, Phico Therapeutics, BioVersys, CARBX, GARDP and NAEJA-RGM. He is (or has recently been) a consultant for Appili Therapeutics, F2G, Spero Therapeutics, NAEJA-RGM, Centauri, Pfizer, Phico Therapeutics and VenataRx. He is a member of the Scientific Advisory Committee for GARDP and the Specialty National Co-lead for Infectious Diseases for the National Institute of Health Research (NIHR). S.D. holds or has recently held research grants with F2G, Spero Therapeutics, Antabio, Pfizer, Bugworks, Phico Therapeutics, BioVersys, CARBX, GARDP and NAEJA-RGM. She is (or has recently been) a consultant for Spero Therapeutics, Centauri, Pfizer and CARBX. She is a member of EUCAST. There is nothing further to declare for all other authors.

### Data availability

The programs ADAPT and PMetrics are publicly available, with instructions, at https://bmsr.usc.edu/software/adapt/ and http://www.lapk.org/ pmetrics.php, respectively.

### Supplementary data

Table S1, Figures S1 to S9 and Texts S1 to S2 are available as Supplementary data at JAC Online.

### References

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