

Generic Name: Meropenem
Trade Name: MERONEM®
CDS Effective Date: January 03, 2024
Supersedes: October 31, 2018
Approved by BPOM:

PT PFIZER INDONESIA
Local Product Document

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Qualitative and quantitative composition

MERONEM® is presented as a sterile white powder containing meropenem; 500 mg or 1 g as the trihydrate blended with anhydrous sodium carbonate for constitution. MERONEM® injection contains 208 mg sodium carbonate for each gram of meropenem (anhydrous potency).

Vial for I.V. injection or infusion	MERONEM® 500 mg	MERONEM® 1 g
Active ingredient:		
Meropenem (as the trihydrate)	570 mg	1.14 g
Equivalent to anhydrous meropenem	500 mg	1 g
Excipient:		
Anhydrous sodium carbonate	104 mg	208 mg

For each gram of meropenem (anhydrous potency) the vial contains 90 mg (3.9 mmol) of sodium.

Pharmaceutical form

Powder for constitution for intravenous administration.

Appearance

White to light yellow powder.

Therapeutic indications

MERONEM® is indicated for treatment, in adults and children, of the following infections caused by single or multiple bacteria sensitive to meropenem.

- Pneumonias and nosocomial pneumonias
- Urinary tract infections
- Intra-abdominal infections
- Gynaecological infections, such as endometritis
- Skin and skin structure infections
- Meningitis
- Septicaemia
- Empiric treatment, for presumed infections in adult patients with febrile neutropenia, used as monotherapy or in combination with anti-viral or anti-fungal agents.

MERONEM® has proved efficacious alone or in combination with other antimicrobial agents in the treatment of polymicrobial infections.

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There is no experience in paediatric patients with neutropenia or primary or secondary immunodeficiency.

Posology and method of administration

Adults:

The dosage and duration of therapy shall be established depending on type and severity of infection and the condition of the patient.

The recommended daily dosage is as follows:

500 mg IV every 8 hours in the treatment of pneumonia, UTI, gynaecological infections such as endometritis, skin and skin structure infections.

1 g IV every 8 hours in the treatment of nosocomial pneumonias, peritonitis, presumed infections in neutropenic patients, septicemia.

In meningitis the recommended dosage is 2 g every 8 hours.

When treating infections known or suspected to be caused by *Pseudomonas aeruginosa*, a dose of at least 1 g every 8 hours in adults (maximum approved dose is 6 g daily given in 3 divided doses) and a dose of at least 20 mg/kg every 8 hours in children (maximum approved dose is 120 mg/kg daily given in 3 divided doses) are recommended.

Regular sensitivity testing is recommended when treating *Pseudomonas aeruginosa* infection.

MERONEM® should be given as an intravenous bolus injection over approximately 5 minutes or by intravenous infusion over approximately 15 to 30 minutes. There is limited safety data available to support the administration of a 2 g bolus dose.

Dosage schedule for adults with impaired renal function

Dosage should be reduced in patients with creatinine clearance less than 51 mL/min, as scheduled below.

Creatinine Clearance (mL/min)	Dose (based on unit doses of 500 mg, 1 g, 2 g)	Frequency
26 – 50	One unit dose	Every 12 hours
10 – 25	One-half unit dose	Every 12 hours
<10	One-half unit dose	Every 24 hours

MERONEM® is cleared by haemodialysis and haemofiltration; if continued treatment with MERONEM® is necessary, the unit dose based on the infection type and severity is recommended at the completion of the haemodialysis procedure to re-institute effective treatment.

There is no experience with peritoneal dialysis.

Dosage in adults with hepatic insufficiency

No dosage adjustment is necessary in patients with hepatic impairment.

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Elderly patients

No dosage adjustment is required for the elderly with normal renal function or creatinine clearance values above 50 mL/min.

Children

For children over 3 months and up to 12 years of age the recommended dose is 10 – 20 mg/kg every 8 hours depending on type and severity of infection, the known or suspected susceptibility of the pathogen(s) and the condition of the patient. In children over 50 kg weight, adult dosage should be used. In meningitis the recommended dose is 40 mg/kg every 8 hours. There is no experience in children with renal impairment.

MERONEM® should be given as an intravenous bolus injection over approximately 5 minutes or by intravenous infusion over approximately 15 to 30 minutes. There is limited safety data available to support the administration of a 40 mg/kg bolus dose.

Constitution and compatibility

MERONEM® to be used for bolus intravenous injection should be constituted with sterile water for injections (10 mL per 500 mg). This provides an approximate available concentration of 50 mg/mL. Constituted solutions are clear or pale yellow.

For intravenous infusion MERONEM® vials may be directly constituted with a compatible infusion fluid (as listed in *Special precautions for storage*) and then further diluted with the compatible infusion fluid, as needed.

Freshly prepared solutions of MERONEM® should be used whenever possible. However, constituted solutions of MERONEM® maintain satisfactory potency at room temperature (15-25°C) or under refrigeration (4°C) as shown in *Special precautions for storage*.

MERONEM® should not be mixed with or physically added to solutions containing other drugs. Solutions of MERONEM® should not be frozen.

Contraindications

MERONEM® is contraindicated in patients who have demonstrated hypersensitivity to this product.

