the infection and the patient's		e.
Patients with renal and hepatic Recommended starting and ma		r nationts with impaired
renal function:	annicinance doses to	patients with impanea
Creatinine Clearance [mL/min/ 1.73 m²]	Serum Creatinine [µmol/L]	Intravenous Dose [mg]
> 60	< 124	See Usual Dosage.
30 - 60	124 to 168	200 - 400 mg every 12 h
< 30	> 169	200 - 400 mg every 24 h
Patients on haemodialysis	> 169	200 - 400 mg every 24 h (after dialysis)
Patients on peritoneal dialysis	> 169	200 - 400 mg every 24 h
In patients with impaired liver Dosing in children with impairs studied. Method of administration Ciprobay 200 mg should be ch used if cloudy. Ciprofloxacin should be admin the infusion duration is 60 mir In adult patients, infusion time and 30 minutes for 200 mg Cip will minimise patient discomfor infusion solution can be infuse compatible infusion solutions Hypersensitivity to the act of the excipients listed in second the excipients listed in second the second than the	ecked visually prior istered by intravenoutes. e is 60 minutes for 4 probay 200 mg. Slow ort and reduce the rised either directly or a (see section 6.6). Eive substance, to ot section of ciprofloxacin arecautions for use	atic function has not been to use. It must not be us infusion. For children, 00 mg Ciprobay 200 mg vinfusion into a large veir sk of venous irritation. The ofter mixing with other ther quinolones or to any and tizanidine (see section
pathogens Ciprofloxacin monotherapy is and infections that might be defined in such infections ciprofloxacin appropriate antibacterial agenstreptococcal Infections (included infections due to inadequate effections due to inadequate effections due to inadequate effections due to inadequate effections. Epididymo-orchitis and pelvic influoroquinolone-resistant Neisfor epididymo-orchitis and pelciprofloxacin should only be compropriate antibacterial agentications.	not suited for treatmue to Gram-positive nust be co-adminits. ding Streptococcus puded for the treatment ficacy. inflammatory disease seria gonorrhoeae iso vic inflammatory disease vic inflammatory disease vic inflammatory disease to be excluded. If	nent of severe infections or anaerobic pathogens. stered with other ineumoniae) nt of streptococcal ses may be caused by plates. seases, empirical ation with another in) unless ciprofloxacinclinical improvement is

Urinary tract infections

Intra-abdominal infections

<u>Travellers' diarrhoea</u>

Inhalational anthrax

Paediatric population

years of age.

Photosensitivity

should be discontinued.

Cardiac disorders

interval such as, for example: congenital long QT syndrome

hypomagnesaemia)

Ren<u>al and urinary system</u>

Impaired renal function

Hepatobiliary system

should be monitored.

necessary (see section 4.5).

Interaction with tests

Injection Site Reaction

content, see section 2).

recommended (see section 4.5).

patients currently taking ciprofloxacin.

Methotrexate

or worsen.

Tizanidine

Methotrexate

Theophylline

Duloxetine

administration.

Clozapine

<u>Sildenafil</u>

Breast-feeding

Other xanthine derivatives

accumulation of ciprofloxacin.

macrolides, antipsychotics)

physicians who are experienced in the treatment of cystic fibrosis and/or severe infections in children and adolescents. Ciprofloxacin has been shown to cause arthropathy in weight-bearing joints of immature animals. Safety data from a randomised double-blind study on ciprofloxacin use in children (ciprofloxacin: n=335, mean age = 6.3 years; comparators: n=349, mean age = 6.2 years; age range = 1 to 17 years) revealed an incidence of suspected drug-related arthropathy (discerned from joint-related clinical signs and symptoms) by Day +42 of 7.2% and 4.6%. Respectively, an incidence of drug-related arthropathy by 1-year follow-up was 9.0% and 5.7%. The increase of suspected drug-related arthropathy cases over time was not statistically significant between groups. Treatment

should be initiated only after a careful benefit/risk evaluation, due to

Clinical trials have included children and adolescents aged 5 - 17 years.

More limited experience is available in treating children between 1 and 5

Ciprofloxacin treatment of urinary tract infections should be considered

Clinical trials have included children and adolescents aged 1 - 17 years.

Other severe infections in accordance with official guidance, or after

careful benefit-risk evaluation when other treatments cannot be used,

or after failure to conventional therapy and when the microbiological

when other treatments cannot be used, and should be based on the results

possible adverse events related to joints and/or surrounding tissue.

