SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Ciprofloxacin 200 mg/100 mL solution for infusion- Nircip

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each 100 mL contains

Ciprofloxacin BP

200mg

Water for Injections

3. PHARMACEUTICAL FORM

Solution for Infusion.

The solution is clear

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Ciprofloxacin 200 mg/100 mL solution for infusion is indicated for the treatment of the following infections (see sections 4.4 and 5.1). Special attention should be paid to available information on resistance to ciprofloxacin before commencing therapy.

Consideration should be given to official guidance on the appropriate use of antibacterial agents.

Adults

- Lower respiratory tract infections due to Gram-negativebacteria
 - exacerbations of chronic obstructive pulmonary disease

 In exacerbation of chronic obstructive pulmonary disease Ciprofloxacin should be used only when it is considered inappropriate to use other antibacterial agents that are commonly recommended for the treatment of these infections.
 - Broncho-pulmonary infections in cystic fibrosis or inbronchiectasis
 - pneumonia
- Chronic suppurative otitismedia
- Acute exacerbation of chronic sinusitis especially if these are caused by Gramnegativebacteria

- Urinary tractinfections
 - Acute pyelonephritis
 - Complicated pyelonephritis
- Bacterial prostatitis
- Genital tract infections
 - epididymo-orchitis including cases due to susceptible Neisseria gonorrhoeae
 - pelvic inflammatory disease including cases due to susceptible Neisseria gonorrhoeae
- Infections of the gastrointestinal tract (e.g. travellers` diarrhoea)
- Intra-abdominal infections
- Infections of the skin and soft tissue caused by Gram-negative bacteria
- Malignant external otitis
- Infections of the bones and joints
- Inhalation anthrax (post-exposure prophylaxis and curative treatment)

Ciprofloxacin may be used in the management of neutropenic patients with fever that issuspected to be due to a bacterial infection.

Children and adolescents

- Broncho-pulmonary infections in cystic fibrosis caused by Pseudomonasaeruginosa
- Complicated urinary tract infections and pyelonephritis
- Inhalation anthrax (post-exposure prophylaxis and curativetreatment)

Ciprofloxacin may also be used to treat severe infections in children and adolescents when this is considered to be necessary.

Treatment should be initiated only by physicians who are experienced in the treatment of cystic ections in children and adolescents (see sections 4.4 and 5.1).

4.2 Posology and method of administration

Posology

The dosage is determined by the indication, the severity and the site of the infection, the susceptibility to ciprofloxacin of the causative organism(s), the renal function of the patient and, in children and adolescents the body weight.

The duration of treatment depends on the severity of the illness and on the clinical and bacteriological course.

After intravenous initiation of treatment, the treatment can be switched to oral treatment with tablet or suspension if clinically indicated at the discretion of the physician. IV treatment should be followed by oral route as soon as possible.

In severe cases or if the patient is unable to take tablets (e.g. patients on enteral nutrition), it is recommended to commence therapy with intravenous ciprofloxacin until a switch to oral administration is possible.

Treatment of infections due to certain bacteria (e.g. Pseudomonas aeruginosa, Acinetobacter or Staphylococci) may require higher ciprofloxacin doses and co-administration with other appropriate antibacterial agents.

Treatment of some infections (e.g. pelvic inflammatory disease, intra-abdominal infections, infections in neutropenic patients and infections of bones and joints) may require co-administration with other appropriate antibacterial agents depending on the pathogens involved.

Adults

			Total duration of
Indications		Daily dose in mg	treatment (including
			switch to oral therapy as
Infections of the	lower respiratory tract	400 mg twice daily	7 to 14 days
		to 400 mg three	
		times a day	
Infections of	Acute exacerbation of	400 mg twice daily	7 to 14 days
the upper	chronic sinusitis	to 400 mg three	
respiratory		times a day	
tract	Chronic suppurative	400 mg twice daily	7 to 14 days
	otitis media	to 400 mg three	
		times a day	
	Malignant external otitis	400 mg three	28 days up to 3 months
		times a day	
Urinary tract	Complicated and	400 mg twice	7 to 21 days, it can be
infections	uncomplicated	daily to 400 mg	continued for longer than 21
	pyelonephritis	three times a	days in some specific
		dav	circumstances (such as
	Prostatitis	400 mg twice	2 to 4 weeks (acute)
		daily to 400 mg	
		three times a	