Special warnings and precautions for use

Patients who have a history of hypersensitivity to carbapenems, penicillins or other β -lactam antibiotics may also be hypersensitive to MERONEM®. As with all β -lactam antibiotics rare hypersensitivity reactions (serious and occasionally fatal) have been reported (see *Undesirable effects*).

Severe cutaneous adverse reactions (SCAR), such as Stevens-Johnson syndrome (SJS), toxic epidermal necrolysis (TEN), drug reaction with eosinophilia and systemic symptoms (DRESS), erythema multiforme (EM) and acute generalized exanthematous pustulosis (AGEP) have been reported in patients receiving MERONEM® (see *Undesirable effect*). If signs and symptoms suggestive of these reactions appear, meropenem should be withdrawn immediately and alternative treatment should be considered.

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Rhabdomyolysis has been reported with the use of meropenem. If signs or symptoms of rhabdomyolysis are observed, meropenem should be discontinued and appropriate therapy initiated.

As with other antibiotics, overgrowth of non-susceptible organisms may occur and repeated evaluation of each patient is necessary.

Use in infections caused by methicillin resistant staphylococci is not recommended.

Rarely, pseudomembranous colitis has been reported on MERONEM® as with practically all antibiotics and may vary in severity from slight to life-threatening. Therefore, antibiotics should be prescribed with care for individuals with a history of gastro-intestinal complaints, particularly colitis.

It is important to consider the diagnosis of pseudomembranous colitis in the case of patients who develop diarrhoea in association with the use of MERONEM®. Although studies indicate that a toxin produced by *Clostridium difficile* is one of the main causes of antibiotic-associated colitis, other causes should be considered.

The concomitant use of valproic acid/sodium valproate and MERONEM® is not recommended. MERONEM® may reduce serum valproic acid levels. Subtherapeutic levels may be reached in some patients (see *Interaction with other medicinal products and other forms of interaction*).

Paediatric use

Efficacy and tolerability in infants under 3 months old have not been established; therefore, MERONEM® is not recommended for use below this age.

Use in patients with renal insufficiency: Refer to dosage recommendations for MERONEM®.

Use in patients with liver disease: Patients with pre-existing liver disorders should have liver function monitored during treatment with MERONEM®.

A positive direct or indirect Coombs test may develop.

Interaction with other medicinal products and other forms of interaction

Probenecid competes with meropenem for active tubular secretion and thus inhibits the renal excretion, with the effect of increasing the elimination half-life and plasma concentration of meropenem. As the potency and duration of action of MERONEM® dosed without probenecid are adequate, the co-administration of probenecid with MERONEM® is not recommended. The potential effect of MERONEM® on the protein binding of other drugs or metabolism has not been studied. However, the protein binding of MERONEM® is so low that no interactions with other compounds would be expected on the basis of this mechanism.

Decreases in blood levels of valproic acid have been reported when it is co-administered with carbapenem agents resulting in a 60-100% decrease in valproic acid levels in about two days. Due to the rapid onset and the extent of the decrease, co-administration of MERONEM® in patients stabilised on valproic acid is not considered to be manageable and therefore should be avoided (see *Special warnings and precautions for use*).

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MERONEM® has been administered concomitantly with other medications without adverse pharmacological interactions. However, no specific drug interactions studies other than probenecid were conducted.

Pregnancy and lactation

Pregnancy

The safety of MERONEM® in human pregnancy has not been established although animal studies have not shown an adverse effect on the developing foetus. MERONEM® should not be used in pregnancy unless the potential benefit justifies the potential risk to the foetus.

Lactation

Meropenem has been reported to be excreted in human milk. MERONEM® should not be used in breast-feeding women unless the potential benefit justifies the potential risk to the baby.

Effect on ability to drive or operate machinery

No studies on the ability to drive and use machines have been performed. However when driving or operating machines, it should be taken into account that headache, paraesthesia and convulsions have been reported for MERONEM®.

Undesirable effects

MERONEM® is generally well tolerated. Adverse reactions rarely lead to cessation of treatment. Serious adverse reactions are rare.

The following adverse reactions have been identified following clinical studies and post-marketing experience with MERONEM®.

ADRs by SOC and CIOMS frequency category listed in order of decreasing medical seriousness or clinical importance within each frequency category and SOC.

System Organ Class	Common ≥1/100 to <1/10	Uncommon ≥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 infestations		Oral candidiasis, Vaginal candidiasis			
Blood and lymphatic system disorders	Thrombocythemia	Thrombocytopenia, Neutropenia, Leucopenia, Eosinophilia	Agranulocytosis*	Haemolytic anaemia*	
Immune system disorders				Manifestations of anaphylaxis*	Angioedema*
Psychiatric disorders			Delirium*		
Nervous	Headache	Paraesthesia	Convulsion		

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System Organ Class	Common ≥1/100 to <1/10	Uncommon ≥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)
system disorders			s		
Gastrointestinal disorders	Diarrhoea, Vomiting, Nausea				Pseudomembranous colitis*
Hepatobiliary disorders	Alanine aminotransferase increased, Aspartate aminotransferase increased, Blood alkaline phosphatase increased, Blood lactate dehydrogenase increased, Gamma-glutamyltransferase increased	Blood bilirubin increased			
Skin and subcutaneous tissue disorders	Rash, Pruritus	Urticaria		Toxic epidermal necrolysis*, Stevens Johnson syndrome*, Erythema multiforme*	Drug Reaction with Eosinophilia and Systemic Symptoms*, Acute generalised exanthematous pustulosis*
Musculoskeletal and connective tissue disorders					Rhabdomyolysis*
General disorders and administration site conditions	Inflammation, Pain	Thrombophlebitis			

*ADR identified post-marketing.