Broncho-pulmonary infections in cystic fibrosis

of the microbiological documentation.

documentation can justify a ciprofloxacin use.

Other specific severe infections

Complicated urinary tract infections and pyelonephritis

Resistance to fluoroquinolones of *Escherichia coli* – the most common

resistance in Escherichia coli to fluoroquinolones.

post-surgical intra-abdominal infections.

Infections of the bones and joints

regarding the treatment of anthrax.

pathogen involved in urinary tract infections – varies across the European

Union. Prescribers are advised to take into account the local prevalence of

There are limited data on the efficacy of ciprofloxacin in the treatment of

resistance to ciprofloxacin in relevant pathogens in the countries visited.

Ciprofloxacin should be used in combination with other antimicrobial

Use in humans is based on in-vitro susceptibility data and on animal

should refer to national and /or international consensus documents

official guidance. Ciprofloxacin treatment should be initiated only by

agents depending on the results of the microbiological documentation.

experimental data together with limited human data. Treating physicians

The use of ciprofloxacin in children and adolescents should follow available

The choice of ciprofloxacin should take into account information on

The use of ciprofloxacin for specific severe infections other than those mentioned above has not been evaluated in clinical trials and the clinical experience is limited. Consequently, caution is advised when treating patients with these infections. **Hypersensitivity** Hypersensitivity and allergic reactions, including anaphylaxis and anaphylactoid reactions, may occur following a single dose (see section 4.8) and may be life-threatening. If such reaction occurs, ciprofloxacin should be discontinued and an adequate medical treatment is required. Musculoskeletal System

Ciprofloxacin should generally not be used in patients with a history of

tendon disease/disorder related to quinolone treatment. Nevertheless, in

very rare instances, after microbiological documentation of the causative

infections, particularly in the event of failure of the standard therapy or bacterial resistance, where the microbiological data may justify the use of

ciprofloxacin. Tendinitis and tendon rupture (especially Achilles tendon),

sometimes bilateral, may occur with ciprofloxacin, even within the first 48

hours of treatment. Inflammation and ruptures of tendon may occur even

concomitantly treated with corticosteroids (see section 4.8). At any sign of

should be discontinued. Care should be taken to keep the affected limb at

up to several months after discontinuation of ciprofloxacin therapy. The

risk of tendinopathy may be increased in elderly patients or in patients

tendinitis (e.g. painful swelling, inflammation), ciprofloxacin treatment

Ciprofloxacin should be used with caution in patients with myasthenia gravis, because symptoms can be exacerbated (see section 4.8).

organism and evaluation of the risk/benefit balance, ciprofloxacin may be prescribed to these patients for the treatment of certain severe

Ciprofloxacin has been shown to cause photosensitivity reactions. Patients taking ciprofloxacin should be advised to avoid direct exposure to either extensive sunlight or UV irradiation during treatment (see section 4.8). Central Nervous System Ciprofloxacin like other quinolones are known to trigger seizures or lower the seizure threshold. Cases of status epilepticus have been reported. Ciprofloxacin should be used with caution in patients with CNS disorders which may be predisposed to seizure. If seizures occur ciprofloxacin should be discontinued (see section 4.8). Psychiatric reactions may occur even after first administration of ciprofloxacin. In rare cases, depression or psychosis can progress to suicidal ideations/thoughts culminating in attempted

suicide or completed suicide. In the occurrence of such cases, ciprofloxacin

Cases of polyneuropathy (based on neurological symptoms such as pain,

burning, sensory disturbances or muscle weakness, alone or in combination)

be discontinued in patients experiencing symptoms of neuropathy, including

concomitant use of drugs that are known to prolong the QT interval

(e.g. Class IA and III anti-arrhythmics, tricyclic antidepressants,

have been reported in patients receiving ciprofloxacin. Ciprofloxacin should

pain, burning, tingling, numbness, and/or weakness in order to prevent the

development of an irreversible condition (see section 4.8).

