

- ns of the bones, joints, soft tissue and skin, and vanis in patients with impaired immune response

demonstrated with many Gram-negative bacteria under experimental conditions, tways predictable, combination should be considered in severe, life-threatening as aeruoinosa. Because of physical incompatibility, the two drups must be

Dosage and administration
Dosage
Dosage and administration
Dosage
and administration
Dosage
and children over 12 years old:
The usual dosage is 1-2 of it ebacted, administered once daily (every 24 hours).
In severe cases or in infection caused by only moderately sensitive organisms, the dosa
Menantae, Infants and children up to 1-2 years old:
Dosage of the dosage

Contrainctations of follocation hydrochloride must be excluded before intramuscular injection of cathrisone when lidocatine hydrochloride is used as a solvent.

Warnings and precautions

Even after forward instructions are produced in the production of the product

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The following undesirable effects, which subsided either spontaneously or after withdrawal of the drug, have been observed during the use of certifications. Or certifications of the grant fact that subsided either spontaneously or after withdrawal of the drug, have been observed during the use of the drugs and the control of the grant fact that subsides a specific common control of the spontaneously of the control of the spontaneously of the spon

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Note: Many strains of 5-laterames-producing Bacteroides app. (notably 8. Anglis) are resissant.

Pharmacokinetics
The pharmacokinetics of confiscence are nonlinear.

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Ceftriaxone penetrates the inflamed meninges of neonates, infants and children.
The average concentration in CSF during bacterial meningitis is 17% of the plasma concentration; in aseptic meningitis it is 4%. 24 hours

The average concentration in CSP cours goacteria minimizes 8 17-6 of the pleastre concentration; in septic meninguis its 4%. 24 hours fater IX. injection of Lebacer in doses of 50-100 mg/kg bodyweight, cethriaxone concentrations > 1.4 mg/l were measured in GSF. In adult patients with meningitis, administration of 50 mg/kg leads within 2-24 hours to CSF concentrations several times higher than the minimum inhibitory concentrations required for the most common causative organisms of meningitis.

Metabolism: Ceftriaxone is not metabolized in the organism itself. Only following billiary excretion into the intestinal lumen does the intestinal flora transform the active incredient into inactive metabolites.

flora transform the active ingredient into inactive metabolites. Ellimination: Plasma clearance is 10-22 ml/min. Renal clearance is 5-12 ml/min. 50-60% of ceftriaxone is excreted unchanged via the kidneys, while 40-50% is excreted unchanged in the bile. The plasma half-life in adults is about 8 hours.

Pharmacokinetics in special patient groups: In neonates, renal elimination accounts for about 70% of the dose.

In infants aged less than 8 days and in persons aged over 75 years, the average plasma half-life is approximately 2-3 times that in healthy young adults.

In patients with mild to moderate renal failure or hepatic dysfunction, the pharmacokinetics of certifications ere only slightly altered. The losten half-life is minimally increased. If kinder yunchot agine is impaired, billiany elimination of certificance is increased, whereast if liver

function alone is impaired, renal elimination is increased. Presentation

Lebacef 0.5 g for I.M. injection:

DPAT Kit: 1 vial containing sterile powder of ceftriaxone sodium equivalent to 0.5 g of ceftriaxone, 1 solvent ampoule containing 2 ml lidocalne solution (lidocalne hydrochloride 1%), sterile syringe, 2 needles, sterile alcohol swab, adhesive bandage.

Lebase 0.5 g for I.V. injection:
OPAT Rit: 1 vial containing sterile powder of ceftriaxone sodium equivalent to 0.5 g of ceftriaxone, 1 solvent ampoule containing 5 ml water for injection, sterile syringe, 2 needles, sterile alcohol swab, adhesive bandage.

Lebacef 0.5 g for I.M./I.V. injection:

Pack of 10 vials each containing sterile powder of ceftriaxone sodium equivalent to 0.5 g of ceftriaxone. Lebacef 1 g for I.M. injection:

OPAT Kit: 1 vial containing sterile powder of ceftriaxone sodium equivalent to 1 g of ceftriaxone, 1 solvent ampoule containing 3.5 ml lidocaine solution (tidocaine hydrochloride 1%), sterile syringe, 2 needies, sterile alcohol swab, adhesive bandage. Lebacef 1 of CVI, kinection:

OPAT Kit: 1 vial containing sterile powder of ceftriaxone sodium equivalent to 1 g of ceftriaxone, 1 solvent ampoule containing 10 ml water for injection, sterile syringe, 2 needles, sterile alcohol swab, adhesive bandage.

Lebacef 1 g for I.M./İ.V. İnjection:
Pack of 10 vials each containing sterile powder of ceftriaxone sodium equivalent to 1 g of ceftriaxone.

Lebacef 2 g for I.V. infusion:
Pack of 1 or 10 vials each containing sterile powder of ceftriaxone sodium equivalent to 2 g of ceftriaxone.

Expiry date and storage conditions
See the expiry date printed on the outer carton.

See the expiry date printed on the outer carton.

This date refers to the product correctly stored in unopened package.

Beware not to use Lebacef after this date. Store below 30°C. Protect from light and heat.

Keep all medicines out of reach of children.

Manufactured by: Mitim S.R.L.

For:

Brescia, Italy

ARWAN Pharmaceutical Industries Lebanon s.a.l.

THIS IS A MEDICAMENT

- Medicament is a product which affects your health, and its consumption contrary to instructions is dangerous for you.
- Follow strictly the doctor's prescription, the method of use and the instructions of the pharmacist who sold the medicament.
 The doctor and the pharmacist are experts in medicines, their henefits and risks.
- . Do not by yourself interrupt the period of treatment prescribed for you.
- . Do not repeat the same prescription without consulting your doctor.

Keep all medicaments out of the reach of children. Council of Arab Health Ministers. Union of Arab Pharmacists