Description of selected adverse reactions

Kounis Syndrome

Kounis syndrome (acute coronary syndrome associated with an allergic reaction) has been reported with other beta-lactam antibiotics.

Reporting of suspected adverse reactions

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Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via:

Pusat Farmakovigilans/MESO Nasional

Direktorat Pengawasan Keamanan, Mutu, dan Ekspor Impor Obat, Narkotika, Psikotropika, Prekursor dan Zat Adiktif

Badan Pengawas Obat dan Makanan

Jl. Percetakan Negara No. 23, Jakarta Pusat, 10560

Email: pv-center@pom.go.id

Phone: +62-21-4244691 Ext.1079

Website: <https://e-meso.pom.go.id/ADR>

PT Pfizer Indonesia

Email: IDN.AEReporting@pfizer.com

Website: www.pfizersafetyreporting.com

Overdose

Intentional overdosing of MERONEM® is unlikely, although overdosing could occur particularly in patients with renal impairment. Limited post-marketing experience indicates that if adverse events occur following over dosage, they are consistent with the adverse event profile described in *Undesirable effects*, are generally mild in severity and resolve on withdrawal or dose reduction. Symptomatic treatments should be considered.

In normal individuals rapid renal elimination will occur. Haemodialysis will remove meropenem and its metabolite.

Pharmacological properties

Pharmacodynamic properties

Meropenem is a carbapenem antibiotic for parenteral use, that is relatively stable to human dehydropeptidase-1 (DHP-1). It is structurally similar to imipenem.

Meropenem exerts its bactericidal action by interfering with vital bacterial cell wall synthesis. The ease with which it penetrates bacterial cell walls, its high level of stability to all serine β -lactamases and its marked affinity for the Penicillin Binding Proteins (PBPs) explain the potent bactericidal action of meropenem against a broad spectrum of aerobic and anaerobic bacteria. The bactericidal concentrations are generally within one doubling dilution of the minimum inhibitory concentrations (MICs).

Meropenem is stable in susceptibility tests and these tests can be performed using normal routine methods. *In vitro* tests show that meropenem acts synergistically with various antibiotics. It has been demonstrated both *in vitro* and *in vivo* that meropenem has a post-antibiotic effect against Gram-positive and Gram-negative organisms.

Mechanisms of resistance

Bacterial resistance to meropenem may result from one or more factors: (1) decreased permeability of the outer membrane of Gram-negative bacteria (due to diminished production of porins) (2) reduced affinity

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of the target PBP (3) increased expression of efflux pump components, and (4) production of β -lactamases that can hydrolyse carbapenems.

Localised clusters of infections due to carbapenem-resistant bacteria have been reported in some regions.

The susceptibility to meropenem of a given clinical isolate should be determined by standard methods. Interpretations of test results should be made in accordance with local infectious diseases and clinical microbiology guidelines.

The antibacterial spectrum of meropenem includes the following species, based on clinical experience and therapeutic guidelines.

Commonly susceptible species: Gram-positive aerobes

Enterococcus faecalis (note that *E. faecalis* can naturally display intermediate susceptibility), *Staphylococcus aureus* (methicillin-susceptible strains only: methicillin-resistant staphylococci including MRSA are resistant to meropenem), *Staphylococcus* species including *Staphylococcus epidermidis* (methicillin-susceptible strains only: methicillin-resistant staphylococci including MRSE are resistant to meropenem), *Streptococcus agalactiae* (Group B streptococcus), *Streptococcus milleri* group (*S. anginosus*, *S. constellatus*, and *S. intermedius*), *Streptococcus pneumoniae*, *Streptococcus pyogenes* (Group A streptococcus)

Commonly susceptible species: Gram-negative aerobes

Citrobacter freundii, *Citrobacter koseri*, *Enterobacter aerogenes*, *Enterobacter cloacae*, *Escherichia coli*, *Haemophilus influenzae*, *Klebsiella oxytoca*, *Klebsiella pneumoniae*, *Morganella morganii*, *Neisseria meningitidis*, *Proteus mirabilis*, *Proteus vulgaris*, *Serratia marcescens*

Commonly susceptible species: Gram-positive anaerobes

Clostridium perfringens, *Peptoniphilus asaccharolyticus*, *Peptostreptococcus* species (including *P. micros*, *P. anaerobius*, *P. magnus*)

Commonly susceptible species: Gram-negative anaerobes

Bacteroides caccae, *Bacteroides fragilis*, *Prevotella bivia*, *Prevotella disiens*

Species for which acquired resistance may be a problem: Gram-positive aerobes

Enterococcus faecium (*E. faecium* can naturally display intermediate susceptibility even without acquired resistance mechanisms).

Species for which acquired resistance may be a problem: Gram-negative aerobes

Acinetobacter species, *Burkholderia cepacia*, *Pseudomonas aeruginosa*

Inherently resistant organisms: Gram-negative aerobes

Stenotrophomonas maltophilia, *Legionella* species

Other inherently resistant organisms

Chlamydophila pneumoniae, *Chlamydophila psittaci*, *Coxiella burnetii*, *Mycoplasma pneumoniae*

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The published medical microbiology literature describes *in vitro* meropenem-susceptibilities of many other bacterial species. However the clinical significance of such in-vitro findings is uncertain. Advice on the clinical significance of *in vitro* findings should be obtained from local infectious diseases and clinical microbiology experts and local professional guidelines.