Caution should be taken when using fluoroquinolones, including ciprofloxacin, in patients with known risk factors for prolongation of the QT

uncorrected electrolyte imbalance (e.g. hypokalaemia,

cardiac disease (e.g. heart failure, myocardial infarction, bradycardia) Elderly patients and women may be more sensitive to QTc-prolonging medications. Therefore, caution should be taken when using fluoroquinolones, including ciprofloxacin, in these populations. (See section 4.2 Elderly patients, section 4.5, section 4.8, section 4.9). Hypoglycemia As with other quinolones, hypoglycemia has been reported most often in diabetic patients, predominantly in the elderly population. In all diabetic patients, careful monitoring of blood glucose is recommended (see section <u>Gastrointestinal System</u> The occurrence of severe and persistent diarrhoea during or after treatment (including several weeks after treatment) may indicate an antibioticassociated colitis (life-threatening with possible fatal outcome), requiring immediate treatment (see section 4.8). In such cases, ciprofloxacin should immediately be discontinued, and an appropriate therapy initiated. Anti-

Crystalluria related to the use of ciprofloxacin has been reported (see section

4.8). Patients receiving ciprofloxacin should be well hydrated and excessive

Since ciprofloxacin is largely excreted unchanged via renal pathway dose

Cases of hepatic necrosis and life-threatening hepatic failure have been

and symptoms of hepatic disease (such as anorexia, jaundice, dark urine,

Haemolytic reactions have been reported with ciprofloxacin in patients with glucose-6-phosphate dehydrogenase deficiency. Ciprofloxacin should

be avoided in these patients unless the potential benefit is considered to

outweigh the possible risk. In this case, potential occurrence of haemolysis

During or following a course of treatment with ciprofloxacin bacteria that

reported with ciprofloxacin (see section 4.8). In the event of any signs

pruritus, or tender abdomen), treatment should be discontinued.

Glucose-6-phosphate dehydrogenase deficiency

in section 4.2 to avoid an increase in adverse drug reactions due to

adjustment is needed in patients with impaired renal function as described

peristaltic drugs are contraindicated in this situation.

alkalinity of the urine should be avoided.

demonstrate resistance to ciprofloxacin may be isolated, with or without a clinically apparent superinfection. There may be a particular risk of selecting for ciprofloxacin-resistant bacteria during extended durations of treatment and when treating nosocomial infections and/or infections caused by Staphylococcus and Pseudomonas species. Cytochrome P450 Ciprofloxacin inhibits CYP1A2 and thus may cause increased serum concentration of concomitantly administered substances metabolised by this enzyme (e.g. theophylline, clozapine, olanzapine, ropinirole, tizanidine, duloxetine). Co-administration of ciprofloxacin and tizanidine is contraindicated. Therefore, patients taking these substances concomitantly with ciprofloxacin should be monitored closely for clinical signs of overdose,

and determination of serum concentrations (e.g. of theophylline) may be

The in-vitro activity of ciprofloxacin against Mycobacterium tuberculosis

might give false negative bacteriological test results in specimens from

Local intravenous site reactions have been reported with the intravenous

administration of ciprofloxacin. These reactions are more frequent if

the infusion time is 30 minutes or less. These may appear as local skin

In patients for whom sodium intake is of medical concern (patients with

additional sodium load should be taken into account (for sodium chloride

Interaction with other medicinal products and other forms of

congestive heart failure, renal failure, nephrotic syndrome, etc.), the

reactions which resolve rapidly upon completion of the infusion. Subsequent

intravenous administration is not contraindicated unless the reactions recur

The concomitant use of ciprofloxacin with methotrexate is not

interaction Effects of other products on ciprofloxacin: Drugs known to prolong QT interval Ciprofloxacin, like other fluoroquinolones, should be used with caution in patients receiving drugs known to prolong QT interval (e.g. Class IA and III anti-arrhythmics, tricyclic antidepressants, macrolides, antipsychotics) (see section 4.4). Probenecid Probenecid interferes with renal secretion of ciprofloxacin. Co-administration of probenecid and ciprofloxacin increases ciprofloxacin serum concentrations. Effects of ciprofloxacin on other medicinal products:

Tizanidine must not be administered together with ciprofloxacin (see section

serum tizanidine concentration (Cmax increase: 7-fold, range: 4 to 21-fold;

AUC increase: 10-fold, range: 6 to 24-fold) when given concomitantly with

ciprofloxacin. Increased serum tizanidine concentration is associated with a

Renal tubular transport of methotrexate may be inhibited by concomitant

levels of methotrexate and increased risk of methotrexate-associated toxic

Concurrent administration of ciprofloxacin and theophylline can cause an undesirable increase in serum theophylline concentration. This can lead to theophylline-induced side effects that may rarely be life threatening or fatal. During the combination, serum theophylline concentrations should be checked and the theophylline dose reduced as necessary (see section 4.4).

administration of ciprofloxacin, potentially leading to increased plasma

reactions. The concomitant use is not recommended (see section 4.4).

