Morphine 10mg/mL (Parenteral)

2021

## Newborn use only

Alert	S8 - High risk medication- may cause significant patient harm when used in error.				
Indication Analgesia / sedation:					
	1. Pre-medication prior to intubation or other procedure				
	2. During assisted ventilati				
	<ol> <li>Procedures and post-sur</li> </ol>				
	4. Neonatal abstinence syr		v to opioid	withdrawal	
Action	mu-opioid analgesic – stimul				
Drug Type	mu-opioid analgesic.	· · ·	·		
Trade Name	DBL Morphine Sulfate (also contains sodium chloride and hydrochloric acid). Juno Morphine Hydrochloride				
Presentation	10 mg/mL (10,000 microgra				
Dosage	ANALGESIA				
	CONTINUOUS IV IN	FUSION			
	Range: 5–40 microg	ram/kg/hour:			
	Ventilated infants of	or after surgery*	1,2,3]		
	Postnatal age <sup>#</sup>	Starting dose		Range	
	0-7 days	10 microgram,	-	5-40 microgram/kg/hour	
	8-30 days	15 microgram,	′kg/hour	5-40 microgram/kg/hour	
	31-90 days	20 microgram,	′kg/hour	5-40 microgram/kg/hour	
	*Infants after cardio	ovascular surgery	may need	lower starting dose and titrated to clinical	
	response.[2]				
Maximum Daily Dose Route Preparation	Doses up to 100 microgram/ with an increase in the durat IV	dose (up to 200 m NDROME –INITIA our titrated to Ne kg/hour have bee ion of mechanica r IV infusion	L TREATM conatal Abs en used in l ventilatic	ENT stinence Syndrome scores. newborns; however this was associated	
	Prescribed amo			Infusion rate	
			1 ml /h o		
	1 mg/kg morphine and mal	to 50 mL	1 mL/not	ur = 20 microgram/kg/hour	
	Step 1:       Draw up 1 mL (10mg morphine in 1mL) and add 9 mL sodium chloride 0.9% to make a volume of 10 mL with a concentration of 1000 microgram/mL.         Step 2:       From the above solution, draw up 1 mL/kg (1000 microgram/kg) and further dilute with glucose 5% or glucose 10% or sodium chloride 0.9% to make a final volume of 50 mL with a concentration of 1 mL/hour = 20 microgram/kg/hour.         IV bolus dose from single strength solution: 2.5 mL =50 microgram/kg.				
	Prescribed amo			Infusion rate	
	2 mg/kg morphine and mal	ke up to 50 mL	1 mL/hou	ır = 40 microgram/kg/hour	
	Step 1: Draw up 1 mL (10mg morphine in 1mL) and add 9 mL sodium chloride 0.9% to make a volume of 10 mL with a concentration of 1000 microgram/mL.				

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	Step 2: From the above solution draw up	2 mL/kg (2000 microgram/kg) and further dilute with		
		pride 0.9% to make a final volume of 50 mL with a		
	concentration of <b>1 mL/hour = 40 microgram</b> /			
	IV bolus dose from double strength solution	-		
	1-STEP DILUTION for IV infusion	(consider for weight 2 kg and over)		
	IV Infusion: SINGLE STRENGTH			
	Prescribed amount	Infusion rate		
	1 mg/kg morphine and make up to 50 mL	1 mL/hour = 20 microgram/kg/hour		
		and add glucose 5% or glucose 10% or sodium chloride concentration of 1 mL/hour = 20 microgram/kg/hour. on: 2.5 mL = 50 microgram/kg.		
	IV Infusion: DOUBLE STRENGTH			
	Prescribed amount	Infusion rate		
	2 mg/kg morphine and make up to 50 mL	1 mL/hour = 40 microgram/kg/hour		
	Draw up 0.2 mL/kg (10mg morphine in 1mL) and add glucose 5% or glucose 10% or sodium chloride 0.9% to make a final volume of 50 mL with a concentration of 1 mL/hour = 40 microgram/kg/hour. For IV bolus dose from double strength solution: 1.25 mL = 50 microgram/kg.			
	IV BOLUS and PRE-MEDICATION			
	Draw up 0.5 mL (5 mg morphine) and add 9.5 mL sodium chloride 0.9% to make a final volume of 10 mL with a concentration of 500 microgram/mL.			
Administration	CONTINUOUS IV INFUSION: Via syringe drive			
	<b>IV BOLUS</b> : Administer over 5 minutes. Flush v injection. Rapid IV administration may increa	vith 1 mL sodium chloride 0.9% before and after se adverse effects.		
	<b>PRE-MEDICATION FOR INTUBATION:</b> As about of action; however for maximum effect wait a	ve for IV bolus. Wait a minimum of 5 minutes for onse 15 minutes after giving the dose.		
Monitoring	All patients should have cardiorespiratory monitoring and be carefully observed, particularly if they			
	are breathing spontaneously. Respiratory depression/apnoea can be reversed with naloxone. Naloxone is contraindicated in opioid dependent infants.			
	Observe for urinary retention, abdominal distension or delay in passage of stool. Withdraw slowly following prolonged use.			
Contraindications	Hypersensitivity to morphine or any excipient	ts.		
Precautions	Potentially toxic serum concentrations of morphine may occur in infants with hypoxic ischaemic			
		and infusion rates >10 microgram/kg per hour. [3] Use		
	with caution in patients with hypersensitivity reactions to other opioids.			
	Hypotension and bradycardia. Respiratory depression.			
	Transient hypertonia. Convulsions. Ileus and delayed gastric emptying time. Urinary retention. Renal or hepatic impairment.			
	Tolerance may develop after prolonged use –			
Drug Interactions		potentiates effects of opioids, increasing risk of		
<b>C</b>	respiratory depression, profound sedation or			
Adverse Reactions		ry depression (levels above 20 ng/mL); decreased		
Adverse Reactions Compatibility	gastrointestinal motility, hypotension at high Compatibility is likely to be similar for morph	er doses, and urinary retention [4].		