Meropenem is active *in vitro* against many strains resistant to other β -lactam antibiotics. This is explained in part by enhanced stability to β -lactamases. Activity *in vitro* against strains resistant to unrelated classes of antibiotics such as aminoglycosides or quinolones is common.

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.

Stenotrophomonas maltophilia, *Enterococcus faecium* and methicillin-resistant staphylococci have been found to be resistant to meropenem.

Pharmacokinetic properties

In healthy subjects the mean plasma half-life is approximately 1 hour; the mean volume of distribution is approximately 0.25 l/kg and the mean clearance is 239 mL/min at 500 mg falling to 205 mL/min at 2 g. Doses of 500, 1000 and 2000 mg doses infused over 30 minutes give mean $C_{\max}$ values of approximately 23, 49 and 115 $\mu\text{g/mL}$ respectively, corresponding AUC values were 39.3, 62.3 and 153 $\mu\text{g.h/mL}$. After infusion over 5 minutes $C_{\max}$ values are 52 and 112 $\mu\text{g/mL}$ after 500 and 1000 mg doses respectively. When multiple doses are administered 8-hourly to subjects with normal renal function, accumulation of meropenem does not occur.

A study of 12 patients administered meropenem 1000 mg 8 hourly post-surgically for intra-abdominal infections showed a comparable $C_{\max}$ and half-life to normal subjects but a greater volume of distribution 27 l.

Distribution

The average plasma protein binding of meropenem was approximately 2% and was independent of concentration. Meropenem has been shown to penetrate well into several body fluids and tissues: including lung, bronchial secretions, bile, cerebrospinal fluid, gynaecological tissues, skin, fascia, muscle, and peritoneal exudates.

Metabolism

Meropenem is metabolised by hydrolysis of the β -lactam ring generating a microbiologically inactive metabolite. *In vitro* meropenem shows reduced susceptibility to hydrolysis by human dehydropeptidase-I (DHP-I) compared to imipenem and there is no requirement to co-administer a DHP-I inhibitor.

Elimination

Meropenem is primarily excreted unchanged by the kidneys; approximately 70% (50 – 75%) of the dose is excreted unchanged within 12 hours. A further 28% is recovered as the microbiologically inactive metabolite. Faecal elimination represents only approximately 2% of the dose. The measured renal

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clearance and the effect of probenecid show that meropenem undergoes both filtration and tubular secretion.

Renal insufficiency

Pharmacokinetic studies in patients with renal insufficiency have shown the plasma clearance of meropenem correlates with creatinine clearance. Dosage adjustments are necessary in subjects with renal impairment.

Hepatic insufficiency

A study in patients with alcoholic cirrhosis has shown no effect of liver disease on the pharmacokinetics of meropenem after repeated doses.

Paediatrics

Studies in children have shown that the pharmacokinetics of MERONEM® IV in children is similar to those in adults. The elimination half-life for meropenem was approximately 1.5 to 2.3 hours in children under the age of 2 years and the pharmacokinetics is linear over the dose range of 10 to 40 mg/kg.

Elderly

Pharmacokinetic studies in the elderly subjects (65-80 years) have shown a reduction in plasma clearance which correlated with age-associated reduction in creatinine clearance and a smaller reduction in non-renal clearance. No dose adjustment is required in elderly patients with normal renal function or creatinine clearance values above 50 mL/min (see *Posology and method of administration*).

Preclinical safety data

Animal studies indicate that meropenem is well tolerated by the kidney. In animal studies meropenem has shown nephrotoxic effects, only at high dose levels (500 mg/kg).

Meropenem is generally well tolerated by the CNS. Effects were seen only at very high doses of 2000 mg/kg and above.

The IV LD₅₀ of meropenem in rodents is greater than 2000 mg/kg. In repeat dose studies of up to 6 months duration only minor effects were seen including a small decrease in red cell parameters and an increase in liver weight in dogs treated with doses of 500 mg/kg.

There was no evidence of mutagenic potential in the 5 tests conducted and no evidence of reproductive and teratogenic toxicity in studies at the highest possible doses in rats and monkeys; the no effect dose level of a (small) reduction in F₁ body weight in rats was 120 mg/kg.

There was no evidence of increased sensitivity to meropenem in juveniles compared to adult animals. The intravenous formulation was well tolerated in animal studies.

The sole metabolite of meropenem had a similar profile of toxicity in animal studies.

Pharmaceutical particulars

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List of excipients

Anhydrous sodium carbonate.

Incompatibilities

MERONEM® is compatible with the infusion fluids listed in *Special precautions for storage*.
MERONEM® should not be mixed with or physically added to solutions containing other drugs.

Special precautions for storage

Do not store above 30°C.
Do not freeze.

A solution for bolus injection is prepared by dissolving the drug product MERONEM® in water for injection to a final concentration of 50 mg/mL. Chemical and physical in-use stability for a prepared solution for bolus injection has been demonstrated for 3 hours at up to 25°C or 16 hours under refrigerated conditions (2-8°C).