4.3). In a clinical study with healthy subjects, there was an increase in

potentiated hypotensive and sedative effect.

On concurrent administration of ciprofloxacin and caffeine or pentoxifylline (oxpentifylline), raised serum concentrations of these xanthine derivatives were reported. Phenvtoin Simultaneous administration of ciprofloxacin and phenytoin may result in increased or reduced serum levels of phenytoin such that monitoring of drug levels is recommended. Cyclosporin A transient rise in the concentration of serum creatinine was observed when ciprofloxacin and cyclosporin containing medicinal products were administered simultaneously. Therefore, it is frequently (twice a week) necessary to control the serum creatinine concentrations in these patients. Vitamin K antagonists Simultaneous administration of ciprofloxacin with a vitamin K antagonist

may augment its anti-coagulant effects. The risk may vary with the underlying infection, age and general status of the patient so that the contribution of ciprofloxacin to the increase in INR (international normalised ratio) is difficult to assess. The INR should be monitored frequently during and shortly after co-administration of ciprofloxacin with a vitamin K antagonist (e.g., warfarin, acenocoumarol, phenprocoumon, or fluindione).

effects can be expected upon concomitant administration (see section 4.4). Ropinirole It was shown in a clinical study that concomitant use of ropinirole with ciprofloxacin, a moderate inhibitor of the CYP450 1A2 isozyme, results in an increase of Cmax and AUC of ropinirole by 60% and 84%, respectively. Monitoring of ropinirole-related side effects and dose adjustment as appropriate is recommended during and shortly after co-administration with ciprofloxacin (see section 4.4). It was demonstrated in healthy subjects that concomitant use of lidocaine containing medicinal products with ciprofloxacin, a moderate inhibitor of CYP450 1A2 isozyme, reduces clearance of intravenous lidocaine by 22%. Although lidocaine treatment was well tolerated, a possible interaction

with ciprofloxacin associated with side effects may occur upon concomitant

N-desmethylclozapine were increased by 29% and 31%, respectively. Clinical

surveillance and appropriate adjustment of clozapine dosage during and

shortly after co-administration with ciprofloxacin are advised (see section

Following concomitant administration of 250 mg ciprofloxacin

with clozapine for 7 days, serum concentrations of clozapine and

In clinical studies, it was demonstrated that concomitant use of duloxetine

with strong inhibitors of the CYP450 1A2 isozyme such as fluvoxamine, may

result in an increase of AUC and Cmax of duloxetine. Although no clinical

data are available on a possible interaction with ciprofloxacin, similar

 $C_{\rm max}$ and AUC of sildenafil were increased approximately twofold in healthy subjects after an oral dose of 50 mg given concomitantly with 500 mg ciprofloxacin. Therefore, caution should be used prescribing ciprofloxacin concomitantly with sildenafil taking into consideration the risks and the benefits. 4.6 Pregnancy and lactation Pregnancy The data that are available on administration of ciprofloxacin to pregnant women indicates no malformative or feto/neonatal toxicity of ciprofloxacin. Animal studies do not indicate direct or indirect harmful effects with respect to reproductive toxicity. In juvenile and prenatal animals exposed to quinolones, effects on immature cartilage have been observed, thus,

it cannot be excluded that the drug could cause damage to articular

damage, ciprofloxacin should not be used during breast-feeding.

Due to its neurological effects, ciprofloxacin may affect reaction time.

4.7 Effects on ability to drive and use machines

cartilage in the human immature organism / foetus (see section 5.3). As

a precautionary measure, it is preferable to avoid the use of ciprofloxacin

Ciprofloxacin is excreted in breast milk. Due to the potential risk of articular

Thus, the ability to drive or to operate machinery may be impaired. 4.8 Undesirable effects The most commonly reported adverse drug reactions (ADRs) are nausea, iniection of ta en-

diarrhoea, von and infusion s ADRs derived f Ciprobay (oral frequency are	ite reactions. rom clinical : , intravenous listed below.	studies and p and sequent The frequence	oost-marketir ial therapy) s cy analysis ta	ng surveilland sorted by cat kes into acco	ce with egories of
from both ora System Organ Class	Common	nous adminis Uncom- mon ≥ 1/1,000 to < 1/100	Rare ≥ 1 / 10,000 to < 1 / 1,000	Very Rare < 1/ 10,000	Frequency not known (cannot be estimated from the available data)
Infections and Infesta- tions		Mycotic superin- fections	Antibiotic associated colitis (very rarely with possible fatal outcome) (see section 4.4)		
Blood and Lymphatic System Disorders		Eosino- philia	Leukope- nia Anaemia Neutrope- nia	Haemo- lytic anaemia Agranulo- cytosis	