Genital tract	Epididymo-orchitis and	400 mg twice	at least 14 days
infections	pelvic inflammatory	daily to 400 mg	
	diseases	three times a	
Infections of	Diarrhoea caused by	400 mg twice	1 day
the gastro-	bacterial pathogens	daily	
intestinal tract	including Shigella spp.		
and intra-	other than		
abdominal	Shigelladysenteriaetype		
infections	1 and empirical		
	treatment of severe		
	travellers' diarrhoea		
	Diarrhoea caused by	400 mg twice	5 days
	Shigelladysenteriae type	daily	
	Diarrhoea caused by	400 mg twice	3 days
	Vibrio cholerae	daily	
	Typhoid fever	400 mg twice	7 days
		daily	
	Intra-abdominal	400 mg twice	5 to 14 days
	infections due to Gram-	daily to 400 mg	
	negative bacteria	three times a	
		day	
Infections of the	skin and soft tissue	400 mg twice	7 to 14 days
		daily to 400 mg	
		three times a day	
Bone and joint in	Bone and joint infections		max. of 3 months
		three times a day	
Neutropenic pat	ients with fever that is	400 mg twice	Therapy should be continued
suspected to be d	lue to a bacterial infection.	daily to 400 mg	over the entire period of
Ciprofloxacin sh	nould be co-administered	three times a day	neutropenia
with appropriate	antibacterial agent(s) in		
accordance to off	icial guidance.		

Inhalation	anthrax	post-exposure	400	mg	twice	60 days from the confirmation
prophylaxis	and curative	treatment for	daily			of Bacillus anthracis exposure
persons requiring parenteral treatment						
Drug administration should begin as soon						
as possible after suspected or confirmed						

Children and adolescents

		Total duration of treatment
Indications	Daily dose in mg	(including switch to oral therapy
		as soon as possible)
Cystic fibrosis	10 mg/kg body weight three times	10 to 14 days
	a day with a maximum of 400 mg	
	per dose.	
Complicated urinary	6 mg/kg body weight three timesa	10 to 21 days
tract infections and	day to 10 mg/kg body weight	
pyelonephritis	three times a day with a maximum	
	of 400 mg perdose.	
Inhalation anthrax	10 mg/kg body weight twice daily	60 days from the confirmation of
post- exposure	to 15 mg/kg body weight twice	Bacillus anthracis exposure
curative treatment for	daily with a maximum of 400 mg	
persons requiring	per dose.	
parenteral treatment		
Drug administration		
should begin as soon as		
possible after suspected		
or confirmed exposure.		
O(1	10 /	A diagram
Other severe infections		According to the type of infections.
	a day with a maximum of 400 mg	
	per dose.	

Elderly patients

Geriatric patients should receive a dose selected according to the severity of the infection and the patient's creatinine clearance.

Renal and hepatic impairment

Recommended starting and maintenance doses for patients with impaired renal function:

Creatinine Clearance	Serum Creat	inine Intravenous Dose [mg]
[mL/min/1.73 m2]	[µmol/l]	
> 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 function no dose adjustment is required.

Dosing in children with impaired renal and/or hepatic function has not been studied.

Method of administration

Ciprofloxacin should be checked visually prior to use. It must not be used if cloudy.

Ciprofloxacin should be administered by intravenous infusion. For children, the infusion duration is 60 minutes.

In adult patients, infusion time is 60 minutes for 400 mg Ciprofloxacin and 30 minutes for 200 mg Ciprofloxacin. Slow infusion into a large vein will minimize patient discomfort and reduce the risk of venous irritation.

The infusion solution can be infused either directly or after mixing with other compatible infusion solutions

4.3 Contraindications

Hypersensitivity to the active substance, to other quinolones or to any of the excipients listen in section 6.1.

Concomitant administration of ciprofloxacin and tizanidine (see section 4.5).

4.4 Special warnings and precautions for use

The use of ciprofloxacin should be avoided in patients who have experienced serious adversereactions in the past when using quinolone or fluoroquinolone containing products (see section

4.8). Treatment of these patients with ciprofloxacin should only be initiated in the absence of alternative treatment options and after careful benefit/risk assessment (see section 4.3).

<u>Severe infections and mixed infections with Gram-positive and anaerobic pathogens</u>Ciprofloxacin monotherapy is not suited for treatment of severe infections and infections that might be due to Gram-positive or anaerobic pathogens. In such infections ciprofloxacin must be co-administered with other appropriate antibacterial agents.

