Clinical pharmacology of cefuroxime and cefuroxime axetil in infants and children

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Abstract
Cefuroxime is a second-generation cephalosporin β-lactamase resistant active against gram-negative and gram-positive organisms and the prodrug cefuroxime axetil is the drug of choice for treatment of Lyme disease. Cefuroxime inhibits bacterial cell wall synthesis, is bactericidal, and is one of the few cephalosporins available for oral, intravenous, and intramuscular administration. Cefuroxime is rapidly absorbed when given by mouth, and its blood concentrations remarkably vary among subjects, but the drug has not adverse-effects even when is given at high doses. The gastrointestinal absorption-rate of cefuroxime axetil is 70%, and after hydrolysis it relays cefuroxime. Parenteral dosing-regimens consist of 25 mg/kg in infants and dosing intervals decrease with increasing postnatal age. Cefuroxime oral dosing-regimens are: 10, 20 mg/kg and 250 mg twice-daily in children aged up to one year, 2 to 11 and 12 to 17 years, respectively. Elimination half-live are: 6 hours in the first week of life and 1.8 hours in children aged up to 6 years. Cefuroxime is efficacy and safe for prophylaxis and treatment of respiratory-tract and acute urinary-tract infections, pneumoniaeae, acute otitis media, and sinusitis in children. Prophylaxis for surgical procedures requires 50 mg/kg before procedure and then 30 mg/kg for up 3 days after intervention. Cefuroxime concentration in cerebrospinal fluid is about 10% of that in plasma and the drug successfully treated meningitis caused by β-haemolytic streptococci, Streptococcus pneumoniae, and Neisseria meningitides, but when Salmonella species are the infective pathogens additional 5 mg dose is given intraventricularly. Some bacteria may become resistant to cefuroxime and antibiotic consumption is associated with increased bacterial-resistance. The aim of this study is to review the published data on cefuroxime and cefuroxime axetil dosing, efficacy, safety, and cefuroxime effects, prophylaxis, treatment, optimization, meningitis, pharmacokinetics and bacterial-resistance.

Introduction
This non-toxic broad-spectrum cefuroxime is one of the few cephalosporins that may be administered parenterally and orally. Cefuroxime is a β-lactamase resistant second-generation cephalosporin active against gram-positive organisms (including group B streptococci and penicillin-resistant staphylococci) and a wide range of gram-negative organisms. It is reasonable active against Haemophilus influenzae and Neisseria gonorrhoeae, but is inactive against Listeria, enterococci, Bacteroides, and Pseudomonas species. It has sometimes been used prophyilactically in infants undergoing abdominal surgery. Cefuroxime penetrates into the cerebrospinal poorly and coagulase-negative staphylococci are increasingly resistant to this antibiotic. Two-thirds of the lipophilic acetoxyethyl ester dose, cefuroxime axetil, is absorbed when is given orally, and after hydrolysis, it releases cefuroxime. There are no published reports of the use of this formulation in children aged < three months, but it has been widely used to treat otitis media and other respiratory infections in older children. It is just as active as treatment with co-amoxiclav and less likely causes troublesome loose stools. Cefuroxime is largely excreted by the kidney. Elimination half-life in plasma falls from six hours at birth to about three hours at two weeks. Infants aged > one month clear cefuroxime from their plasma almost as fast as adults (half-life one hour), but dosage intervals should be extended in infants with severe renal failure. Toxic adverse-effects are rare but oral treatment does sometimes cause nausea and vomiting and a change in stool frequency, and pseudomembranous colitis has occasionally been reported. Other problems, as with cephalosporin in general, are uncommon. Lyme disease is caused by the spirochete Borrelia burgdorferi and human infection is caused by the bite of an infected animal tick. Illness was once rare in the United Kingdom but does not seem common in Europe and North America. While a migrating annular skin lesion (erythema migrans) is the classic presentation, symptoms are very variable. Foetal infection was first recognized in 1985, and it is now realized that the risk to the foetus is comparable to that of congenital syphilis. Cefuroxime axetil is the antimicrobial agents of choice for early disease in young children. Intravenous ceftriaxone, cefotaxime, or benzylpenicillin are recommended if there are cardiac or neurological complications. The duration of treatment is usually two to four weeks [1,2].

Cefuroxime inhibits bacterial cell wall in a manner similar to that of penicillin [3]. Peptidoglycan is a heteropolymeric component of the bacterial cell wall that provides rigid mechanical stability. β-Lactam antibiotics inhibit the last step in the peptidoglycan synthesis, the cell wall is 50 to 100 molecules thick in gram-negative bacteria and it is only 1 or 2 molecules thick in gram-negative organisms. The peptidoglycan is composed by glycan chains, which are linear strands of two alternating amino sugars (N-acetylglucosamine and N-acetylglucosamuramic) that are cross-linked by peptide chains. Peptidoglycan precursor formation takes place in the cytoplasm. The synthesis of UDP-acetylmuramylpentapeptide is completed with the addition of a dipeptide, D-alanyl-D-alanine (formed by racemization and condensation of L-alanine). UDP-