Leukocyto- Pancyto-

sis Thrombo-

cytopenia

Thrombo-

cytaemia

penia (life-

threate-

marrow depression (lifethreate-

ning)

Bone

ning)

			4.4) Vertigo	hyperten- sion and pseudotu- mor cerebri	
Eye Disorders			Visual disturban- ces (e.g. diplopia)	Visual colour distortions	
Ear and Labyrinth Disorders			Tinnitus Hearing loss / Hearing impaired		
Cardiac Disorders			Tachycar- dia		Ventricular arrhythmia, torsades de pointes (reported predominantly in patients with risk factors for QT prolon- gation), ECG QT prolon- ged (see sections 4.4 and
Vascular Disorders Respira- tory, Thoracic and Media- stinal			Vasodi- latation Hypo- tension Syncope Dyspnoea (including asthmatic condition)	Vasculitis	4.9)
Disorders Gastroin- testinal Disorders	Nausea Diarrhoea	Vomiting Gastroin- testinal and abdominal pains		Pancrea- titis	
Hepato- biliary Disorders		Dyspepsia Flatulence Increase in transa- minases Increased bilirubin	Hepatic impair- ment Choles- tatic icterus Hepatitis	Liver necrosis (very rarely progres- sing to life-threa- tening hepatic failure) (see section 4.4)	
Skin and Subcu- taneous Tissue Disorders		Rash Pruritus Urticaria	Photosen- sitivity reactions (see section 4.4)	Petechiae Erythema multi- forme Erythema nodosum Stevens- Johnson syndrome (poten- tially life- threate- ning) Toxic epidermal necrolysis (poten- tially life-threa- tening)	Acute genera- lised exanthe- matous pustu- losis (AGEP)
Musculo- skeletal and Connective Tissue Disorders		Musculo- skeletal pain (e.g. extremity pain, back pain, chest pain) Arthralgia	Myalgia Arthritis Increased muscle tone and cramping	Muscular weakness Tendinitis Tendon rupture (predominantly Achilles tendon) (see section 4.4) Exacerbation of symptoms of myasthenia gravis (see section	
Renal and Urinary Disorders		Renal impair- ment	Renal failure Haema- turia Crystal- luria (see section 4.4) Tubulo- inters- titial nephritis	4.4)	
General Disorders and Adminis- tration Site Conditions	Injection and infusion site reactions (only intra- venous adminis- tration)	Asthenia Fever	Oedema Sweating (hyper- hidrosis)		
Inves- tigations	Undariza ki	Increase in blood alkaline phosphatase	Increased amylase	Ipnov est-	International normalised ratio increased (in patients treated with Vitamin kantagonists)
subgroups of poral) treatmen	atients receiv				
Common Uncommon	Vomiting, Tra Thrombocyto disorientatio Seizures, Verl	openia, Thror n, Hallucina tigo, Visual d	nbocytaemia tions, Par- ar listurbances,	, Confusion a nd dysaesthes Hearing loss	and sia, ,
	Tachycardia, impairment			l failure, Oed	

resistance to many or all active substances within the class. Impermeability and/or active substance efflux pump mechanisms of resistance may have a variable effect on susceptibility to fluoroquinolones, which depends on the physiochemical properties of the various active substances within the class and the affinity of transport systems for each active substance. All invitro mechanisms of resistance are commonly observed in clinical isolates. Resistance mechanisms that inactivate other antibiotics such as permeation barriers (common in Pseudomonas aeruginosa) and efflux mechanisms may affect susceptibility to ciprofloxacin. Plasmid-mediated resistance encoded by qnr-genes has been reported. Spectrum of antibacterial activity

Susceptible

S ≤ 0.5 mg/L

 $S \le 0.5 \text{ mg/L}$

 $S \le 1 \text{ mg/L}$

 $S \le 1 \text{ mg/L}$

 $S \le 0.5 \text{ mg/L}$

S ≤ 0.03 mg/L

S ≤ 0.03 mg/L

 $S \le 0.5 \text{ mg/L}$

1 Staphylococcus spp. – breakpoints for ciprofloxacin relate to high dose

Non-species-related breakpoints have been determined mainly on the

basis of PK/PD data and are independent of MIC distributions of specific species. They are for use only for species that have not been given a

species-specific breakpoint and not for those species where susceptibility

The prevalence of acquired resistance may vary geographically and with

haemodialysis or peritoneal dialysis.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

transcription, repair and recombination.

area under the curve (AUC) and the MIC.