A solution for infusion is prepared by dissolving the drug product MERONEM® in either 0.9% sodium chloride solution for infusion or 5% glucose (dextrose) solution for infusion to a final concentration of 1 to 20 mg/mL. Chemical and physical in-use stability for a prepared solution for infusion using 0.9% sodium chloride solution has been demonstrated for 3 hours at up to 25°C or 15 hours under refrigerated conditions (2-8°C). Constituted solutions of MERONEM® in 5% glucose (dextrose) solution should be used immediately.

The constituted solutions should not be frozen.

From a microbiological point of view, unless the method of opening/constitution/dilution precludes the risk of microbiological contamination, the product should be used immediately.

If not used immediately, in-use storage times and conditions are the responsibility of the user.

Instructions for use/handling

Refer to *Posology and method of administration* and *Special precautions for storage*. Shake constituted solution before use.

All vials are for single use only.

Standard aseptic technique should be employed during constitution and administration.

Supply

MERONEM® (500 mg): vial in boxes of 10 (Reg. No.: DKI1814500644A1)
MERONEM® (1.0 g): vial in boxes of 10 (Reg. No.: DKI1814500644B1)

HARUS DENGAN RESEP DOKTER

Manufactured by:

ACS Dobfar SpA, Tribiano, Milano, Italy

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Packed and released by:

Zambon Switzerland Ltd., Cadempino, Switzerland

Imported by:

PT. Pfizer Indonesia
Jakarta, Indonesia

Leaflet kemasan: Informasi bagi pengguna

MERONEM® 500 mg, 1 g serbuk untuk larutan infus Meropenem

Baca semua bagian leaflet ini dengan cermat sebelum mulai menggunakan obat ini karena berisi informasi penting bagi Anda.

- Simpan leaflet ini. Anda mungkin perlu membacanya kembali.
- Jika Anda memiliki pertanyaan lebih lanjut, tanyakan kepada dokter, apoteker, atau perawat Anda.
- Obat ini telah diresepkan hanya untuk Anda. Jangan berikan kepada orang lain. Obat ini dapat membahayakan mereka, sekali pun tanda-tanda penyakit mereka sama dengan Anda.
- Jika Anda mengalami efek samping apa pun, konsultasikan dengan dokter, apoteker, atau perawat Anda. Termasuk setiap kemungkinan efek samping yang tidak tercantum dalam leaflet ini. Lihat bagian 13.

Isi leaflet ini:

1. Nama obat
2. Bentuk sediaan
3. Deskripsi obat
4. Apa kandungan obat ini?
5. Kekuatan obat
6. Apa kegunaan obat ini?
7. Berapa banyak dan seberapa sering Anda seharusnya menggunakan obat ini? Apa yang harus dilakukan jika ada dosis terlewat?
8. Kapan seharusnya Anda tidak menggunakan obat ini?
9. Apa yang harus dipertimbangkan saat menggunakan obat ini?
10. Apa saja obat lain atau makanan yang harus dihindari selama menggunakan obat ini?
11. Apakah obat ini aman untuk perempuan hamil dan menyusui?
12. Apakah pasien diperbolehkan mengemudi dan mengoperasikan mesin selama menggunakan obat ini?
13. Apakah potensi efek samping yang tidak diinginkan karena menggunakan obat ini?
14. Tanda-tanda dan gejala-gejala overdosis
15. Apa yang harus dilakukan jika Anda menggunakan lebih dari dosis yang dianjurkan?
16. Bagaimana cara menyimpan obat ini?
17. Berapa lama umur simpan obat setelah wadahnya dibuka untuk pertama kali?
18. Petunjuk penggunaan/penanganan
19. Bagaimana cara melarutkan obat ini?
20. Nomor otorisasi pemasaran
21. Nama dan alamat pemohon dan/atau pemilik obat sesuai dengan ketentuan yang berlaku
22. Tanggal revisi PIL
23. Peringatan khusus

1. Nama obat

Meropenem®

2. Bentuk sediaan

Serbuk untuk larutan injeksi atau infus.

3. Deskripsi obat

Merone[®]m merupakan serbuk putih steril yang mengandung meropenem; 500 mg atau 1 g sebagai trihidrat yang dicampur dengan natrium karbonat anhidrat untuk konstitusi.

Pemerian

Serbuk putih sampai kuning muda.

4. Apa kandungan obat ini?

Vial untuk injeksi atau infus intravena	Merone [®] m 500 mg	Merone [®] m 1 g
Bahan aktif:		
Meropenem (sebagai trihidrat)	570 mg	1,14 g
Setara dengan meropenem anhidrat	500 mg	1 g
Eksipien:		
Natrium karbonat anhidrat	104 mg	208 mg

Untuk setiap gram meropenem (potensi anhidrat), vial berisi 90 mg (3,9 mmol) natrium.

5. Kekuatan obat

500 mg, 1 g

6. Apa kegunaan obat ini?

Merone[®]m diindikasikan untuk pengobatan infeksi pada orang dewasa dan anak-anak yang disebabkan oleh bakteri tunggal atau banyak bakteri yang sensitif terhadap meropenem.

- Pneumonia dan pneumonia nosokomial
- Infeksi saluran kemih
- Infeksi intraabdomen
- Infeksi ginekologis, misalnya endometritis
- Infeksi kulit dan struktur kulit
- Meningitis
- Septikemia
- Pengobatan empirik untuk dugaan infeksi pada pasien dewasa yang disertai demam neutropenia, yang digunakan sebagai terapi tunggal atau dalam kombinasi dengan agen antivirus atau antijamur.