<u>Streptococcal Infections (including Streptococcus pneumoniae)</u>

Ciprofloxacin is not recommended for the treatment of streptococcal infections due to inadequate efficacy.

Genital tract infections

Epididymo-orchitis and pelvic inflammatory diseases may be caused by fluoroquinolone-resistant Neisseria gonorrhoeae. For epididymo-orchitis and pelvic inflammatory diseases, empirical ciprofloxacin should only be considered in combination with another appropriate antibacterial agent (e.g. a cephalosporin) unless ciprofloxacin-resistant Neisseria gonorrhoeae can be excluded. If clinical improvement is not achieved after 3 days of treatment, the therapy should be reconsidered.

<u>Urinary tract infections</u>

Resistance to fluoroquinolones of Escherichia coli – the most common pathogen involved in urinary tract infections – varies across the European Union. Prescribers are advised to take into account the local prevalence of resistance in Escherichia coli to fluoroquinolones.

Intra-abdominal infections

There are limited data on the efficacy of ciprofloxacin in the treatment of post-surgical intraabdominal infections.

Travelers' diarrhea

The choice of ciprofloxacin should take into account information on resistance to ciprofloxacin in relevant pathogens in the countries visited.

<u>Infections of the bones and joints</u>

Ciprofloxacin should be used in combination with other antimicrobial agents depending on the results of the microbiological documentation.

Inhalational anthrax

Use in humans is based on in-vitro susceptibility data and on animal experimental data together with limited human data. Treating physicians should refer to national and /or international consensus documents regarding the treatment of anthrax.

Paediatric population

Ciprofloxacin treatment should be initiated only by 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 possible adverse events related to joints and/or surrounding tissue.

Broncho-pulmonary infections in cystic fibrosis

Clinical trials have included children and adolescents aged 5-17 years. More limited experience is available in treating children between 1 and 5 years of age.

Complicated urinary tract infections and pyelonephritis

Ciprofloxacin treatment of urinary tract infections should be considered when other treatments cannot be used, and should be based on the results of the microbiological documentation.

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

Other specific severe infections

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 documentation can justify a ciprofloxacin use.

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.

Prolonged, disabling and potentially irreversible serious adverse drug reactions. Very rare cases of prolonged (continuing months or years), disabling and potentially irreversible serious adverse drug reactions affecting different, sometimes multiple, body systems (musculoskeletal, nervous, psychiatric and senses) have been reported in patients receiving quinolones and fluoroquinolones irrespective of their age and pre-existing risk factors.

Ciprofloxacin should be discontinued immediately at the first signs or symptoms of any serious adverse reaction and patients should be advised to contact their prescriber for advice.

Tendinitis and tendon rupture

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 organism and evaluation of the risk/benefit balance, ciprofloxacin may be prescribed to these patients for the treatment of certain severe 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 but not limited to Achilles tendon), sometimes bilateral, may as early as within 48 hours of starting treatment with quinolones and fluoroquinolones and have been reported to occur even up to several months after discontinuation of treatment (see section 4.8). The risk of tendinitis and tendon rupture is increased in older patients, patients with renal impairment, patients with solid organ transplants, and those treated concurrently with corticosteroids. Therefore, concomitant use of corticosteroids should be avoided.

At the first sign of tendinitis (e.g. painful swelling, inflammation), the treatment with ciprofloxacin should be discontinued and alternative treatment should be considered. The affected limb(s) should be appropriately treated (immobilisation). Corticosteroids should not be used if signs of tendinopathy occur.

Patients with myasthenia gravis

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

Aortic aneurysm and dissection, and heart valve regurgitation/incompetence

Epidemiologic studies report an increased risk of aortic aneurysm and dissection, particularly in elderly patients, and of aortic and mitral valve regurgitation after intake of fluoroquinolones. Cases of aortic aneurysm and dissection, sometimes complicated by rupture (including fatal ones), and of regurgitation/incompetence of any of the heart valves have been reported in patients receiving fluoroquinolones (see section 4.8).