Pharmacokinetic/pharmacodynamic relationship

susceptibility and the latter from resistant strains:

theoretically reduce the absorption of ciprofloxacin in overdoses. Only a small quantity of ciprofloxacin (<10%) is eliminated by

Pharmacotherapeutic group: Fluoroquinolones, ATC code: J01MA02

As a fluoroquinolone antibacterial agent, the bactericidal action of

Efficacy mainly depends on the relation between the maximum

ciprofloxacin results from the inhibition of both type II topoisomerase

(DNA-gyrase) and topoisomerase IV, required for bacterial DNA replication,

concentration in serum (Cmax) and the minimum inhibitory concentration

(MIC) of ciprofloxacin for a bacterial pathogen and the relation between the

In-vitro resistance to ciprofloxacin can be acquired through a stepwise

IV. The degree of cross-resistance between ciprofloxacin and other

process by target site mutations in both DNA gyrase and topoisomerase

fluoroquinolones that results is variable. Single mutations may not result

in clinical resistance, but multiple mutations generally result in clinical

4.9 Overdose

has been reported.

interval prolongation.

Mechanism of action

Mechanism of resistance

EUCAST Recommendations

Enterobacteriaceae

Pseudomonas spp.

Acinetobacter spp.

Staphylococcus spp.1

Moraxella catarrhalis

Neisseria gonorrhoeae

Neisseria meningitidis

Non-species-related

breakpoints*

Microorganisms

Haemophilus influenzae and

testing is not recommended.

Streptococcus species see section 4.4).

COMMONLY SUSCEPTIBLE SPECIES

Bacillus anthracis (1)

Aeromonas spp.

Citrobacter koseri

Francisella tularensis

Haemophilus ducreyi

Enterobacter cloacae

Klebsiella pneumoniae* Morganella morganii*

Neisseria gonorrhoeae*

Pseudomonas aeruginosa*

Pseudomonas fluorescens

Anaerobic micro-organisms

INHERENTLY RESISTANT ORGANISMS

Aerobic Gram-positive micro-organisms

Serratia marcescens*

Peptostreptococcus spp.

Propionibacterium acnes

Escherichia coli*

Klebsiella oxytoca

Proteus mirabilis*

Proteus vulgaris*

Providencia spp.

Actinomyces

compound.

enzymes.

Elimination

extent, faecally.

Cinroflovacin

Enteroccus faecium

Haemophilus influenzae*

Brucella spp.

Aerobic Gram-positive micro-organisms

Aerobic Gram-negative micro-organisms

Legionella spp. Moraxella catarrhalis* Neisseria meningitidis Pasteurella spp. Salmonella spp.* Shigella spp. Vibrio spp. Yersinia pestis Anaerobic micro-organisms Mobiluncus Other micro-organisms Chlamydia trachomatis (\$) Chlamydia pneumoniae (\$) Mycoplasma hominis (\$) Mycoplasma pneumoniae (\$) SPECIES FOR WHICH ACQUIRED RESISTANCE MAY BE A PROBLEM Aerobic Gram-positive micro-organisms Enterococcus faecalis (\$) Staphylococcus spp. *(2) Aerobic Gram-negative micro-organisms Acinetobacter baumannii+ Burkholderia cepacia + Campylobacter spp.+' Citrobacter freundii* Enterobacter aerogenes

