

otic; Ceflor; Cloracef; Especlor; Forifek; Medikoncef; Soclor; **Irl.**: Cefager; Distaclor; Kefid; Pinaclor; **Israel**: Ceclor; Cefalor; **Ital.**: Altaclor; Bacticef; Bactigram; Cefulton; Citidlor; Clorad; Clorazer; Dorf; Emredor; Eurocef; Fuclode; Genidor; Klicaf; Lafarlor; Macovan; Necloral; Omaspir; Oracef; Panacef; Performer; Selanor; Selvidor; Takecef; Tibifor; Valacior; **Malaysia**: Distaclor; Sifalor; Sofidor; Vercef; **Mex.**: Arcefal; Cec; Ceclor; Cefalan; Ceflacid; Fascilor; Fermed; Ranclor; Serviclora; Teraclor; **Neth.**: Ceclor; **NZ**: Clorotir; **Philipp.**: Aczebi; Brelox; Ceclobid; Ceclor; Clorotir; Ephron; Lorcef; Vefarol; Vercef; Verzat; Xelent; Xeztron; **Pol.**: Ceclor; Cek; Kloracef; Pandor; Serviclora; Vercef; **Port.**: **Rus.**: Ceclor (Llewoop); Vercef (Bepuef); **S.Afr.**: Cec; Ceclor; CloraCef; Vercef; **Singapore**: Cleancef; Distaclor; Sofidor; Vercef; **Spain**: Ceclor; **Switz.**: Ceclor; **Thai**: Celco; Clorotir; Distaclor; Kefador; Sifalor; Tefalor; Vercef; **Turk.**: Ceclor; Kefsid; Losefar; **UAE**: Recocef; **UK**: Bacticlora; Distaclor; Kefid; **USA**: Ceclor; Ranclor; **Venez.**: Ceclor.

Multi-ingredient. Mex.: Ceclorlox.

Cefadroxil (BAN, USAN, pINN)

BL-S578; Cefadroksilis monohidratas; Cefadroksyl jednowodny; Cefadroxil; Cefadroxil monohydrát; Cefadroxil monohydraté; Cefadroxilmonohydrát; Cefadroxilo; Cefadroxilum; Cefadroxilum monohydricum; Cephadroxil; Kefadroksili; Kefadroksilimonohydratti; MJF-11567-3; Sefadroksil. (7R)-7-(α -D-4-Hydroxyphenylglycylamino)-3-methyl-3-cephem-4-carboxylic acid monohydrate.

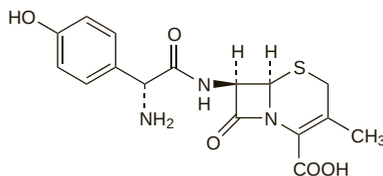
Цефадроксил

$C_{16}H_{17}N_3O_5S \cdot H_2O = 381.4$.

CAS — 50370-12-2 (anhydrous cefadroxil); 119922-85-9 (cefadroxil hemihydrate); 66592-87-8 (cefadroxil monohydrate).

ATC — J01DB05.

ATC Vet — QJ01DB05.



Pharmacopoeias. In *Chin.*, *Eur.* (see p.vii), and *US. Jpn* includes the anhydrous substance.

Ph. Eur. 6.2 (Cefadroxil Monohydrate). A white or almost white powder. Slightly soluble in water; very slightly soluble in alcohol. A 5% suspension in water has a pH of 4.0 to 6.0. Protect from light.

USP 31 (Cefadroxil). A white to off-white crystalline powder. Slightly soluble in water; practically insoluble in alcohol, in chloroform, and in ether. pH of a 5% suspension in water is between 4.0 and 6.0. Store in airtight containers.

Adverse Effects and Precautions

As for Cefalexin, p.218.