Therefore, fluoroquinolones should only be used after a careful benefit/risk assessment and after consideration of other therapeutic options in patients with positive family history of aneurysm disease or congenital heart valve disease, or in patients diagnosed with pre-existing aortic aneurysm and/or dissection or heart valve disease, or in presence of other risk factors or conditions predisposing

- for both aortic aneurysm and dissection and heart valve regurgitation/incompetence (e.g. connective tissue disorders such as Marfan syndrome or Ehlers-Danlos syndrome, Turner syndrome, Behcet's disease, hypertension, rheumatoid arthritis) or additionally
- for aortic aneurysm and dissection (e.g. vascular disorders such as Takayasu arteritis or giant cell arteritis, or known atherosclerosis, or Sjögren's syndrome) or additionally
- for heart valve regurgitation/incompetence (e.g. infective endocarditis).

The risk of aortic aneurysm and dissection, and their rupture may also be increased in patients treated concurrently with systemic corticosteroids.

In case of sudden abdominal, chest or back pain, patients should be advised to immediately consult a physician in an emergency department.

Patients should be advised to seek immediate medical attention in case of acute dyspnoea, new onset of heart palpitations, or development of oedema of the abdomen or lower extremities.

Vision disorders

If vision becomes impaired or any effects on the eyes are experienced, an eye specialist should be consulted immediately.

Photosensitivity

Ciprofloxacin has been shown to cause photosensitivity reactions. Patients taking ciprofloxacin should be advised to avoid direct exposure to either extensive sunlight or UV on 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

Psychiatric reactions may occur even after the 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 should be discontinued.

Peripheral neuropathy

Cases of polyneuropathy (based on neurological symptoms such as pain, burning, sensory disturbances or muscle weakness, alone or in combination) have been reported in patients receiving ciprofloxacin. Ciprofloxacin should be discontinued in patients experiencing

symptoms of neuropathy, including pain, burning, tingling, numbness, and/or weakness in order to prevent the development of an irreversible condition.

Cardiac disorders

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

- congenital long QT syndrome
- concomitant use of drugs that are known to prolong the QT interval (e.g. Class IA and III anti-arrhythmics, tricyclic antidepressants, macrolides, antipsychotics)
- uncorrected electrolyte imbalance (e.g. hypokalaemia, hypomagnesaemia)
- 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).

Dysglycaemia

As with all quinolones, disturbances in blood glucose, including both hypoglycaemia and hyperglycaemia have been reported (see section 4.8), usually in elderly diabetic patients, receiving concomitant treatment with an oral hypoglycaemic agent (e.g. glibenclamide) or with insulin. Cases of hypoglycaemic coma have been reported. In diabetic patients, careful monitoring of blood glucose is recommended.

Gastrointestinal System

The occurrence of severe and persistent diarrhoea during or after treatment (including several weeks after treatment) may indicate an antibiotic-associated 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. Antiperistaltic drugs are contraindicated in this situation.

Renal and urinary system

Crystalluria related to the use of ciprofloxacin has been reported (see section 4.8). Patients receiving ciprofloxacin should be well hydrated and excessive alkalinity of the urine should be avoided.

Impaired renal function

Since ciprofloxacin is largely excreted unchanged via renal pathway dose adjustment is needed in patients with impaired renal function as described in section 4.2 to avoid an increase in adverse drug reactions due to accumulation of ciprofloxacin.

Hepatobiliary system

Cases of hepatic necrosis and life-threatening hepatic failure have been reported with ciprofloxacin (see section 4.8). In the event of any signs and symptoms of hepatic disease (such as anorexia, jaundice, dark urine, pruritus, or tender abdomen), treatment should be discontinued.

Glucose-6-phosphate dehydrogenase deficiency

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 should be monitored.

Resistance

During or following a course of treatment with ciprofloxacin bacteria that 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, agomelatine). Co-administration of ciprofloxacin and tizanidine is contra-indicated. 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 necessary (see section 4.5).

Methotrexate

The concomitant use of ciprofloxacin with methotrexate is not recommended (see section 4.5).

Interaction with tests

The in-vitro activity of ciprofloxacin against Mycobacterium tuberculosis might give false negative bacteriological test results in specimens from patients currently taking ciprofloxacin.

<u>Injection Site Reaction</u>

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 reactions which resolve rapidly upon completion of the infusion. Subsequent intravenous administration is not contraindicated unless the reactions recur or worsen.

NaCl Load

In patients for whom sodium intake is of medical concern (patients with congestive heart failure, renal failure, nephrotic syndrome, etc.), the additional sodium load should be taken into account (for sodium chloride content, see section 2).