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acetylmuramyl-pentapeptide and UDP-acetylglucosamine are linked (with the release of the uridine nucleotides) to form a long polymer. The cross-link is completed by a transpeptidation reaction that occurs outside the cell membrane. The β-lactam antibiotics inhibit last step in peptidoglycan synthesis, presumably by acylating the transpeptidase via cleavage of the –CO-N- bond of the β-lactam ring. The targets for the actions of β-lactam antibiotics are collectively termed penicillin-binding-proteins. The transpeptidase responsible for synthesis of the peptidoglycan is one of these penicillin-binding-proteins. The lethality of penicillins for bacteria appears to involve both lytic and nonlytic mechanisms [2]. Cefuroxime has good activity against Haemophilus influenzae Neisseria meningitides and Streptococcus pneumonia. Activity against Escherichia coli and Klebsiella species is modest. Anti-staphylococcal activity is inferior to first-generation cephalosporins. Unlike cefoxitin, cefotetan, and cefmetazole, cefuroxime lacks activity against Bacillus fragilis. Cefuroxime can be given orally, intravenously, or intramuscularly twice-daily to thrice-daily in children and adult patients. Concentrations in cerebrospinal fluid are about 10% of those in plasma, and the drug is effective but inferior to ceftriaxone for treatment of meningitis due to susceptible organisms [4].

Children aged 12 to 17 years: give 250 mg twice-daily, and the dose intervals may be doubled for severe lower respiratory-tract infections and for pneumonia [3].

Cefuroxime administered parenterally
Give 20 mg/kg thrice-daily (maximum per dose=750 mg); increase the dose up to 50-60 mg/kg thrice-daily or 4 times-daily (maximum per dose=1.5 grams), to treat severe infections and cystic fibrosis [3].

Cefuroxime administered orally to treat lower urinary-tract infection

Children aged 12 to 17 years: give 125 mg twice-daily [3].

Cefuroxime administered by intravenous injection for surgical prophylaxis
Give 50 mg/kg (maximum per dose=1.5 grams) administered 30 min before procedure, then 30 mg/kg thrice-daily (maximum per dose=750 mg) if required for up to three doses for high-risk procedures [3].

Cefuroxime axetil oral treatment schedules to treat Lyme disease

Children aged 3 months to 11 years: give 15 mg/kg twice-daily (maximum per dose=500 mg) for 14 to 21 days (for 28 days to treat Lyme arthritis) [3].

Children aged 12 to 17 years: give 500 mg twice-daily for 14 to 21 days (for 28 days to treat Lyme arthritis) [3].

Literature search
The literature search was performed electronically using PubMed database as search engine and the cut-off point was June 2020. The following key words: "cefuroxime infants effects", "cefuroxime children effects", "cefuroxime infants metabolism", "cefuroxime children metabolism", "cefuroxime infants pharmacokinetics", "cefuroxime children pharmacokinetics", "cefuroxime infants resistance", and "cefuroxime children resistance" were used. In addition, the books Neonatal Formulary [1], The British National Formulary for Children [3], and The Pharmacological Basis of Therapeutics [4] were consulted. The manuscript is written according to "Instructions for Authors".

Results

Cefuroxime efficacy and safety in infants and children

Powell et al. [6] assessed the efficacy, safety and tolerability of cefuroxime axetil suspension in 36 infants and children, aged 3 months to 12 years, and doses were: 10, 15, or 20 mg/kg twice-daily or thrice-daily. Three of 36 patients (8.3%) were withdrawn from therapy because of adverse-effects. Of 32 patients who completed therapy, 9 (28.1%) developed mild-reactions including oral thrust, diarrhoea, or diaper dermatitis, and none were withdrawn from therapy. Complete clinical cure occurred in 28 subjects (87.5%), and 4 subjects (12.5%) were clinically improved but still required an additional antibiotic after one week to complete therapy, and treatment was effective, safety and well tolerated.
Reed et al. [7] studied efficacy, safety and tolerability of cefuroxime axetil suspension in 35 children, aged 3 months to 12 years. The drug was administered at doses of: 10, 15, or 20 mg/kg twice-daily or thrice-daily. Three of 35 subjects (8.6%) were withdrawn from therapy because of drug-related reactions. Of 32 subjects who completed therapy, 9 (28.1%) developed mild oral thrush, diarhoea, or diaper dermatitis; but none were withdrawn from therapy and this treatment was efficacious, safe, and well tolerated.

Barson et al. [8] treated 36 children, aged 3.5 to 4.7 years, suffering from infections in the face, epiglottis, or buccal cellulitis who were treated with cefuroxime at a dose of 75 mg/kg daily. Blood culture was performed in 22 children (61.1%) and the infective pathogens were: Haemophilus influenzae type b, Streptococcus pneumoniae, and nontypeable Haemophilus influenzae, and this treatment was efficacy and safe.

Syrogiannopoulos et al. [9] administered cefuroxime axetil at a dose of 250 mg twice-daily for 10 days to 55 children, aged 5 to 11 years, suffering from acute otitis media. A tympanogram was obtained on enrolment and 2 days after completion of therapy. Complete cure occurred in 74.1% children and clinical improvement was observed in 25.9% children. There was no clinical failures and none children experienced relapse of acute otitis media infection one month after end of therapy.