Merone[®]m dapat digunakan, dengan atau tanpa agen antimikroba lainnya, dalam tata laksana infeksi polimikroba.

Tidak ada pengalaman pada pasien pediatrik dengan neutropenia atau imunodefisiensi primer atau sekunder.

Diperlukan pertimbangan dalam menggunakan panduan resmi penggunaan agen antibakteri yang sesuai.

7. Berapa banyak dan seberapa sering Anda seharusnya menggunakan obat ini? Apa yang harus dilakukan jika ada dosis terlewat?

Merone[®]m diinjeksikan ke dalam pembuluh vena Anda. Obat ini hanya boleh diberikan oleh dokter atau perawat. Banyak orang yang menerima Merone[®]m di rumah sakit akan

menerimanya melalui drip (saluran intravena). Meronem® dapat diberikan secara langsung ke dalam pembuluh vena atau melalui drip sehingga tidak perlu melakukan injeksi melalui kulit. Obat ini diberikan baik sebagai injeksi lambat selama sekitar 5 menit atau pemberian melalui drip secara lambat selama 15 hingga 30 menit. Dokter akan memutuskan metode terbaik bagi Anda.

Dokter Anda akan memutuskan dosis Meronem® yang Anda butuhkan bergantung pada faktor tertentu seperti jenis infeksi dan usia Anda. Dosis normal adalah 500 mg hingga 1 g yang diinjeksikan setiap 8 jam. Jika Anda menderita meningitis, Anda mungkin memerlukan dosis yang lebih tinggi, sementara dosis yang lebih rendah dapat digunakan pada anak-anak atau jika Anda menderita gangguan ginjal.

Meronem® harus diberikan sebagai serangkaian injeksi dalam beberapa hari. Dokter Anda akan memutuskan berapa hari Anda akan menjalani pengobatan dengan Meronem®.

Jika Anda lupa menggunakan Meronem®

Jika ada injeksi yang terlewat, maka Anda harus sesegera mungkin melakukan injeksi. Namun demikian, jika sudah mendekati waktu injeksi Anda berikutnya, maka abaikan injeksi yang terlewat.

Jangan memberikan dosis ganda (dua injeksi secara bersamaan) sebagai pengganti injeksi yang terlupa.

8. Kapan seharusnya Anda tidak menggunakan obat ini?

Jangan menggunakan Meronem® jika Anda alergi terhadap meropenem atau bahan-bahan lain dalam obat ini.

9. Apa yang harus dipertimbangkan saat menggunakan obat ini?

Konsultasikan dengan dokter atau perawat Anda sebelum menggunakan Meronem®:

- jika Anda memiliki riwayat hipersensitivitas terhadap karbapenem, penisilin, atau antibiotik β -laktam lainnya
- jika Anda memiliki riwayat keluhan pencernaan, khususnya kolitis
- jika Anda sedang menjalani pengobatan dengan asam valproat/natrium valproat
- jika Anda memiliki gangguan ginjal atau hati

Konsultasikan dengan dokter atau perawat Anda dengan segera selama Anda menjalani pengobatan dengan Meronem®:

- jika Anda mengalami gangguan kulit, seperti ruam, gatal-gatal, atau kulit melepuh
- jika Anda mengalami diare

Jika Anda mengalami nyeri otot, nyeri tekan, atau kelemahan tanpa alasan yang jelas dan/atau urine berwarna gelap, segera beri tahu dokter Anda. Kondisi tersebut bisa saja menandakan adanya kerusakan otot (disebut rabdomiolisis) yang dapat menyebabkan gangguan ginjal.

Seperti halnya antibiotik lainnya, pertumbuhan berlebihan organisme nonrentan dapat terjadi sehingga evaluasi berulang perlu dilakukan.

Tidak dianjurkan untuk digunakan dalam infeksi yang disebabkan oleh stafilokokus yang resistan terhadap metisilin.

Nama Generik: Meropenem
Nama Dagang: MERONEM®
Tanggal Berlaku CDS: 3 Januari 2024
Menggantikan: 31 Oktober 2018
Disetujui oleh BPOM:

Penggunaan pada anak-anak

Merone^m® tidak dianjurkan untuk digunakan pada bayi berusia di bawah 3 bulan.

Penggunaan pada pasien dengan gangguan hati

Pasien yang sebelumnya sudah mengalami gangguan hati harus terus dipantau fungsi hatinya selama pengobatan dengan Merone^m®.

Tes lab

Beri tahu dokter Anda bahwa Anda menggunakan Merone^m® jika Anda akan menjalani suatu tes. Ini karena Anda mungkin mendapatkan hasil positif dalam tes yang disebut “Coombs”. Tes ini digunakan untuk menemukan antibodi yang melawan sel darah merah Anda.

Merone^m® mengandung natrium.

Untuk setiap gram meropenem (potensi anhidrat), vial berisi 90 mg (3,9 mmol) natrium.

10. Apa saja obat lain atau makanan yang harus dihindari selama menggunakan obat ini?

Beri tahu dokter atau perawat Anda jika Anda sedang menggunakan, baru-baru ini menggunakan, atau mungkin akan menggunakan obat-obatan lain.