4.5 Interaction with other medicinal products and other forms of interaction

Effects of other medicinal 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

Tizanidine must not be administered together with ciprofloxacin. In a clinical study with healthy subjects, there was an increase in 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 potentiated hypotensive and sedative effect.

Methotrexate

Renal tubular transport of methotrexate may be inhibited by concomitant administration of ciprofloxacin, potentially leading to increased plasma levels of methotrexate and increased risk of methotrexate-associated toxic reactions. The concomitant use is not recommended.

Theophylline

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.

Other xanthine derivatives

On concurrent administration of ciprofloxacin and caffeine or pentoxifylline (oxpentifylline), raised serum concentrations of these xanthine derivatives were reported.

Phenytoin

Simultaneous administration of ciprofloxacin and phenytoin may result in increased or reduced serum levels of phenytoin such that monitoring of drug levels is recommended.

Oral anticoagulants

Simultaneous administration of ciprofloxacin with warfarin may augment its anti-coagulant effects. There have been many reports of increases in oral anti-coagulant activity in patients receiving antibacterial agents, including fluoroquinolones. The risk may vary with the underlying infection, age and general status of the patient so that the contribution of the fluoroquinolone to the increase in INR (international normalised ratio) is difficult to assess. It is recommended that the INR should be monitored frequentl during and shortly after coadministration of ciprofloxacin with an ora anticoagulant agent.

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 anticoagulant 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 normalized ratio) is difficult to assess. The INR should be monitored frequently during and shortly after coadministration of ciprofloxacin with a vitamin K antagonist (e.g., warfarin, acenocoumarol, phenprocoumon, or fluindione).

Duloxetine

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 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).

Lidocaine

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 administration.

Clozapine

Following concomitant administration of 250 mg ciprofloxacin with clozapine for 7 days, serum concentrations of clozapine and 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.

Sildenafil

Cmax 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.

Agomelatine

In clinical studies, it was demonstrated that fluvoxamine, as a strong inhibitor of the CYP450 1A2 isoenzyme, markedly inhibits the metabolism of agomelatine resulting in a 60-fold increase of agomelatine exposure. Although no clinical data are available for a possible interaction with ciprofloxacin, a moderate inhibitor of CYP450 1A2, similar effects can be expected upon concomitant administration ('Cytochrome P450' in section 'Special warnings and precautions for use).

Zolpidem

Co-administration ciprofloxacin may increase blood levels of zolpidem, concurrent use is not recommended.

4.6 Fertility, pregnancy and lactation

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 cartilage in the human immature organism / foetus (see section 5.3). As a precautionary measure, it is preferable to avoid the use of ciprofloxacin during pregnancy.

Lactation

Ciprofloxacin is excreted in breast milk. Due to the potential risk of articular damage, ciprofloxacin should not be used during breast-feeding.

4.7 Effects on ability to drive and use machines

Due to its neurological effects, ciprofloxacin may affect reaction time. 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 and diarrhea, vomiting, transient increase in transaminases, rash, and injection and infusion site reactions. ADRs derived from clinical studies and post-marketing surveillance with ciprofloxacin (oral, intravenous and sequential therapy) sorted by categories of frequency are listed below. The frequency analysis takes into account data from both oral and intravenous administration of ciprofloxacin.

System Class	Common	Uncommon	Rare	Very	Frequency not
Organ	(≥1/100	(≥1/1,000 To	(≥1/10,000	Torare(<1/10,000)	known (cannot be
	То	<1/100)	<1/1,000)		estimated from
	<1/10)				the available
					data)
Infections and		Mycotic	Antibiotic		
infestations		superinfections	associated colitis		
			(very rarely with		
			possible fatal		
			outcome) (see		
			section 4.4)		

Blood and	Eosinophilia	Leukopenia	Haemolyticanaemi	
lymphatic system		Anaemia	a Agranulocytosis	
disorders		Neutropenia	Pancytopenia	
		Leukocytosis	(life-threatening)	
		Thrombocytopenia	Bonemarrow	
		Thrombocytaemia	depression (life-	
			threatening)	
Immune system		Allergic reaction	Anaphylactic	
disorders		Allergic oedema /	reaction	
		angioedema	Anaphylactic	
			shock (life-	
			threatning) Serum	
			sickness- like	
			reaction	
Metabolism and	Anorexia	Hyperglycaemia		
nutrition disorders				
Psychiatric	Psychomotor	Confusion and	Psychotic	Mania, hypomania
disorders	hyperactivity /	disorientation	reactions (see	
	agitation	Anxiety reaction	section 4.4)	
		Abnormal dreams		
		Depression		
		(potentially		
		culminating in		
		suicidal		
		ideations/thoughts or		
Nervous system	Headache	Par- and	Migraine	Peripheral
disorders	Dizziness Sleep	dysaesthesiaHypoaest	disturbed	neuropathy (see
	disorders	hesia Tremor	coordination Gait	section 4.4)
	Tastedisorders	Seizures (see	disturbance	
		section 4.4) Vertigo	Olfactory nerve	
			disorders	
			Intracranial	
			hypertension	