Olivier [10] observed that treatment of community-acquired pneumonia caused by Mycoplasma, Chlamydia, or Legionella species required a drug which penetrates respiratory tissues effectively such as cefuroxime, and she reviewed literature on treatment of this disease. Cefuroxime axetil has been extensively used in infants aged > 3 months, suffering from acute otitis media. A tympanogram was obtained on enrolment and 2 days after completion of therapy. Complete cure occurred in 74.1% children and clinical improvement was observed in 25.9% children. There was no clinical failures and none children experienced relapse of acute otitis media infection one month after end of therapy.

Burman et al. [11] observed that Enterobacter cloacae strains dominated aerobic flora in faeces of 79 of 953 infants (8.3%) discharged from neonatal intensive care units. Cefuroxime yielded slight increase of MICs of ampicillin, cephalexin, and cephalothin compared to isolates obtained from untreated infants (P-value=0.02).

Zachariassen et al. [12] investigated cefuroxime pharmacokinetics in infants born to mothers undergoing caesarean section who received cefuroxime for prophylaxis. The drug migrated into infants and its plasma concentration and elimination half-life remarkably varied in infants.

Dierikx et al. [13] assessed the effect of cefuroxime on bacterial composition in gut of infants born to mothers treated with cefuroxime at single dose of 1.5 grams during, prior caesarean section and after umbilical clamping. Cefuroxime migrated into infants and impaired gut intestinal microbiota.

Arnez et al. [14] determined efficacy and adverse-effects caused by cefuroxime axetil administered at a dose of 30 mg/kg daily to 46 children, aged<15 years, suffering from erythema migrans. Efficacy and adverse-effects were compared to those of 44 children treated with penicillin at a dose of 100,000 IU/kg daily. Both treatments were equally efficacious, whereas adverse-effect-rates were: 26.1% and 6.8% (P-value=0.0301) following cefuroxime axetil and penicillin administration, respectively.

Cefuroxime adverse-effects in children

In literature there is only one study on adverse-effects caused by cefuroxime in children and it was reported by Gold and Rodriguez [15], they stated that cefuroxime is free from adverse-effects.

Cefuroxime and cefuroxime axetil prophylaxis for urinary tract infection and surgery in infants and children

Gurevich et al. [16] observed that recurrent urinary-tract infections occurred in 19.8% infants, aged<1 month, and the first episode of infection occurred in 6 infants (5.9%); peak incidence occurred 2 to 6 months after birth. One-hundred-one infants, aged<2 months, with the first and recurrent episodes of urinary-tract infections were treated with cefuroxime axetil. Infective pathogens were: Escherichia coli, Klebsiella species, Enterococcus species, Morganella morganii, Proteus and Enterobacter species. Prophylaxis significantly reduced the frequency of urinary-tract infection episodes up to 8.3%.

Bitsori et al. [17] determined effects of prophylaxis in 638 uropathogens isolated from the urine of children. Infective uropathogens were: Escherichia coli, Klebsiella species, Pseudomonas aeruginosa, Enterococcus and Proteus species. Bacterial-resistance rate of these organisms to several antibiotics increased following their administrations, whereas that to cefuroxime decreased (P-value<0.016). Continuous surveillance, re-evaluation of empiric dosing regimens, and further assessment of prophylaxis are needed to optimize prophylaxis of urinary-tract infection.

Bi et al. [18] characterized pathogens and their antibiotic susceptibility in 895 isolated obtained from children with urinary-tract infections caused by catheter indwelling in order to optimize prophylaxis. Escherichia coli was the most frequent pathogen, followed by Staphylococcus aureus, Staphylococcus epidermidis, and Enterococcus species. Gram-negative species were the predominant infective uropathogens and bacterial-susceptibility patterns must be determined in order to optimize prophylaxis.

Knoderer et al. [19] evaluated efficacy of two prophylaxis regimens with cefuroxime in 210 children, aged ≤ 18 years, undergoing cardiovascular surgery. Prolonged prophylaxis was performed before surgery and short-term prophylaxis was performed on the first day after intervention. Short-term post-operative prophylaxis did not increase infection outcomes.

Cefuroxime treatment of infants and children

Brogden et al. [20] stated that staphylococci β-lactamase producing and gram-negative bacteria are not resistant to ceftriaxone. This antibiotic was given parenterally to infants and children and effectively cured infections caused by gram-positive and gram-negative aerobes. Cefuroxime was also efficacy to treat respiratory infections caused by Haemophilus influenzae, Streptococcus pneumoniae, Klebsiella pneumoniae cephalosporin-resistant and Enterobacter.

Nelson et al. [21] administered cefuroxime intravenously at a dose of 25 mg/kg thrice-daily to 100 infants, with mean age of 15 months, suffering from pneumoniae caused by Streptococcus pneumoniae, Haemophilus influenzae type b, Staphylococcus aureus, Neisseria meningitides B and pneumococcus species. All microorganisms were susceptible and had a MIC ≤ 1.25 μg/mL. Mean duration treatment was 3.1 days in afebrile infants and 5.1 days until respiratory symptoms were gone. Eosinophilia occurred in only 10 infants (10.0%), and cefuroxime successfully treat pneumonia.
Pichichero [22] stated that antibiotic-resistance, caused by antecedent antibiotic treatment, is increasing in pathogens causing acute otitis media infections in infants. Amoxicillin is the first choice antibiotic to treat acute otitis media although a high dose of 80 mg/kg is needed to eradicate resistant-pathogens. Oral cefuroxime is a second-line antibiotic to treat this disease.