Listeria monocytogenes Aerobic Gram-negative micro-organisms Stenotrophomonas maltophilia Anaerobic micro-organisms Excepted as listed above Other micro-organisms Mycoplasma genitalium Ureaplasma urealitycum Clinical efficacy has been demonstrated for susceptible isolates in approved clinical indications Resistance rate ≥ 50% in one or more EU countries Natural intermediate susceptibility in the absence of acquired mechanism of resistance Studies have been conducted in experimental animal infections due to inhalations of Bacillus anthracis spores; these studies reveal that antibiotics starting early after exposition avoid the occurrence of the disease if the treatment is made up to the decrease of the number of spores in the organism under the infective dose. The recommended use in human subjects is based primarily on in-vitro susceptibility and on animal experimental data together with limited human data. Two-month treatment duration in adults with oral ciprofloxacin given at the following dose, 500 mg bid, is considered as effective to prevent anthrax infection in humans. The treating physician should refer to national and /or international consensus documents regarding treatment of anthrax. Methicillin-resistant S. aureus very commonly express co-resistance to fluoroquinolones. The rate of resistance to methicillin is around 20 to 50% among all staphylococcal species and is usually higher in nosocomial isolates. Pharmacokinetic properties Absorption Following an intravenous infusion of ciprofloxacin the mean maximum serum concentrations were achieved at the end of infusion. Pharmacokinetics of ciprofloxacin were linear over the dose range up to 400 mg administered intravenously. Comparison of the pharmacokinetic parameters for a twice a day and three times a day intravenous dose regimen indicated no evidence of drug accumulation for ciprofloxacin and its metabolites. A 60-minute intravenous infusion of 200 mg ciprofloxacin or the oral administration of 250 mg ciprofloxacin, both given every 12 hours, produced an equivalent area under the serum concentration time curve (AUC). A 60-minute intravenous infusion of 400 mg ciprofloxacin every 12 hours was bioequivalent to a 500 mg oral dose every 12 hours with regard to AUC. The 400 mg intravenous dose administered over 60 minutes every 12 hours resulted in a Cmax similar to that observed with a 750 mg oral dose. A 60-minute infusion of 400 mg ciprofloxacin every 8 hours is equivalent with respect to AUC to 750 mg oral regimen given every 12 hours. Distribution Protein binding of ciprofloxacin is low (20 - 30%). Ciprofloxacin is present in plasma largely in a non-ionised form and has a large steady state distribution volume of 2 - 3 L/kg body weight. Ciprofloxacin reaches high concentrations in a variety of tissues such as lung (epithelial fluid, alveolar macrophages, biopsy tissue), sinuses, inflamed lesions (cantharides blister fluid), and the urogenital tract (urine, prostate, endometrium) where total concentrations exceeding those of plasma concentrations are reached. Biotransformation

Ciprofloxacin	61.5	15.2			
Metabolites (M1-M4)	9.5	2.6			
Renal clearance is betwee is between 480 - 600 mL/l filtration and tubular secrincreased half lives of cipinon-renal clearance of cipinecretion and metabolism Ciprofloxacin is present in Paediatric patients	kg/h. Ciprofloxacin undergretion. Severely impaired r rofloxacin of up to 12 h. profloxacin is mainly due 1. 1% of the dose is excreto	goes both glomerular enal function leads to to active trans-intestinal ed via the biliary route.			
	in naodiatric nationts ar	limitad			
The pharmacokinetic data In a study in children C _{max} of age). No notable increa	and AUC were not age-dese in C_{max} and AUC upon r	pendent (above one year			
kg three times daily) was	observed.				
In 10 children with severe after a 1-hour intravenou 1 year compared to 7.2 m 1 and 5 years of age. The mg*h/L) and 16.5 mg*h/L groups.	sepsis C _{max} was 6.1 mg/L s infusion of 10 mg/kg in g/L (range 4.7 - 11.8 mg/L AUC values were 17.4 mg	children aged less than .) for children between *h/L (range 11.8 - 32.0			
These values are within the Based on population phan various infections, the pre hours and the bioavailabi	rmacokinetic analysis of pedicted mean half-life in c	aediatric patients with hildren is approx. 4 - 5			
5.3 Preclinical safety d	ata				
Non-clinical data reveal n		nans based on			
conventional studies of single dose toxicity, repeated dose toxicity,					
carcinogenic potential, or toxicity to reproduction.					
- •					

comparable to that of other gyrase inhibitors.

6. PHARMACEUTICAL PARTICULARS

pH (pH of ciprofloxacin solutions: 3.9 - 4.5).

are the responsibility of the user.

6.4 Special precautions for storage

Articular tolerability

6.3 Shelf life

36 months.

