Alert	May cause hypotension. Caution advised when using loading dose. Reduce infusion rate for infants with renal impairment and prematurity.						
Indication	Inotrope and vasodilator for:						
iliuication	1. Treatment of low cardiac output and as an adjunct to inhaled nitric oxide in neonates with						
	persistent pulmonary hypertension of the neonate ¹ .						
	2. Prevention of low cardiac output syndrome (LCOS) post cardiac surgery ^{2, 3} .						
	 Prevention of low cardiac output syndrome (LCOS) post cardiac surgery. Treatment of myocardial dysfunction in neonates and children with shock particularly in context of 						
	enteroviral 71 infection ⁴ .	oraniculon in neona	es and emidren w	iti shock particularly in context c	٠.		
Action		P phosphodiestera	se in cardiac and v	vascular muscle.			
Drug type	Selective inhibitor of type 3 cAMP phosphodiesterase in cardiac and vascular muscle. Inotrope and vasodilator.						
Trade name	Primacor, Milrinone GH, Milrinone-Baxter						
Presentation	10mg/10mL (1000 microgram/m						
Dose							
Dose	RECOMMENDED - Regimen with NO loading dose Term infant Preterm infant						
	Maintenance NO loading	0.33 – 0.75 micro	gram/kg/minute	0.2 microgram/kg/minute			
	dose	0.55 0.75 micro	gram, kg/mmate	0.2 microgram, kg/minace			
	uose						
	OPTIONAL - Regimen with loading	ng dose					
	Caution: Risk of hypotension with	-					
	,	Term infant		Preterm infant			
	OPTIONAL Loading dose	Loading: 75 micro	ogram/kg over 1	Loading: 45 microgram/kg			
	Followed by maintenance	hour		over 1 hour			
	dose	0.33 – 0.75 micro	gram/kg/minute	0.2 microgram/kg/minute			
Dose adjustment	Renal impairment (including hypoplastic left heart syndrome undergoing surgery) 0.2 -0.33 microgram/kg/minute IV infusion						
Maximum dose	Maximum IV Infusion rate for the		e is 1 microgram/k	g/minute and 0.5			
	microgram/kg/minute for term a	nd preterm infants	respectively – cau	ution as risk of drug accumulation	n		
	over time.						
Total cumulative							
dose							
Route	IV infusion.						
Preparation	Term infant						
	Regimen with NO loading dose						
	Infusion strengt		Prescribed amount				
	1 mL/kg milrinone and make up to 50mL						
	Draw up 1mL/kg (1000 microgram/kg of milrinone) and add sodium chloride 0.9% or glucose 5% to make a final volume of 50mL. Infusing at a rate of 1mL/hour = 0.33 microgram/kg/minute. For term infants – if loading is not given, higher maintenance infusion may be required to reach the						
	steady drug range of 0.5–0.75 microgram/kg/minute.						
	, , , , ,						
	Preterm infant and renal impair	<u>ment</u>					
	Regimen with NO loading dose						
·	Infusion strengt	<u>th</u>		rescribed amount			
1 mL/hour = 0.2 microgram/kg/minute 0.6 mL/kg milrinone and make up to				one and make up to 50mL			
	Draw up 0.6mL/kg (600 microgram/kg of milrinone) and add sodium chloride 0.9% or glucose 5% to						
	make a final volume of 50mL. Inf	using 1mL/hour = 0	0.2microgram/kg/	minute.			
	For preterm infants – if loading dose is not given, titrate the maximal infusion rate to 0.5						
	microgram/kg/minute if required. Avoid prolonged infusion > 0.2 microgram/kg/minute in very preterm						
	infants.						

	Term infant
	OPTIONAL Regimen with loading dose
	Give a loading dose of 3.75 mL (75 microgram/kg) over 1 hour (Note: risk of hypotension with loading
	dose).
	Preterm infant
	OPTIONAL Regimen with loading dose
	Give a loading dose of 3.75 mL (45 microgram/kg) over 1 hour (Note: risk of hypotension with loading
	dose).
Administration	Continuous IV infusion preferably via central line. Change solution every 24 hours.