Shafina et al. [23] enrolled 101 infants and children, aged 1 day to 13 years, suffering from acute urinary-tract infections caused by Escherichia coli, Klebsiella and Enterococcus species which were treated with ampicillin. The resistance-rate to ampicillin was high for Escherichia coli, Klebsiella and Enterococcus whereas that to cefuroxime was low and cefuroxime is an appropriate antimicrobial agent for treatment of acute urinary-tract infections caused by these bacteria.

Wald [24] stated that Streptococcus pneumoniae, Haemophilus influenzae, and Moraxella catarrhalis are the commonest organisms causing acute and subacute sinusitis in infants and children. Amoxicillin is the appropriate treatment for infections caused by pathogens producing β-lactamase. If a patient does not respond to this treatment cefuroxime-axetil should be used as it has a broad-spectrum of activity.

Cefuroxime administered to mothers before caesarean section causes alteration of microbiota composition in gut of their infants

Kamal et al. [25] assessed alteration of microbiota in infant gut caused by cefuroxime prophylaxis performed in pregnant women undergoing caesarean section. Faecal samples were collected 10 days after birth and also after 9 months. Composition of gram-positive cocci and Enterobacteriaceae was assessed in infant gut born to these mothers. No alteration of microbiota composition was observed 10 days after birth whereas bacteria composition was altered 9 months later.

Optimization of cefuroxime and cefuroxime axetil treatments in children

Carlier et al. [26] optimized treatment with cefuroxime at a dose of 750 mg thrice-daily in 20 critically ill children who had creatinine clearance<20 ml/min. Optimization of treatments were performed according to different cefuroxime dosing-regimens and also to creatinine clearance of: 50, 100, 200 and 300 ml/min. Five-hundred children were simulated by Monte Carlo method for optimizing dosing-regimens which was defined as the blood concentration > MIC for at least 65% of dosing interval (65% dosing-regimen which was defined as the blood concentration > MIC). Sixty-seven isolates (3.3%) were β-Lactamase producing and the infection sites were: acute otitis media 38%, sinusitis 36%, and lower respiratory-tract 21%. Among 10 antibiotic tested, cefuroxime was the most active agent for prophylaxis and cure of children with acute otitis media caused by Haemophilus influenzae.

Cefuroxime axetil treatment of lymph disease in children

Lyme disease is caused by the spirochete Borrelia burgdorferi. Wormser et al. [29] reported guidelines for treatment of Lyme disease in children. Suggested treatments were cefuroxime axetil at a dose of 15 mg/kg twice-daily (maximum 500 mg) for 14 to 21 days. Recommended duration of treatments was for: erythema migrans (14 to 21 days), acute neurological disease, meningitis, radiculopathy or cranial-nerve (14 to 28 days), cardiac disease 1<sup>st</sup> and 2<sup>nd</sup> degree (14 to 21 days), 3<sup>rd</sup> degree (14 to 213 days), and arthritis (14 to 28 days). Occurrence-rates of the first and multiple lesions of erythema migrans were: 98.3 and 16.8%, respectively, in subjects treated with cefuroxime axetil and the size of the primary lesion was 79+10 cm<sup>2</sup> [30]. Gerber et al. [31] reported the frequency of treatment for infections caused by Borrelia burgdorferi bit and was: fever 24%, fatigue 58%, headache 42%, arthralgia 33%, neck pain 26%, nausea 21%, abdominal pain 17%, diarrhoea 16%, myalgia 16%, sore throat 12%, cough and rhinorrhoea 8%, vomiting and conjunctivitis 5%.

Ceftriaxone penetration into the cerebrospinal fluid of infants and children

Rochette [32] measured the cerebrospinal volume in 54 preterm and term infants, with a postmenstrual age of 30 to 60 weeks, and the volume was 1.97+0.54 ml/kg. The Pearson's rank correlation test revealed that cerebrospinal fluid volume correlated with both the infant weight and the postmenstrual age (r=0.78, P-value<0.0001).

Pfenninger et al. [33] enrolled 33 infants, aged > 3 months, suffering from meningitis caused by: Haemophilus influenzae, Neisseria meningitides, and Streptococcus pneumoniae. Cefuroxime was intravenously infused at a dose of 50 mg/kg 4-times-daily, the duration of treatment was 12 to 13 days, and cerebrospinal fluid concentration ranged from 1.1 to 18.8 µg/ml (mean, 7.0).

Edwards et al. [34] studied 7 children, aged 7 months to 9 years, suffering from meningitis caused by Staphylococcus epidermis, Staphylococcus aureus, Staphylococcus hominis, and Staphylococcus sciuri. Cefuroxime was intravenously infused at doses of 67 and 76 mg/kg thrice-daily for 2 to 14 days; cefuroxime crossed the brain blood

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barrier easily and achieved concentrations from <2.0 to 22.5 µg/ml in cerebrospinal fluid. The wide interindividual variability of cefuroxime concentration reflected both the site of local inflammation and the time after dosing.