Beri tahu dokter Anda sebelum Anda menggunakan Merone^m® jika Anda sedang menggunakan obat-obatan berikut ini:

- probenesid
- asam valproat/natrium valproat

11. Apakah obat ini aman untuk perempuan hamil dan menyusui?

Jika Anda hamil atau sedang menyusui, menduga bahwa diri Anda sedang hamil, atau berencana untuk hamil, mintalah saran dokter Anda sebelum menggunakan obat ini.

12. Apakah pasien diperbolehkan mengemudi dan mengoperasikan mesin selama menggunakan obat ini?

Merone^m® dapat membuat Anda merasa sakit kepala, kesemutan, atau gemetar. Ini dapat memengaruhi kemampuan Anda dalam mengemudi, menggunakan alat, atau mesin.

13. Apa saja potensi efek samping yang tidak diinginkan karena menggunakan obat ini?

Seperti semua obat-obatan yang ada, obat ini bisa menimbulkan efek samping, meskipun tidak semua orang mengalaminya. Efek samping berikut dapat terjadi dengan obat ini:

Efek samping serius

Segera beri tahu dokter Anda jika Anda mengamati adanya efek samping serius berikut - Anda mungkin membutuhkan perawatan medis yang mendesak:

- Reaksi alergi berat yang meliputi
 - Ruam, gatal-gatal. atau kaligata yang parah pada kulit.
 - Pembengkakan wajah, bibir, lidah, atau bagian tubuh lainnya.
 - Sesak napas, mengi, atau kesulitan bernapas.
- Reaksi kulit berat yang meliputi

- Reaksi hipersensitivitas serius yang mengakibatkan demam, ruam kulit, dan perubahan hasil tes darah untuk memeriksa fungsi hati (peningkatan kadar enzim hati) dan peningkatan sejenis sel darah putih (eosinofilia) dan pembesaran kelenjar getah bening. Hal ini bisa jadi menunjukkan tanda-tanda gangguan sensitivitas multiorgan yang dikenal dengan istilah sindrom DRESS.
- Ruam bersisik merah yang parah, benjolan pada kulit yang berisi nanah, kulit melepuh atau mengelupas, yang dapat dikaitkan dengan demam tinggi dan nyeri sendi.
- Ruam kulit parah yang dapat terlihat sebagai bercak-bercak melingkar kemerahan pada badan seringkali dengan bagian tengah yang melepuh, pengelupasan kulit, tukak pada mulut, tenggorok, hidung, alat kelamin, dan mata dan dapat didahului dengan demam dan gejala menyerupai flu (sindrom Stevens-Johnson) atau dalam bentuk yang lebih berat (nekrolisis epidermal toksik).

Kerusakan otot

- Nyeri otot, nyeri tekan, atau kelemahan tanpa alasan yang jelas, dan/atau urine berwarna gelap.

Jika Anda mengalami tanda-tanda atau gejala ini, segera beri tahu dokter Anda.

Kemungkinan efek samping yang lain:

Beri tahu dokter atau perawat Anda jika teramati adanya efek samping berikut ini:

Umum: (dapat dialami hingga 1 di antara 10 orang)

- Peningkatan kadar platelet dalam darah
- Sakit kepala
- Diare
- Muntah
- Mual
- Peningkatan enzim hati
- Ruam
- Gatal-gatal
- Inflamasi
- Nyeri

Tidak umum: (dapat dialami hingga 1 di antara 100 orang)

- Infeksi jamur, termasuk infeksi pada mulut dan vagina
- Penurunan kadar platelet dalam darah
- Penurunan jumlah sel darah
- Peningkatan sel asidofil
- Abnormalitas indera peraba
- Peningkatan bilirubin darah
- Kaligata
- Inflamasi pembuluh vena

Langka: (dapat dialami hingga 1 di antara 1000 orang)

- Terlalu bersemangat dan tidak dapat berpikir atau berbicara dengan jelas
- Tubuh gemetar yang tidak dapat dikendalikan

- Nyeri dada tiba-tiba, yang mungkin merupakan tanda reaksi alergi yang berpotensi serius yang disebut sindrom Kounis telah dicatat terjadi dengan obat-obatan lain dari jenis yang sama. Jika ini terjadi, segera konsultasikan dengan dokter atau perawat.

Sangat langka: (dapat dialami hingga 1 di antara 10000 orang)

- Jumlah sel darah merah lebih rendah dikarenakan sel darah merah dihancurkan lebih cepat dibandingkan normal
- Reaksi alergi
- Timbulnya ruam parah atau lecet atau terkelupasnya kulit secara mendadak, yang dapat disertai demam tinggi atau nyeri persendian (ini mungkin merupakan tanda-tanda kondisi medis yang lebih serius seperti nekrolisis epidermal toksik, sindrom Stevens-Johnson, eritema multiformis, atau kondisi yang dikenal sebagai DRESS (Drug Reaction with Eosinophilia and Systemic Symptoms/Reaksi Obat dengan Eosinofilia dan Gejala Sistemik)

Tidak diketahui: (frekuensi tidak dapat diestimasi dari data yang tersedia)

- Pembengkakan pembuluh darah
- Inflamasi usus disertai diare

Beri tahu dokter atau perawat Anda jika teramati adanya efek samping yang tercantum di atas.

Melaporkan efek samping

Jika Anda mengalami efek samping apa pun, konsultasikan dengan dokter, apoteker, atau perawat Anda. Termasuk setiap kemungkinan efek samping yang tidak tercantum dalam leaflet ini. Dengan melaporkan efek samping, Anda bisa membantu memberikan informasi lebih lanjut mengenai keamanan obat ini.