	Adjust infusion rate based on haemodynamic and clinical response.
Monitoring	For Loading dose: IV infusion over ONE hour Heart rate, ECG and blood pressure
Monitoring	Urine output and peripheral perfusion frequently.
	Fluid and electrolytes.
	Liver function.
	Platelets
Contraindications	Severe obstructive aortic or pulmonary valvular disease or hypertrophic subaortic stenosis.
	Hypersensitivity to milrinone, other 3,4'-bipyridines (inamrinone) or any other ingredient of the
	formulation.
Precautions	Ensure adequate circulating blood volume prior to commencement.
	Loading dose: Considered optional depending on clinical circumstances. May cause hypotension.
	Monitor BP and heart rate closely and ensure adequate volume replacement.
	Prematurity: Long half-life reported (10 hours) in very preterm infants. ⁵ Avoid prolonged higher rate
	infusion ≥0.2 microgram/kg/minute.
	Renal impairment: Significantly increases half-life of milrinone. A reduction in the infusion rate in
	patients with renal impairment to prevent drug accumulation is advised.
	Patient recovery: Improvement in cardiac output with resultant diuresis may necessitate a reduction in
	the dose of diuretic. Potassium loss due to excessive diuresis may predispose digitalised patients to
	arrhythmias.
Drug interactions	None known.
Adverse	Ventricular arrhythmias in cardiac patients.
reactions	Patent ductus arteriosus.
6 111 1111	May cause hypotension.
Compatibility	Fluids: Glucose 5%, sodium chloride 0.9%.
	PN at Y site: compatible with 2 in 1 solution (Amino acid-glucose-trace element mixture)
	Y-site: Aciclovir, adrenaline (epinephrine) hydrochloride, amikacin, amiodarone, amphotericin B
	liposome, ampicillin, anidulafungin, atenolol, atracurium, azithromycin, aztreonam, bivalirudin, caffeine
	citrate, calcium chloride, calcium gluconate, caspofungin, cefazolin, cefepime, cefiderocol, cefotaxime, cefotetan, cefoxitin, ceftazidime, ceftizoxime, ceftriaxone, cefuroxime, ciprofloxacin, cisatracurium,
	clindamycin phosphate, cloxacillin, dexamethasone sodium phosphate, dexmedetomidine, digoxin,
	dobutamine, dopamine, doripenem, doxycycline, enalaprilat, epinephrine, erythromycin lactobionate,
	fentanyl, fluconazole, fluorouracil, Fosfomycin, ganciclovir, gentamicin sulfate, glyceryl trinitrate,
	glycopyrrolate, heparin, hydralazine, hydrocortisone sodium succinate, insulin (short-acting), ketamine,
	labetalol, linezolid, lorazepam, magnesium sulfate, meropenem, methadone, methylprednisolone
	sodium succinate, metoprolol, metronidazole, midazolam, morphine sulfate, naloxone, nicardipine,
	nitroglycerin, noradrenaline (norepinephrine), octreotide, pamidronate, pancuronium, pentobarbital,
	phenobarbital, piperacillin/tazobactam, potassium acetate, potassium chloride, propofol, propranolol,
	ranitidine, remifentanil, rocuronium, sildenafil, sodium acetate, sodium bicarbonate, sodium
	nitroprusside, succinylcholine, sulfamethoxazole/trimethoprim, tacrolimus, ticarcillin,
	ticarcillin/clavulanate, tobramycin, vancomycin, vasopressin, vecuronium, verapamil, voriconazole,
	zidovudine.
Incompatibility	Fluids: No information.
	Y-site: Alprostadil, Amphotericin B, Amphotericin B lipid complex, esmolol, furosemide (frusemide),
	lide seine andersetuen mentenuerale abouteir
	lidocaine, ondansetron, pantoprazole, phenytoin

	Milrinone GH: If storage is necessary, diluted solution may be stored at 2-8°C and use within 24 hours.
	Milrinone-Baxter: Diluted solution should be used immediately or as soon as practical to reduce
	microbiological hazard.
Storage	Primacor and Milrinone Baxter: Store below 30°C. Do not freeze.