Kuzemko and Walker [35] measured cefuroxime concentration in cerebrospinal fluid of 6 children, aged 1.5 to 6 years, with meningitis caused by Neisseria meningitides (MIC=0.025 µg/ml), Haemophilus influenzae (MIC=0.5 µg/ml), and Streptococcus pneumoniae (MIC was not determined). Cefuroxime was intravenous infused at a dose of 25 mg/kg twice-daily in three children and other 3 children received this drug by lumbar puncture. Cefuroxime concentration ranged from 1.5 to 6.0 µg/ml in cerebrospinal fluid and was > MICs for Neisseria meningitides and Haemophilus influenzae indicating that cefuroxime is an appropriate antimicrobial agent to cure meningitis caused by these organisms.

Ceftriaxone treatment of bacterial meningitis in infants and children

Sirinavin et al. [36] intravenously infused cefuroxime at a dose of 50 mg/kg daily for 14 to 15 days to 7 infants and children, aged 25 days to 12 years, suffering from meningitis caused by β-haemolytic streptococci, Streptococcus pneumoniae, Neisseria meningitides, Salmonella krefeld and Salmonella typhimurium. Cerebrospinal fluid and blood specimens were sampled 2 hours after dosing for the fourth or successive treatment days. Other 6 infants were suffering from meningitis caused by Haemophilus influenzae and all were cured. Five infants of these were initially treated with cefuroxime co-administered with penicillin bud cefuroxime treatment was discontinued after two days because etiological agents were: hospital-acquired Pseudomonas cepacia and Pseudomonas aeruginosa strains which are cefuroxime-resistant. Cefuroxime concentration was measured in cerebrospinal fluid and serum of other 9 subjects on the second treatment day and was: 6.4±1.7 and 35.5±9.5 µg/ml, respectively, and the ratio was 18.8±3.0%. Cefuroxime concentration was also measured in cerebrospinal fluid and plasma of 6 subjects on the 14th treatment day and was: 3.6±2.2 and 32.2±23.3 µg/ml, respectively, and the ratio was 12.1±2.5%. In children without meningitis, cefuroxime concentrations were: 1.3±1.3 and 32.5±14.7 µg/ml in cerebrospinal fluid and serum, respectively, and the ratio was 3.8±2.4%. Cerebrospinal fluid drug concentration was > 10-fold higher than the MICs of β-haemolytic streptococci, Streptococcus pneumoniae, Haemophilus influenzae, Streptococcus pneumoniae, and Neisseria meningitides, whereas MICs of Salmonella strains were similar to those of cefuroxime. Meningitis caused by β-haemolytic streptococci, Streptococcus pneumoniae, Neisseria meningitides and Haemophilus influenzae type b, was cured with 50 mg/kg cefuroxime given 4 times-daily, whereas meningitis caused by Salmonella species required an additional dose of 5 mg cefuroxime injected intraventricularly for 4 days. Other ten subjects without meningitis received single cefuroxime dose of 50 mg/kg and cerebrospinal fluid concentration was 3 to 6 times lower than those measured in subjects with meningitis, thus infamened m eninges improve cefuroxime penetration-rate into cerebrospinal fluid.

Del Rio et al. [37] evaluated susceptibility of Haemophilus influenzae, Neisseria meningitides, and Streptococcus pneumoniae to cefuroxime in 38 infants and children, aged 1.3±1.3 years (range, 4 weeks to 6.5 years), suffering from meningitis. Twenty-seven subjects (71.0%) had meningitis caused by Haemophilus influenzae type b, (β-lactamase negative and β-lactamase positive), Haemophilus influenzae type e β-lactamase-negative, Streptococcus pneumoniae, Neisseria meningitides, and group B Streptococcus, whereas 4 subjects were not infected. Cefuroxime plasma concentration was measured after single daily intravenous doses of 50 and 75 mg/kg of cefuroxime for a minimum of 5 days in 5 children. Table 1 shows susceptibility of bacteria to cefuroxime. Cefuroxime plasma concentration correlated with the protein content following a dose of 50 mg/kg (P-value=0.01) and 75 µg/ml (P-value=0.039), and the concentration was higher following 75 mg/kg at each time interval. Mean cefuroxime exposition in cerebrospinal fluid, expressed as the AUC in cerebrospinal fluid to that in plasma ratio, ranged from 6.4 to 10.0% following 50 and 75 mg/kg doses, respectively. Cefuroxime concentration (µg/ml) was measured in cerebrospinal fluid on the first and 6th doses at various times after administration and was respectively: 0.96 and 2.1 at 0.5 hours, 17.3 and 14.5 at 1 hour, 2.5 and 3.4 at 2 hours, and 2.1 and 0.94 at 4 hours. Both MICs and MBCs tested for these 5 organisms were lower than cefuroxime concentrations in cerebrospinal fluid suggesting that this drug is an appropriate antimicrobial agent to treat bacterial meningitis caused by these organisms.