Excretion of ciprofloxacin (% of dose)

Urine

Intravenous Administration

large weight-bearing joints in immature animals. The extent of the cartilage

damage varies according to age, species and dose; the damage can be

reduced by taking the weight off the joints. Studies with mature animals

beagle dogs, ciprofloxacin caused severe articular changes at therapeutic

(rat, dog) revealed no evidence of cartilage lesions. In a study in young

doses after two weeks of treatment, which were still observed after 5

freeze. Not to be stored above 30°C. 6.5 Nature and contents of container Pack sizes of 5 bottles (N2) containing 100 ml of solution for infusion each 6.6 Special precautions for disposal and other handling The ciprofloxacin infusion solution is compatible with Ringer solution, Ringer lactate solution, 5 % and 10 % glucose solutions, and 5 % and 10 % fructose solutions. When ciprofloxacin infusion solutions are mixed with compatible infusion solutions, for microbial reasons and light sensitivity these solutions must be administered shortly after admixture.

As the infusion solution is sensitive to light, only remove the bottles from the folding box for use. In daylight the full efficacy of the solution is guaranteed over a period of 3 days. For single use only. At cool temperatures precipitation may occur, which will re-dissolve at room temperature (15°C - 25°C). For ease of use the infusion vial stopper should be penetrated in the central ring. Penetration of the outer ring may result in damage to the vial stopper. Any unused solution should be disposed off. 7. Manufacturer Bayer Pharma AG Site: 51368 Leverkusen, Germany. 8. DATE OF REVISION OF THE TEXT September, 2013. 9. GENERAL CLASSIFICATION FOR SUPPLY

A medicament is a product which affects your health and its consumption contrary to instructions is dangerous for you.

instructions of the pharmacist who sold the medicament.

Medicinal product subject to medical prescription

Keep medicament out of reach of children.

il of Arab Health Minister

Bayer Pharma AG, Germany

This is a medicament

Bayer

Faeces

Keep in the outer carton in order to protect from light. Do not refrigerate or

Do not by yourself interrupt the period of treatment prescribed. Do not repeat the same prescription without consulting your doctor. Union of Arab Pharmacists Bayer

Chemical and physical in-use stability has been demonstrated for 24 hours view, unless the method of opening and mixing with co-infusion solutions precludes the risk of microbial contamination, the product should be used immediately. If not used immediately, in-use storage times and conditions

Follow strictly the doctor's prescription, the method of use and the

An overdose of 12 g has been reported to lead to mild symptoms of toxicity. An acute overdose of 16 g has been reported to cause acute renal failure. Symptoms in overdose consist of dizziness, tremor, headache, tiredness, seizures, hallucinations, confusion, abdominal discomfort, renal and hepatic impairment as well as crystalluria and haematuria. Reversible renal toxicity Apart from routine emergency measures, e.g. ventricular emptying followed by medical carbon, it is recommended to monitor renal function, including urinary pH and acidify, if required, to prevent crystalluria. Patients should be kept well hydrated. Calcium or magnesium containing antacids may In the event of overdose, symptomatic treatment should be implemented. ECG monitoring should be undertaken, because of the possibility of QT

Breakpoints separate susceptible strains from strains with intermediate Resistant R > 1 mg/LR > 1 mg/L

R > 1 mg/L

R > 1 mg/L

R > 0.5 mg/L

R > 0.06 mg/L

R > 0.06 mg/L

R > 1 mg/L

time for selected species and local information on resistance is desirable. particularly when treating severe infections. As necessary, expert advice should be sought when the local prevalence of resistance is such that the utility of the agent in at least some types of infections is questionable. Groupings of relevant species according to ciprofloxacin susceptibility (for

Like a number of other quinolones, ciprofloxacin is phototoxic in animals photocarcinogenicity show a weak photomutagenic or phototumorigenic effect of ciprofloxacin in-vitro and in animal experiments. This effect was

The doctor and the pharmacist are experts in medicine, its benefits

Low concentrations of four metabolites have been reported, which were identified as: desethyleneciprofloxacin (M 1), sulphociprofloxacin (M 2), oxociprofloxacin (M 3) and formylciprofloxacin (M 4). The metabolites display in-vitro antimicrobial activity but to a lower degree than the parent Ciprofloxacin is known to be a moderate inhibitor of the CYP 450 1A2 iso-Ciprofloxacin is largely excreted unchanged both renally and, to a smaller

at clinically relevant exposure levels. Data on photomutagenicity/ As reported for other gyrase inhibitors, ciprofloxacin causes damage to the

at room temperature (15 °C to 25 °C). From a microbiological point of