Untuk melaporkan efek samping, hubungi www.pfizersafetyreporting.com atau email di IDN.AEReporting@pfizer.com.

14. Tanda-tanda dan gejala-gejala overdosis

Karena Meronem® diberikan kepada Anda di bawah pengawasan dokter, maka sangat kecil kemungkinan Anda akan mengalami kelebihan dosis. Namun demikian, jika Anda mengalami efek samping serius setelah diberi Meronem®, beri tahu dokter Anda atau datang ke Unit Gawat Darurat di rumah sakit terdekat. Anda mungkin memerlukan penanganan medis segera.

15. Apa yang harus dilakukan jika Anda menggunakan lebih dari dosis yang dianjurkan?

Meronem® akan diberikan kepada Anda oleh dokter atau perawat, jadi kecil kemungkinan untuk diberikan dengan dosis yang salah.

Namun, jika Anda mengalami efek samping atau beranggapan telah diberi terlalu banyak Meronem®, segera beri tahu dokter atau perawat Anda.

16. Bagaimana cara menyimpan obat ini?

Jauhkan obat ini dari pandangan dan jangkauan anak-anak.

Jangan gunakan obat ini setelah melewati tanggal kedaluwarsa yang tertera pada wadahnya.

Jangan menyimpannya di atas suhu 30 °C.

Jangan dibekukan.

Jangan buang obat melalui saluran pembuangan air atau bersama sampah rumah tangga. Tanyakan kepada apoteker cara membuang obat yang sudah tidak dibutuhkan. Langkah-langkah ini akan membantu melindungi lingkungan.

17. Berapa lama umur simpan obat setelah wadahnya dibuka untuk pertama kali?
Jangan menyimpannya di atas suhu 30 °C.

Jangan dibekukan.

Setelah rekonstitusi:

Pemberian injeksi bolus intravena

Larutan untuk injeksi bolus disiapkan dengan melarutkan produk obat Meronem® dalam air untuk injeksi hingga konsentrasi akhir 50 mg/ml. Stabilitas kimia dan fisika selama penggunaan untuk larutan injeksi bolus yang disiapkan telah ditunjukkan selama 3 jam pada suhu maksimal 25 °C atau selama 16 jam jika disimpan di lemari pendingin (2–8 °C).

Pemberian infus intravena

Larutan untuk infus yang disiapkan dengan melarutkan produk obat Meronem® baik dalam larutan natrium klorida 0,9% untuk infus atau larutan glukosa (dekstrosa) 5% untuk infus hingga konsentrasi akhir 1 hingga 20 mg/ml. Stabilitas kimia dan fisika selama penggunaan untuk infus menggunakan larutan natrium klorida 0,9% yang disiapkan telah ditunjukkan selama 3 jam pada suhu maksimal 25 °C atau selama 15 jam jika disimpan di lemari pendingin (2–8 °C). Larutan Meronem® yang dikonsstitusi dalam larutan glukosa (dekstrosa) 5% harus digunakan secepatnya.

Larutan yang dikonsstitusi tidak boleh dibekukan.

Dari sudut pandang mikrobiologi, kecuali jika metode dalam pembukaan/konsstitusi/pengenceran menghindarkan risiko kontaminasi mikrobiologis, produk harus digunakan secepatnya.

18. Petunjuk penggunaan/penanganan

Kocok larutan yang dikonsstitusi sebelum digunakan.

Semua vial hanya untuk sekali pakai.

Teknik aseptik standar harus diterapkan saat konsstitusi dan pemberian.

19. Bagaimana cara melarutkan obat ini?

Meronem® yang akan digunakan untuk injeksi intravena harus dikonsstitusi dengan air steril untuk injeksi (10 ml per 500 mg). Cara ini akan menghasilkan konsentrasi sekitar 50 mg/ml. Larutan yang dikonsstitusi akan terlihat bening atau kuning pucat.

Untuk infus intravena, vial Meronem® dapat dikonsstitusi langsung dengan cairan infus yang kompatibel dan diencerkan lebih lanjut dengan cairan infus yang kompatibel, sesuai kebutuhan.

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Nama Dagang: MERONEM®
Tanggal Berlaku CDS: 3 Januari 2024
Menggantikan: 31 Oktober 2018
Disetujui oleh BPOM:

Larutan Meronem® yang baru disiapkan harus segera digunakan bila memungkinkan. Namun demikian larutan Meronem® yang dikonsitusi tetap stabil secara kimia dan fisika selama 3 jam pada suhu maksimal 25°C atau 15 jam jika disimpan dalam lemari pendingin (2°C - 8°C).

Meronem® tidak boleh dicampur dengan atau ditambahkan secara fisik ke larutan yang mengandung obat lain. Larutan Meronem® tidak boleh dibekukan.

20. Nomor otorisasi pemasaran

MERONEM IV (500 mg): vial dalam dus isi 10 (No. Reg.: DKI1814500644A1)

MERONEM IV (1,0 g): vial dalam dus isi 10 (No. Reg.: DKI1814500644B1)

21. Nama dan alamat pemohon dan/atau pemilik obat sesuai dengan ketentuan yang berlaku

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22. Tanggal revisi PIL

06/2025

23. Peringatan khusus

HARUS DENGAN RESEP DOKTER