Cefuroxime plasma and serum concentrations in infants

Philipson and Sternstedt [38] measured cefuroxime plasma concentration in infants at birth, born to mothers who received cefuroxime at a dose of 750 mg, and it ranged from 1.4 to 3.6 µg/ml. Reference 5 is above reported under the heading cefuroxime efficacy and safety: de Louvois [5] measured cefuroxime serum concentration in 28 preterm and term infants with a body weight of 2.0±0.85 kg who received cefuroxime parenterally at a dose of 25 mg/kg twice-daily. Median concentrations were: 35, 45, 42, 35, and 10µg/ml at: 0.25, 0.5, 1, 3, 5, 12 hours after dosing, and the median elimination half-life was 6.0±2.1 hours.

Cefuroxime pharmacokinetics in infants undergoing cardio pulmonary bypass

Gertler et al. [39] studied cefuroxime pharmacokinetics in 42 infants, aged<1 year, of whom 36 had congenital heart defect and underwent cardio pulmonary bypass, whereas 6 did not. Cefuroxime single dose of 50 mg/kg was intravenously infused before cardiac pulmonary bypass and 75 mg/kg was injected in the prime volume. The impact of age (Fmat) on total body clearance had a sigmoid maturation function and was calculated according to the following equation:

\[ Fmat = \frac{PMA_{\text{HILL}}}{(PMS_{\text{HILL}} + PMA_{50})^{HILL}} \]

Table 1. Susceptibility of 5 bacteria to cefuroxime which were isolated from to cerebrospinal fluid. Susceptibility is expressed as both the minimum inhibitory concentration (MIC) and the minimum bactericidal concentration (MBC). Figures are the range and (median), by Del Rio et al. [37]

| Strain tested                     | Number of isolates | MIC (µg/ml) | MBC (µg/ml) |
|-----------------------------------|--------------------|-------------|-------------|
| Haemophilus influenzae β-lactamase negative | 26 | 0.156 - 0.625 (0.312) | 0.312 - 0.625 (0.312) |
| Haemophilus influenzae β-lactamase positive | 7 | 0.312 - 0.625 (0.312) | 0.312 - 0.625 (0.625) |
| Neisseria meningitides            | 3 | <0.02 - 0.05 | 0.02 - 1.0 |
| Streptococcus pneumoniae          | 2 | 0.02 - 0.156 | 0.02 - 0.156 |
| Group B streptococcus             | 1 | 0.312 | 0.625 |
Cefuroxime pharmacokinetics in infants and children

Reference 37 is above reported under the heading cefuroxime penetration into the cerebrospinal fluid. Del Rio et al. [37] investigated cefuroxime pharmacokinetics in plasma of 5 infants and children on the first and 6th treatment days. Cefuroxime was intravenously infused at doses of 50 and 75 mg/kg. Following a dose of 50 mg/kg, plasma concentrations (µg/ml) were: 105±36 at the end of infusion and 4.4±4.5 6 hours later, and after a dose of 75 mg/kg, these concentrations were: 152±43.5 and 5.4±3.0, respectively. Table 3 summarizes pharmacokinetic parameters obtained in 5 subjects following multiple cefuroxime dosing of 50 mg/kg, on the first and the 6th administration days. Absorption and elimination half-life, distribution volume, and the total body clearance are not statistically different according the two sampling days suggesting that cefuroxime does not accumulate in blood.

Powell et al. [40] assessed cefuroxime axetil pharmacokinetics in 28 infants and children, aged 3 months to 12 years (mean, 1.2 years) with bacterial infection, and were treated with a single cefuroxime axetil suspension at doses of 10, 15, or 20 mg/kg. Table 4 shows that cefuroxime axetil serum peak concentrations were not statistically different according to the 3 doses. In addition, Tmax, AUC, and elimination half-life are not statistically different according the dosages indicating that cefuroxime axetil pharmacokinetic parameters are not dosing-dependent.

Cefuroxime pharmacokinetics in children undergoing cardio pulmonary bypass

Knoderer et al. [41] assessed cefuroxime pharmacokinetics in 16 children, aged 1.2±0.3 years with a body-weight of 9.7±2.8 kg, requiring cardio pulmonary bypass for cardiac surgical repair. Cefuroxime was intravenously infused at a dose of 25 mg/kg one hour before skin incision and another dose of 12.5 mg/kg was injected in the cardio pulmonary bypass prime fluid. The distribution volume of the peripheral compartment was similar to that of the central compartment and this contrasts with the results obtained in infants undergoing cardio pulmonary bypass in whom peripheral distribution volume is larger than the central distribution volume [39]. Likely, total protein content is lower in plasma of infants than that of children, thus the extent of cefuroxime bound plasma proteins is lower in infants with consequent higher migration-rate into body-tissues causing a larger peripheral distribution volume.

Mechanisms bacterial-resistance to cefuroxime

Källman et al. [42] determined cefuroxime susceptibility by disc-diffusion and by E-test methods. The ability of bacteria to grow in

| Parameters estimated in all 42 infants | Typical value | %RSE | Bootstrap: median | %RSE |
|---------------------------------------|---------------|------|-------------------|------|
| *Individual parameters                |               |      |                   |      |
| *Central distribution volume (L/kg)   | 11.2 (8.3)    | 0.178 (10.3) | 11.3 (8.8) | 0.170 (10.4) |
| *Peripheral distribution volume (L/kg)| 17.1 (4.2)    | 0.037 (1.1)  | 17.1 (4.3) | 0.037 (13.5) |
| *Total body clearance (L/kg)          | 3.1 (6.4)     | 0.048 (1.2)  | 5.1 (9.1)  | 0.046 (13.0) |
| HILL coefficient                      | 4.1 (24)      | ---          | 4.7 (39.1) | ---           |
| Elimination constant (L/hr')          | 18.2 (9.5)    | 0.093 (3.7)  | 168 (10.5) | ---           |
| Residual variability, additive error  | 17.9 (7.6)    | ---          | 17.7 (7.8) | ---           |