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	Morphine hydrochloride – glucose 5%, sodium chloride 0.9% Morphine sulfate – glucose 2.5%, 5% and 10%, glucose in sodium chloride solutions,	
	Hartmann's, sodium chloride 0.45% and 0.9%	
	Y-site :	
	Morphine hydrochloride – some information is available. Consult the pharmacist,	
	pharmacy department or medicines information service for more advice.	
	Morphine sulfate – adrenaline hydrochloride, amifostine, amikacin, amiodarone,	
	ampicillin, anidulafungin, atracurium, atropine, aztreonam, bivalirudin, caspofungin,	
	cefazolin, cefotaxime, cefoxitin, ceftazidime, ceftriaxone, cisatracurium, clindamycin,	
	dexamethasone, digoxin, dopamine, eptifibatide, erythromycin, esmolol, filgrastim,	
	fluconazole, foscarnet, gentamicin, granisetron, haloperidol lactate (in glucose), heparin	
	sodium, hyoscine hydrobromide, insulin (short-acting), ketorolac, labetalol, lignocaine,	
	linezolid, magnesium sulfate, methylprednisolone sodium succinate, metoclopramide,	
	metoprolol, metronidazole, midazolam, milrinone, noradrenaline, palonosetron,	
	paracetamol, piperacillin-tazobactam (EDTA-free), posaconazole, potassium chloride,	
	remifentanil, sodium nitroprusside, tacrolimus, tigecycline, tirofiban, tobramycin,	
	trimethoprim-sulfamethoxazole, vancomycin, vecuronium, zidovudine.	
Incompatibility	<b>Fluids:</b> Morphine may precipitate out of solution when the final pH is greater than 6.4.	
	Drugs :	
	Morphine hydrochloride – esomeprazole	
	<b>Morphine sulfate</b> – Aminophylline, azathioprine, azithromycin, flucloxacillin, folic acid, ganciclovir, indometacin, pentamidine, pethidine, promethazine, sodium nitrite, thiopental	
	sodium.	
Stability	Diluted solution for continuous IV infusion is stable for 48 hours.	
Storage	Ampoule: Store below 25°C. Protect from light.	
	Discard remainder after use (in line with schedule 8 drug legislation).	
	Store in Dangerous Drug (DD) safe and record use in DD register.	
Special Comments	Prolonged use (> 5–7 days) may be associated with dependence.	
	Morphine hydrochloride and sulfate contain approximately equivalent amounts of morphine base	
	per milligram.	
Evidence	Efficacy:	
	Premedication: Morphine 0.2 mg/kg bolus did not reduce the occurrence of severe hypoxia with	
	bradycardia during intubation, in comparison with placebo.[5] [LOE II] Morphine 0.1 mg/kg –	
	atropine 10 microgram/kg and suxamethonium 1 mg/kg premedication reduced the total time and	
	number of attempts taken to achieve successful nasotracheal intubation of neonates compared to	
	awake intubation;[6] [LOE II] Morphine 0.1 mg/kg – atropine 10 microgram/kg and suxamethonium	
	2 mg/kg was less effective than propofol with longer time to intubation, increased oxygen desaturations and nasal trauma and increased time to recovery [7]. (LOE II] No difference in time,	
	number of attempts and duration of intubation has been reported in trials comparing morphine-	
	midazolam versus remifentanil with or without midazolam combination [8, 9]. (LOE II) Conclusion:	
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	selectively, when indicated by clinical judgment and evaluation of pain indicators. If sedation is
	required, morphine is safer than midazolam [11]. (LOE I GOR B)
	Analgaesia: Recommended procedural analgesic doses for neonates are: Intermittent Dose -
	Morphine sulfate 0.05-0.1 mg/kg intravenously; <i>Infusion Dose -</i> 0.01-0.03 mg/kg per hour. It is
	advised that neonatal intensive care units use only 1 opioid analgesic agent to ensure familiarity
	with its use. The opioid doses are only applicable for opioid-naive patients. All patients should be