	Milrinone GH: Store below 25°C. Do not freeze. Protect from light.
Excipients	Primacore, Milrinone GH, Milrinone-Baxter: Glucose (monohydrate or anhydrous), lactic acid or sodium
	hydroxide (for pH adjustment), and water for injections.
Special	Discard mixtures exhibiting colour change.
comments	
Evidence	Efficacy
	Treatment of pulmonary hypertension in near term infants: Case series report improvements in
	pulmonary and systemic haemodynamics and oxygenation in infants with pulmonary hypertension
	treated with nitric oxide. ^{1, 6, 7} (LOE IV GOR C) Treatment of very pre-term infants: An RCT found no difference in measures of systemic blood flow
	when used preventatively in extremely premature infants. 8 Case series reported improvement in
	oxygenation and a fall in blood pressure in pre-term infants with pulmonary hypertension treated with
	nitric oxide. ⁹ There are insufficient data to determine the efficacy and safety of milrinone in pre-term
	infants with pulmonary hypertension and/or myocardial dysfunction. ¹⁰ (LOE II ⁸ , GOR C)
	Neonates and infants undergoing cardiac surgery: A single RCT found high dose milrinone reduced the
	risk of LCOS post cardiac surgery. ^{2, 3} (LOE II, GOR B) An historical control study reported use of milrinone
	post ductal ligation improved ventilation and reduced inotrope use ¹¹ (LOE IV, GOR C).
	Infants and children with shock associated with myocardial dysfunction: An RCT found milrinone 0.5
	microgram/kg/min reduced mortality in children with enterovirus 71-induced pulmonary oedema and/or
	shock. A loading dose was not used. ⁴ (LOE II, GOR B)
	Safety
	Reports of arrhythmias, tachycardia, hypotension and hypokalaemia, bronchospasm, headaches,
	thrombocytopenia, anaemia and elevated serum liver enzymes. In neonates treated with milrinone,
	hypotension and intraventricular haemorrhage have been observed. ^{2, 6} (LOE IV)
	Phomogophication
	Pharmacokinetics Extremely pre-term infants for prevention of low systemic blood flow: T _½ averaged 10 hours. Milrinone
	loading infusion 0.75 microgram/kg/min for 3 hours followed by maintenance infusion 0.2
	microgram/kg/min achieved target (180–300 nanogram/mL). ⁵ (LOE IV GOR C)
	Term infants with pulmonary hypertension: Half-life (t½) averaged 4 hours. Loading dose 50
	microgram/kg resulted in sub-therapeutic concentrations. Maintenance infusion 0.33–0.99
	microgram/kg/min resulted in concentrations above target range (180–300 nanogram/mL). (LOE IV GOR
	(c)
	Term newborns with hypoplastic left heart undergoing surgery: Neonates received an initial dose of
	either a 100 or 250 microgram/kg of milrinone into the cardiopulmonary bypass circuit. A constant
	infusion of 0.5 microgram/kg/min resulted in drug accumulation during the initial 12 h of drug
	administration. Postoperatively, milrinone clearance was significantly impaired. Initial loading dose of
	100 microgram/kg on cardiopulmonary bypass resulted in plasma concentrations similar to those
	observed in other therapeutic settings. In the postoperative setting of markedly impaired renal function,
	an infusion rate of 0.2 microgram/kg/min should be considered. ¹²
	Paediatric patients with septic shock: T½ averaged 1.47 hours (range, 0.62 to 10.85 hours). Loading dose
	75 microgram/kg and starting infusion rates 0.75–1.0 microgram/kg/min for patients with normal renal function recommended. ¹³
	Prevention of low cardiac output syndrome post cardiac surgery in infants: Loading dose 50
	microgram/kg then infusion 3 microgram/kg/min for 30 minutes and then a maintenance infusion 0.5
	microgram/kg/min, with adjustment for age. 14 (LOE IV GOR C).
Practice points	This option, and man adjustment for about 12 to 14 doll of.
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Welel elices	with persistent pulmonary hypertension of the newborn and suboptimal response to inhaled nitric oxide.
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Newborn use only

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