Eye disorders			Visual disturbances	Visual colour	
			(e.g. diplopia)	distortions	
Ear and			Tinnitus		
labyrinth			Hearing loss /		
disorders			hearingimpaired		
Cardiac disorders			Tachycardia		Ventricular
			-		arrhythmia,
					QT
					prolongation,
					torsades de
					pointes *
					pointes .
Vascular			Vasodilatation	Vasculitis	
disorders			Hypotension		
			Syncope		
Respiratory,			Dyspnoea		
thoracic and			(including		
mediastinal			asthmatic		
disorders	3. 7	T.7	condition)	D	
Gastrointestinal	Nausea	Vomiting	Antibiotic	Pancreatitis	
disorders	Diarrhoea	Gastrointestinal	associated colitis		
			(very rarely with		
		pains Dyspepsia			
		Flatulence	outcome)		
Hepatobiliary		Increase in	Hepatic	Liver necrosis	
disorders		transaminases	impairment	(very rarely	
		Increased	Cholestatic icterus	progressing to	
		bilirubin		life-threatening	
				hepatic failure)	
				(see section 4.4)	
				(See Section 4.4)	

Skin and	Rash	Photosensitivity	Petechia	Acute generalised
subcutaneous	Pruritus	reactions (see	Erythema	exanthematouspus
tissue disorders	Urticaria	section 4.4)	multiforme	tulosis (AGEP),
			Erythema	DRESS
			nodosum	
			Stevens-Johnson	
			syndrome	
			(potentially life-	
			threatening)	
			Toxic epidermal	
			necrolysis	
			(potencially life-	
			threatening)	
Musculoskeletal,	Musculoskelete	Myalgia Arthritis	Muscular	
connective tissue	lal pain	Increased muscle	weakness	
and bone	(e.g.extremity	tone andcramping	Tendinitis	
disorders	pain, back pain,		Tendon rupture	
	chest pain)		(predominantly	
	Arthralgia		Achilles	
			tendon) (see	
			section 4.4)	
			Exacerbation of	
			symptoms of	
			myasthenia	
			gravis (see	
			section 4.4)	
D 1 1	D 1	D 16.7	,	
Renal and	Renal	Renal failure		
urinarydisorders	impairment	Haematuria		
		Crystalluria (see		
		section 4.4)		

General disorders	Injection	Asthenia	Oedema Sweating	
and administration	and	Fever	(hyperhidrosis)	
siteconditions	infusion			
	site			
	reactions			
	(only			
	intraveno			
	us			
	administr			
	ation)			
Investigations		Increase in	Increase amylase	International
		blood alkaline		normalised ratio
		phosphatase		increased
				(in patients
				treated with
				Vitamin K
				antagonists)

*Very rare cases of prolonged (up to months or years), disabling and potentially irreversible serious drug reactions affecting several, sometimes multiple, system organ classes and senses (including reactions such as tendonitis, tendon rupture, arthralgia, pain in extremities, gait disturbance, neuropathies associated with paraesthesia, depression, fatigue, memory impairment, sleep disorders, and impairment of hearing, vision, taste and smell) have been reported in association with the use of quinolones and fluoroquinolones in some cases irrespective of pre-existing risk factors (see section 4.4).

**Cases of aortic aneurysm and dissection, sometimes complicated by rupture (including fatal ones), and of regurgitation/incompetence of any of the heart valves have been reported in patients receiving fluoroquinolones (see section 4.4).