	monitored and carefully observed, particularly if they are breathing spontaneously. Consider slow
	intravenous opioid infusion (morphine sulfate or fentanyl citrate) for: central venous line
	placement, endotracheal intubation and suction; chest tube insertion and for ventilated infants.
	[Consensus statement for the International Evidence-Based Group for Neonatal Pain] [4].
	Postoperative pain relief: Continuous and intermittent morphine infusions have been trialled in
	postoperative patients. A continuous morphine 10 microgram/kg per hour or intermittent morphine
	30 microgram/kg per 3 hours were equally effective and safe in neonates. (LOE II] A morphine
	continuous infusion to a targeted morphine concentration of 20 ng/ml provided more reliable
	analgesia than an intermittent bolus doses as needed. The average infusion rate was 20.6 ± 8.7
	microgram/kg/hour. [16]. [LOE II] Postoperative morphine use can be reduced by paracetamol
	infusion [17]. [LOE II]
	<b>Neonatal abstinence syndrome secondary to opioids:</b> There are no trials of intravenous morphine
	for NAS secondary to opioids although its use has been reported including for seizure control [18,
	19]. [LOE IV] Recommended oral dose for initial treatment of NAS in opioid dependent infants 0.5
	mg/kg/day [20]. Estimated oral morphine bioavailability 48.5% in neonates [21]. (LOE IV GOR C)
	Pharmacodynamics / Pharmacokinetics:
	Effective morphine concentrations in the range of 10–20 ng/L have been reported [1, 22].
	Concentrations above 20 ng/L have been associated with respiratory depression [2]. The mean
	morphine half-life is age related, reported as around 9 hours in ventilated preterm infants [23, 24],
	6 hours in term infants [24, 25] and 2 hours for infants beyond 11 days age [24].
	Pharmacodynamic assessment found median (IQR) average morphine infusion rate for pain relief in
	was 4.4 (4.0-4.8) microgram/kg/hour in postoperative term neonates <10 days versus 14.4 (11.3-
	23.4) microgram/kg/hour in older infants (p < 0.001) [26]. Also in postoperative term infants,
	morphine concentrations suggested neonates <7 days require significantly less morphine
	postoperatively than older neonates. The recommended dosage for continuous morphine infusions
	were 7 microgram/kg/h in full-term neonates; 10 microgram/kg/hour in infants >4 weeks of age
	[27]. (LOE II GOR B)
	Lynn et al estimated morphine infusion rates to achieve a steady-state concentration ≤20 ng/mL for
	non-cardiovascular surgery are: 0-7 days: 10 microgram/kg/hour; 8-30 days: 15 microgram/kg/hour;
	31-90 days: 20 microgram/kg/hour [1]. For infants after cardiovascular surgery clearance was
	reduced with the following modelled rates: 0-7 days: 5 microgram/kg/hour; 8-30 days: 5
	microgram/kg/hour; 31-90 days: 10 microgram/kg/hour [2].[LOE II GOR B]
	More restricted dosing recommendations have been suggested in neonates targeting morphine
	concentrations of ≤10 microgram/L [26, 27].
	Infants with hypoxic ischemic encephalopathy have reduced morphine clearance and elevated
	serum morphine concentrations when morphine infusion rates are based on clinical state.
	Potentially toxic serum concentrations of morphine may occur with moderate hypothermia and
	infusion rates >10 microgram/kg per hour [3].
	Safety
	There is no compelling evidence to support severe long-term harm, but subtler behavioural changes
	have been noted. Morphine use should continue to be based on clinical judgment, carefully
	weighing the benefits of acute interventions against the potential for long-term harm.[28]
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