Breast feeding. Although higher concentrations of cefadroxil were reported in breast milk compared with cefalexin, cefalotin, cefapirin, and cefotaxime,¹ no detectable cefadroxil would be expected in breast-fed infants and no adverse effects have been seen in infants whose mothers were receiving cefadroxil. Accordingly, the American Academy of Pediatrics considers² that cefadroxil is usually compatible with breast feeding.

- Kafetzis DA, et al. Passage of cephalosporins and amoxicillin into the breast milk. *Acta Paediatr Scand* 1981; **70**: 285-8.
- American Academy of Pediatrics. The transfer of drugs and other chemicals into human milk. *Pediatrics* 2001; **108**: 776-89. Correction. *ibid.*: 1029. Also available at: <http://aapolicy.aappublications.org/cgi/content/full/pediatrics%3b108/3/776> (accessed 25/05/04)

Interactions

As for Cefalexin, p.218.

Antimicrobial Action

As for Cefalexin, p.218.

Pharmacokinetics

Cefadroxil is almost completely absorbed from the gastrointestinal tract. After oral doses of 500 mg and 1 g, peak plasma concentrations of about 16 and 30 micrograms/mL respectively are obtained after 1.5 to 2 hours. Although peak concentrations are similar to those of cefalexin, plasma concentrations are more sustained. Dosage with food does not appear to affect the absorption of cefadroxil. About 20% of cefadroxil is reported to be bound to plasma proteins. The plasma half-life of cefadroxil is about 1.5 hours and is prolonged in patients with renal impairment.

Cefadroxil is widely distributed to body tissues and fluids. It crosses the placenta and appears in breast milk.

More than 90% of a dose of cefadroxil may be excreted unchanged in the urine within 24 hours by glomerular filtration and tubular secretion; peak urinary concentrations of 1.8 mg/mL have been reported after a dose of 500 mg. Cefadroxil is removed by haemodialysis.

References

- Tanrisever B, Santella PJ. Cefadroxil: a review of its antibacterial, pharmacokinetic and therapeutic properties in comparison with cephalexin and cephradine. *Drugs* 1986; **32** (suppl 3): 1-16.
- Wise R. The pharmacokinetics of the oral cephalosporins—a review. *J Antimicrob Chemother* 1990; **26** (suppl E): 13-20.
- Garrigues TM, et al. Dose-dependent absorption and elimination of cefadroxil in man. *Eur J Clin Pharmacol* 1991; **41**: 179-83.

Uses and Administration

Cefadroxil is a first-generation cephalosporin antibacterial that is the para-hydroxy derivative of cefalexin (p.219), and is used similarly in the treatment of mild to moderate susceptible infections. It is given orally, and doses are expressed in terms of the anhydrous substance; 1.04 g of cefadroxil monohydrate is equivalent to about 1 g of anhydrous cefadroxil. The usual adult dose is 1 to 2 g daily as a single dose or in two divided doses. The following doses are used in children weighing less than 40 kg: 500 mg twice daily for those over 6 years of age, 250 mg twice daily for those aged 1 to 6 years, and 25 mg/kg daily in divided doses for infants under 1 year. For details of reduced doses of cefadroxil in patients with renal impairment, see below.

Cefadroxil has also been used as the lysine derivative.

Administration in renal impairment. Following an initial loading dose of 0.5 to 1 g, dosage of cefadroxil should be adjusted in patients with renal impairment according to creatinine clearance (CC):

- CC 26 to 50 mL/minute per 1.73 m²: 0.5 to 1 g every 12 hours
- CC 11 to 25 mL/minute per 1.73 m²: 0.5 to 1 g every 24 hours
- CC 10 mL/minute per 1.73 m² or less: 0.5 to 1 g every 36 hours.

Preparations

BP 2008: Cefadroxil Capsules; Cefadroxil Oral Suspension; **USP 31:** Cefadroxil Capsules; Cefadroxil for Oral Suspension; Cefadroxil Tablets.