The following undesirable effects have a higher frequency category in the subgroups of patients receiving intravenous or sequential (intravenous to oral) treatment:

Common	Vomiting, Transien,t increase in transaminases, Rash
Uncommon	Thrombocytopenia
	Thrombocytaemia Confusion and

	disorientation Hallucinations			
	Par- and dysaesthesia			
	Seizures			
	Vertigo			
	Visual disturbances			
	Hearing loss			
	Tachycardia			
	Vasodilatation			
	Hypotension			
	Transient hepatic impairment			
	Cholestatic icterus			
	Renal failure			
	Oedema			
Rare	Pancytopenia			
	Bone marrow depression			
	Anaphylactic shock			
	Psychotic reactions			
	Migraine			
	Olfactory nerve disorders			
	Hearing impaired			
	Vasculitis			
	Pancreatitis			
	Liver necrosis			
	Petechiae			
	Tendon rupture			

Paediatric population

The incidence of arthropathy, mentioned above, is referring to data collected in studies with adults. In children, arthropathy is reported to occur commonly. (see section 4.4).

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorization of the medicinal product is important. Report ADR on info@aculife.co.in

4.9 Overdose

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 has been reported.

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 theoretically reduce the absorption of ciprofloxacin in overdoses.

Only a small quantity of ciprofloxacin (< 10%) is eliminated by haemodialysis or peritoneal dialysis.

In the event of overdose, symptomatic treatment should be implemented. ECG monitoring should be undertaken, because of the possibility of QT interval prolongation.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Fluoroquinolones, ATC code: J01MA02

Mechanism of action

As a fluoroquinolone antibacterial agent, the bactericidal action of ciprofloxacin results from the inhibition of both type II topoisomerase (DNA-gyrase) and topoisomerase IV, required for bacterial DNA replication, transcription, repair and recombination.

Pharmacokinetic/pharmacodynamic relationship

Efficacy mainly depends on the relation between the maximum concentration in serum (Cmax) and the minimum inhibitory concentration (MIC) of ciprofloxacin for a bacterial pathogen and the relation between the area under the curve (AUC) and the MIC.

Mechanism of resistance

In-vitro resistance to ciprofloxacin can be acquired through a stepwise process by target site mutations in both DNA gyrase and topoisomerase IV. The degree of cross-resistance between ciprofloxacin and other fluoroquinolones that results is variable. Single mutations may not result in clinical resistance, but multiple mutations generally result in clinical 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 in-vitro 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:

Breakpoints separate susceptible strains from strains with intermediate susceptibility and the latter from resistant strains:

EUCAST Recommendations

Microorganisms	Susceptible	Resistant
Enterobacteria	$S \le 0.5 \text{ mg/L}$	R > 1 mg/L
Pseudomonas	$S \le 0.5 \text{ mg/L}$	R > 1 mg/L
Acinetobacter	$S \le 1 \text{ mg/L}$	R > 1 mg/L
Staphylococcus spp. 1	$S \le 1 \text{ mg/L}$	R > 1 mg/L
Haemophilus influenzae	$S \le 0.5 \text{ mg/L}$	R > 0.5 mg/L
and Moraxella catarrhalis		
Neisseria gonorrhoeae	$S \leq 0.03$	R > 0.06
Neisseria meningitidis	S ≤ 0.03	R > 0.06
Non-species-related	$S \le 0.5 \text{ mg/L}$	R > 1 mg/L

¹ Staphylococcus spp. - breakpoints for ciprofloxacin relate to high dose therapy.

The prevalence of acquired resistance may vary geographically and with 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 Streptococcus species see section 4.4)

COMMONLY SUSCEPTIBLE SPECIES		
Aerobic Gram-positive micro-organisms		
Bacillus anthracis (1)		

^{*} 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 testing is not recommended.

Aerobic Gram-negative micro-organisms
Aeromonasspp.
Brucella spp.
Citrobacterkoseri
Francisellatularensis
Haemophilusducreyi
Haemophilusinfluenzae*
Legionella spp.
Moraxella catarrhalis*
Neisseria meningitidis
Pasteurellaspp.
Salmonella spp.*
Shigellaspp. *
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+
Burkholderiacepacia +*
Campylobacter spp.+*
Citrobacterfreundii*
Enterobacter aerogenes
Enterobacter cloacae *
Escherichia coli*

Klebsiellaoxytoca		
Klebsiella pneumoniae*		
Morganellamorganii*		
Neisseria gonorrhoeae*		
Proteus mirabilis*		
Proteus vulgaris*		
Providencia spp.		
Pseudomonas aeruginosa*		
Pseudomonas fluorescens		
Serratiamarcescens*		
Anaerobic micro-organisms		
coccusspp.		
Propionibacterium acnes		
INHERENTLY RESISTANT ORGANISMS		
Aerobic Gram-positive micro-organisms		
Actinomyces		
Enteroccusfaecium		
Listeria monocytogenes		
Aerobic Gram-negative micro-organisms		
Stenotrophomonasmaltophilia		
<u>Anaerobic micro-organisms</u>		
Excepted as listed above		
Other micro-organisms		
Mycoplasma genitalium		
Ureaplasmaurealitycum		
* 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		

- 1. 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.