*Parameters corrected for 70 kg body-weight. %RSE=percent relative standard error.

| Half-life (hours) | Dose (50 mg/kg) | Absorption | Elimination | Distribution volume (L/kg) | Total body clearance (ml/min.1.73 m²) | AUC0-∞ (µg.h/ml) |
|-------------------|-----------------|------------|------------|---------------------------|---------------------------------------|-----------------|
| First dose        | 0.39±0.03 | 1.8±0.4   | 0.60±0.08  | 141±60                    | 233±79                                |
| 6th dose          | 0.16±0.03 | 1.7±0.3   | 0.60±0.15  | 131±60                    | 215±85                                |
| *Value            | 0.5000     | 0.4124     | 0.5000     | 0.4124                    | 0.4124                               |
| Mann Whitney test |                 |            |            |                           |                                       |

*Table 4: Pharmacokinetics of cefuroxime was assessed following 10, 15, or 20 mg/kg single oral dose of cefuroxime axetil. Pharmacokinetic parameters were obtained in 28 infants and children aged 3 months to 12 years (mean, 1.9 years). Figures are the mean±SD, by Powell et al. [40]

**Table 2: Cefuroxime final population pharmacokinetic parameters obtained in 36 infants undergoing cardiopulmonary bypass, whereas 6 infants did not. Parameters are estimated for the entire population of 42 infants who were aged < 1 year; figures are the typical value (%RSE) median, and %relative standard error (%RSE), by Gertler et al. [39]

**Table 3: Cefuroxime pharmacokinetic parameters obtained in plasma of 5 infants and children following cefuroxime single and multiple doses of 50 mg/kg. Values are reported according to the first and 6th dose. Figures are the mean±SD, by Del Rio et al. [37]
presence of organic solvents was assessed as a phenotypically measure of efflux. Cefuroxime susceptibility was also determined in presence of a known efflux pump inhibitor MC-207,110. Escherichia coli strains, that were organic solvent tolerant, had higher MIC values (4 to > 256 µg/ml; median, 24) than non-tolerant strains. In strains with most pronounced organic solvent-tolerance (i.e. cyclohexane-tolerant strains) cefuroxime MICs were decreased by 2-fold (P-value<0.03) supporting the hypothesis that efflux pump contributes to cefuroxime resistance in Escherichia coli.

Källman et al. [43] studied whether efflux or down-regulation of porins contributes to cefuroxime resistance in Klebsiella pneumoniae. Transcription-rate of acrA, ompK35, ramA, and soxS was determined by quantitative RT-PCR. All clinical isolates had similar non-susceptibility to cefuroxime. Phenylalanine arginine β-naphthylamide did not increase susceptibility to cefuroxime. Increased acrA transcription-rate and decreased ompK35 transcription-rate was seen in all strains. Multidrug-resistant phenotype of Klebsiella pneumoniae is associated with increased acrA and ramA gene transcription-rate and decreased ompK35 gene transcription-rate. Since cefuroxime resistance was not reversed by β-naphthylamide, this is attributable to decreased levels of OmpK35 rather than to efflux.

Alam et al. [44] reported the occurrence of antibiotic-resistance and production of β-lactamases including expended-spectrum β-lactamas in enteric bacteria isolated from hospital wastewater. Among 69 isolates tested for antibiotic sensitivity, 73.9% strains had cefuroxime-resistant rate of 39.1%. MIC showed a high resistance-level (800 to 1,600 µg/ml) tested for antibiotic sensitivity, 73.9% strains had cefuroxime-resistant rate of 39.1%. MIC showed a high resistance-level (800 to 1,600 µg/ml) of β-lactamases including expended-spectrum β-lactamas producing. Four extended-spectrum β-lactamase producing organisms (5.8%) could transfer their R-plasmid to the recipient strain of Escherichia coli K-12 with conjugation frequency ranging from 7.0*10^-4 to 8.8*10^-4. Enteric bacteria extended-spectrum β-lactamide producing is common in the wastewater, and such isolates may disseminate multidrug-resistance caused by genetic exchange mechanisms and requires immediate attention.

Gunduz and Altun [45] evaluated causative pathogens and their antibiotic-resistance in 850 positive strains isolated from the urine of children aged 3.0±3.7 years. Commonest causative pathogens were: Escherichia coli (64.2%) followed by Klebsiella pneumoniae (14.9%). Overall resistance-rate to three antibiotics ranged from 62.6 to 29.8% and that to cefuroxime was 28.7%. Escherichia coli was the commonest bacteria isolated which had the highest resistance-rate to cefuroxime. Several authors observed that antibiotic consumption including cefuroxime was associated with measurable and statistical significant increase in the resistance-rate [46-48].