Proprietary Preparations (details are given in Part 3)

Arg.: Cefabiot; Cefacar; Cefacilina; Cefadrox; Cefamar; Cefasin; Cefatenk; Droxil; Kandinc; Klonadroxil; Versatic; **Austria:** Biodroxil; Duracef; **Belg.:** Duracef; Moxacef; **Braz.:** Cefadroxon; Cefamox; Celoxin; Droxifal; Neo Cefadri; **Canad.:** Duricef; **Chile:** Adroxef; Biodroxil; Cefamox; Sedafex; **Cz.:** Biodroxil; Cedrox; Cefadrox; Duracef; **Fin.:** Duracef; **Fr.:** Oracefal; **Ger.:** Cedrox; Gruncef; **Gr.:** Cefalom; Kleotrat; Moxacef; Nefalox; **Hong Kong:** Amben; Androxyl; Biodroxil; Duracef; Qualidrox; Sofidrox; **Hung.:** Biodroxil; Duracef; **India:** Cefadrox; Cefadur; Lactocef; Lydroxil; Odoxil; Pendrox; Vepan; Vistadrox; **Indon.:** Abxil; Ancefa; Bidicef; Biodroxil; Cefat; Dexacef; Doxide; Duricef; Erphadrox; Ethicef; Kelfex; Lapicef; Librocef; Longcef; Opicef; Osadrox; Pyricef; Q Cef; Qidrox; Renasistin; Roksicaf; Sedrofen; Stafonin; Tisacef; Vldrox; **Irl.:** Ultracef; **Israel:** Biodroxil; Duracef; **Ital.:** Cefadri; Ceoxil; Cephos; Foxil; Oradroxil; **Malaysia:** Kefloxin; Sofidrox; **Mex.:** Cefamox; Cepotec; Duracef; Teroxina; **Philipp.:** Droxex; Drozi; Lexitad; **Pol.:** Biodroxil; Duracef; **Port.:** Biofalil; Cefacile; Ceforal; Cefarid; **S.Afr.:** Cipadur; Dacef; Duracef; **Singapore:** Duricef; Sofidrox; **Spain:** Duracef; **Swed.:** Cefamox; **Thai.:** Cefadri; **Turk.:** Cefadur; Duricef; **UK:** Baxan; **USA:** Duricef; **Venez.:** Bidroxyl; Cedroxim; Cefadri; Cefaval; Cefonax; Drocef; Droxifan; Grunicef; Sanodil.

Multi-ingredient. Arg.: Cefacar Mucolítico; Cefacilina Bronquial; **Mex.:** Duracef Expec.

Cefalexin (BAN, pINN)

66873; Cefaleksinas monohidratas; Cefaleksyna; Cefalexin monohydrát; Cefalexina; Cefalexine; Cefalexine monohydraté; Cefalexinmonohydrát; Cefalexinum; Cefalexinum monohydricum; Cephalexin (USAN); Kefaleksini; Kefaleksiniimonohydratti; Sefaleksin. (7R)-3-Methyl-7-(α -D-phenylglycylamino)-3-cephem-4-carboxylic acid monohydrate.

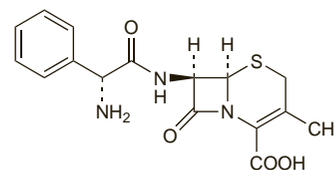
Цефалексин

$C_{16}H_{17}N_3O_4S \cdot H_2O = 365.4$.

CAS — 15686-71-2 (anhydrous cefalexin); 23325-78-2 (cefalexin monohydrate).

ATC — J01DB01.

ATC Vet — QJ01DB01; QJ51DA01.



Pharmacopoeias. In *Chin.*, *Eur.* (see p.vii), *Jpn*, *US*, and *Viet*.

Ph. Eur. 6.2 (Cefalexin Monohydrate). A white or almost white crystalline powder. Sparingly soluble in water; practically insoluble in alcohol. A 0.5% solution in water has a pH of 4.0 to 5.5. Protect from light.