5.2 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

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 compound.

Ciprofloxacin is known to be a moderate inhibitor of the CYP 450 1A2 iso-enzymes.

Elimination

Ciprofloxacin is largely excreted unchanged both renally and, to a smaller extent, faecally.

Excretion of ciprofloxacin (% of dose)				
	Intravenous admir	Intravenous administration		
	Urine	Faeces		
Ciprofloxacin	61.5	15.2		
Metabolites (M1 – M4)	9.5	2.6		

Renal clearance is between 180-300 mL/kg/h and the total body clearance is between 480-600 mL/kg/h. Ciprofloxacin undergoes both glomerular filtration and tubular secretion. Severely impaired renal function leads to increased half lives of ciprofloxacin of up to 12 h. Non-renal clearance of ciprofloxacin is mainly due to active trans-intestinal secretion and metabolism. 1% of the dose is excreted via the biliary route. Ciprofloxacin is present in the bile in high concentrations.

Paediatric population

The pharmacokinetic data in paediatric patients are limited.

In a study in children Cmax and AUC were not age-dependent (above one year of age). No notable increase in Cmax and AUC upon multiple dosing (10 mg/kg three times daily) was observed.

In 10 children with severe sepsis Cmax was 6.1 mg/l (range 4.6-8.3 mg/l) after a 1-hour intravenous infusion of 10 mg/kg in children aged less than 1 year compared to 7.2 mg/l (range 4.7-11.8 mg/l) for children between 1 and 5 years of age. The AUC values were

17.4 mg h/l (range 11.8-32.0 mg h/l) and 16.5 mg h/l (range 11.0-23.8 mg h/l) in the respective age groups.

These values are within the range reported for adults at therapeutic doses. Based on population pharmacokinetic analysis of paediatric patients with various infections, the predicted mean half-life in children is approx. 4-5 hours and the bioavailability of the oral suspension ranges from 50 to 80%.

5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of single dose toxicity, repeated dose toxicity, carcinogenic potential, or toxicity to reproduction.

Like a number of other quinolones, ciprofloxacin is phototoxic in animals at clinically relevant exposure levels. Data on photomutagenicity / photocarcinogenicity show a weak photomutagenic or phototumorigenic effect of ciprofloxacin in-vitro and in animal experiments. This effect was comparable to that of other gyrase inhibitors.

Articular tolerability

As reported for other gyrase inhibitors, ciprofloxacin causes damage to the 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 (rat, dog) revealed no evidence of cartilage lesions. In a study in young beagle dogs, ciprofloxacin caused severe articular changes at therapeutic doses after two weeks of treatment, which were still observed after 5 months.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Sodium Chloride BP

Di-sodium EDTA

Citric Acid Monohydrate BP

Lactic Acid BP

Sodium Hydroxide BP

Hydrochloric Acid BP

Water for Injection BP

6.2 Incompatibilities

This medicinal product must not be mixed with other medicinal products.

Unless compatibility with other solutions/drugs has been confirmed, the infusion solution must always be administered separately. The visual signs of incompatibility are e.g.

precipitation, clouding, and discoloration.

Incompatibility appears with all infusion solutions/drugs that are physically or chemically unstable at the pH of the solutions (e.g. penicillins, heparin solutions), especially in combination with solutions adjusted to an alkaline pH (pH of ciprofloxacin solutions: 3.9 – 4.5).

6.3 Shelf life

36 Months from the date of manufacture

6.4 Special precautions for storage

Store below 30°C.Do not Freeze. Protect from light.

6.5 Nature and contents of container <and special equipment for use, administration or implantation>

100 mL plastic bottle

6.6 Special precautions for disposal <and other handling>

7. MARKETING AUTHORISATION HOLDER

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8. MARKETING AUTHORISATION NUMBER(S)

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Date of latest renewal: 06.12.2022

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Date: 12.07.2023