### Discussion

Cefuroxime is a second-generation cephalosporin β-lactamase resistant which is active against gram-positive and most gram-negative organisms, and bacteria highly sensitive to cefuroxime are: Haemophilus influence (even strains ampicillin-resistant), Neisseria meningitides, and Streptococcus pneumoniae. Cefuroxime is available for oral, intravenous, and intramuscular administration; it is rapidly absorbed when given by mouth, the gastrointestinal absorption-rate of the prodrug cefuroxime axetil is 70%, and after hydrolysis it releases cefuroxime [1,3]. Cefuroxime and cefuroxime axetil are efficacious and safe in infants and children [3-10] and adverse-effects do not occur even when high doses are given [15]. Parenteral dosing-regimens consist of 25 mg/kg, twice-daily, thrice-daily or 4-times-daily and dose intervals decrease with infant maturation [1]. Oral dosing-regimens are: 10, 15 mg/kg and 250 mg for children aged up to 1 year, 2 to 11 and 12 to 17 years, respectively-tract [3]. Prophylaxis reduces the frequency of infection episodes of urinary-tract [3,23] and respiratory infections [20], surgery [16-19], pneumonia [21] acute otitis media [22], and sinusitis [24], and it has been used to decrease bacteria burden before surgery [6-19]. Cefuroxime axetil is the drug of choice for treatment of erythema migrans [14] and Lyme disease [29]. Treatment with cefuroxime has been used to treat generalized infections [20], urinary-tract infections [22, 23], and also pneumonia [21], and sinusitis [24], caused by gram-negative and gram-positive organisms. Optimization of dosing-regimens consists in maintaining cefuroxime plasma concentrations above MICs of infective pathogens for at least 50% to 60% of dosing intervals [26-28]. Cefuroxime crosses the brain-blood-barrier easily and concentration in cerebrospinal fluid is about 10% of that in plasma [32-35], and successfully treated meningitis caused by β-haemolytic streptococci, Streptococcus pneumonia, and Neisseria meningitides [36,37], whereas meningitis caused by Salmonella species requires an additional dose of 5 mg injected intravenicularly. Cefuroxime pharmacokinetics have been assessed in infants [39], in infants and children [37,41] and those, following cefuroxime axetil administration have been reported in infants and children [40]. Cefuroxime half-life is 1.8 hours in children aged up to 6 years [37] and 6 hours in infants during the first week of life [38]. Following three doses of cefuroxime axetil, cefuroxime Cmax and AUC∞ were not statistically different indicating that cefuroxime does not accumulate in blood [40]. Consistent results were obtained following cefuroxime administration on the first and 6th dose, thus cefuroxime does not accumulate in blood even for prolonged treatment [37]. Distribution volume of cefuroxime is 0.60 L/kg in infants and children; thus it migrates into body tissues in relevant amounts and diffuses in more than one half of the body volume [37]. Escherichia coli [42,44,45] and Klebsiella pneumoniae [43,45] may become cefuroxime-resistant; efflux pump [42] is responsible for increased-resistance of Escherichia coli [42] whereas resistance of Klebsiella pneumonia is caused by increased
transcription-rates of acrA, ramA genes and decreased transcription-rate of the gene OmpK35 [43]. Multidrug-resistant of Enteric extended-spectrum β-lactamase producing organism to cefuroxime has been observed in about 20% of isolates. Resistance is caused by genetic exchange mechanisms, consequently disseminate multidrug-resistance, and requires immediate attention [44]. Cefuroxime consumption is associated with increased bacteria resistance [46–48], continuous surveillance of resistance-rates and antibiotic consumption are of utmost importance for optimizing antibiotic dosing strategy.

In conclusion, cefuroxime is a second-generation cephalosporin β-lactamase resistant; it inhibits bacterial cell wall synthesis of gram-positive and gram-negative organisms, and is rapidly bactericidal. Administration forms are available for oral and parenteral administrations, cefuroxime is rapidly absorbed when given orally, and the absorption-rate of the prodrug cefuroxime axetil is about 70%. Cefuroxime axetil works by releasing cefuroxime after hydrolysis. Oral administration of cefuroxime and cefuroxime axetil yields high blood concentrations ensuring eradication of pathogenic microorganisms. These drugs, even given at high doses, do not cause adverse-effects. Cefuroxime axetil is the drug of choice to treat erythema migrans [14] and Lyme [29] diseases. Cefuroxime has been found efficacy against infections caused by gram-negative and gram-positive bacteria. Cefuroxime elimination half-live are 6 hours in infants during the first week of life and 1.8 hours in children aged up to 6 years. Cefuroxime penetrates into the cerebrospinal fluid where achieves a dose of about 10% of that in plasma concentration, and successfully treated meningitis caused by β-haemolytic streptococci species, Haemophilus influenzae, and Neisseria meningitides, whereas meningitis caused by Salmonella species requires an additional dose of 5 mg. Some bacteria, including Escherichia coli and Klebsiella pneumoniae, may become cefuroxime-resistant, and antibiotic consumption, including cefuroxime, is associated with increased resistance, and judicious use of this antibiotic ensures efficacy against bacteria.

Conflict of interests

The authors declare no conflicts of financial interest in any product or service mentioned in the manuscript, including grants, equipment, medications, employment, gifts, and honoraria.

This article is a review and drugs have not been administered to men or animals.

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