USP 31 (Cefalexin). A white to off-white crystalline powder. Slightly soluble in water; practically insoluble in alcohol, in chloroform, and in ether. pH of a 5% suspension in water is between 3.0 and 5.5. Store in airtight containers.

Cefalexin Hydrochloride (BANM, pINN)

Céfaléxine, Chlorhydrate de; Cefalexini Hydrochloridum; Cephalexin Hydrochloride (USAN); Hidrocloruro de cefalexina; LY-061188.

Цефалексина Гидрохлорид

$C_{16}H_{17}N_3O_4S \cdot HCl \cdot H_2O = 401.9$.

CAS — 105879-42-3.

ATC — J01DB01.

ATC Vet — QJ01DB01.

Pharmacopoeias. In *US*.

USP 31 (Cefalexin Hydrochloride). A white to off-white crystalline powder. Soluble 1 in 100 in water, in acetone, in acetonitrile, in alcohol, in dimethylformamide, and in methyl alcohol; practically insoluble in chloroform, in ether, in ethyl acetate, and in isopropyl alcohol. pH of a 1% solution in water is between 1.5 and 3.0. Store in airtight containers.

Adverse Effects and Precautions

As for Cefalotin Sodium, p.219.

The most common adverse effects of cefalexin and other oral cephalosporins are generally gastrointestinal disturbances and hypersensitivity reactions. Pseudomembranous colitis has been reported.

References

- Dave J, et al. Cefalexin induced toxic epidermal necrolysis. *J Antimicrob Chemother* 1991; **28**: 477-8.
- Baran R, Perrin C. Fixed-drug eruption presenting as an acute paronychia. *Br J Dermatol* 1991; **125**: 592-5.
- Clark RF. Crystalluria following cephalexin overdose. *Pediatrics* 1992; **89**: 672-4.
- Murray KM, Camp MS. Cephalexin-induced Stevens-Johnson syndrome. *Ann Pharmacother* 1992; **26**: 1230-3.
- Czechowicz RT, et al. Bullous pemphigoid induced by cephalexin. *Australas J Dermatol* 2001; **42**: 132-5.
- Longstreth KL, et al. Cephalexin-induced acute tubular necrosis. *Pharmacotherapy* 2004; **24**: 808-11.
- Skoog SM, et al. Cephalexin-induced cholestatic hepatitis. *J Clin Gastroenterol* 2004; **38**: 833.
- Penttilä J, et al. Delirium in an adolescent patient during treatment with cephalexin. *J Adolesc Health* 2006; **39**: 782-3.

Porphyria. Cefalexin is considered to be unsafe in patients with porphyria although there is conflicting experimental evidence of porphyrinogenicity.

Interactions

The renal excretion of cefalexin, and many other cephalosporins, is delayed by probenecid.

Hormonal contraceptives. There have been isolated reports of cefalexin decreasing the efficacy of oestrogen-containing oral contraceptives.¹ For a discussion of decreased efficacy of oral contraceptives and the need for additional contraceptive methods in patients taking broad-spectrum antibacterials, see under Hormonal Contraceptives, p.2068.

- Friedman M, et al. Cefalexin and Microgynon-30 do not go well together. *J Obstet Gynaecol* 1982; **2**: 195-6.

Antimicrobial Action

As for Cefalotin Sodium, p.220, although cefalexin is generally less potent. Some strains of Gram-negative bacteria may be inhibited only by the high concentrations achievable in the urinary tract. *Haemophilus influenzae* is moderately resistant to cefalexin.

Pharmacokinetics

Cefalexin is almost completely absorbed from the gastrointestinal tract and produces a peak plasma concentration of about 18 micrograms/mL 1 hour after a 500-mg oral dose. If cefalexin is taken with food, absorption may be delayed, but the total amount absorbed is