Dopamine - Standard Concentration

Newborn use only

Alert	Ensure dopamine has a dedicated line. DO NOT BOLUS.
Indication	Hypotension. (1-3)
	May also be used to improve renal perfusion. (4-6)
Action	Catecholamine with alpha and beta adrenergic, dopaminergic and serotoninergic actions
	Haemodynamic effects are dose dependent. (7)
	• Low dose: 1 to 5 microgram/kg/min – increases renal blood flow and glomerular filtration rate. (4)
	• Intermediate dose: 5 to 10 microgram/kg/min – increases cardiac output and blood pressure in addition
	to renal blood flow.
	 High dose: 10 to 20 microgram/kg/min – systemic vasoconstrictor effect outweighs all other effects. (8) Reduces renal blood flow. (7)
Drug turo	Sympathomimetic, Inotropic vasopressor.
Drug type	
Trade name Presentation	Dopamine (DBL) concentrate 200mg/5mL ampoule
Dose	Hypotension*
Dose	5-20 microgram/kg/minute
	Initiate at 5-10 microgram/kg/minute. Titrate dose as per response.
	Doses higher than 10 microgram/kg/minute require caution. Discuss with neonatologist.
	Clinical response is expected within a few minutes after entry of the drug into circulation. *
	If response is suboptimal, dose can be increased every 10-30 minutes until desired response is
	obtained or maximum dose is reached. (9-12)
	Renal perfusion
	1-5 microgram/kg/min.
	*NOTE: The time from the initiation of infusion to the entry of the drug into circulation may influence the
	time it takes to see the clinical effect. This lag time can be reduced by (a) starting temporarily at a higher dose
	by increasing the infusion rate, and/or (b) priming the line as close to the entry point as possible to reduce the dead space – however, care should be taken not to deliver excess volume that may result in tachycardia
	and hypertension."
Dose adjustment	Therapeutic hypothermia: Limited data in neonates to guide dose adjustments.
···· , ··· ·	ECMO: Limited data in neonates to guide dose adjustments.
	Renal impairment: Limited data in neonates to guide dose adjustments.
	Hepatic impairment: Limited data in neonates to guide dose adjustments.
Maximum dose	20 microgram/kg/minute
Total cumulative	
dose	
Route	Continuous IV infusion.
Preparation	Note: Refer to Appendix for tables to assist with concentration selection.
	20mL Syringe
	1 mg/mL infusion (suggested for a dose ≤5 microgram/kg/minute)
	Draw up 0.5 mL (20 mg) of dopamine and add 19.5 mL glucose 5% or sodium chloride 0.9% [#] to make a final
	volume of 20 mL.
	10 microgram/kg/minute = 0.6 mL/kg/hour.
	2 mg/mL infusion (suggested weight <2 kg)
	Draw up 1 mL (40 mg) of dopamine and add 19 mL glucose 5% or sodium chloride 0.9% [#] to make a final
<i>•</i>	volume of 20 mL.
	10 microgram/kg/minute = 0.3 mL/kg/hour.
	5 mg/mL infusion (suggested weight ≥2 kg)
	Draw up 2.5 mL (100 mg) of dopamine and add 17.5 mL glucose 5% [#] to make a final volume of 20 mL.
	10 microgram/kg/minute = 0.12 mL/kg/hour.

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	50mL Syringe
	1 mg/mL infusion (suggested for a dose ≤5 microgram/kg/minute)
	Draw up 1.25 mL (50mg) of dopamine and add 48.75 mL glucose 5% or sodium chloride 0.9% [#] to make a
	final volume of 50 mL.
	10 microgram/kg/minute = 0.6 mL/kg/hour.
	2 mg/mL infusion (suggested weight <2 kg)
	Draw up 2.5 mL (100mg) of dopamine and add 47.5 mL glucose 5% or sodium chloride 0.9% [#] to make a final
	volume of 50 mL.
	10 microgram/kg/minute = 0.3 mL/kg/hour.
	5 mg/mL infusion (suggested weight ≥2 kg)
	Draw up 6.25 mL (250mg) of dopamine and add 43.75 mL glucose 5% [#] to make a final volume of 50 mL.
	10 microgram/kg/minute = 0.12 mL/kg/hour.
	[#] Sodium chloride 0.9% can be used as a diluent, but only to make a maximum concentration of 3.2 mg/mL
	dopamine solution. ²⁴
Administration	Continuous intravenous infusion via a central line. Use with caution via a peripheral line (preferably low dose and for short duration).
Monitoring	Continuous heart rate, ECG and blood pressure
	Assess urine output and peripheral perfusion frequently.
	Observe intravenous site closely for blanching and extravasation.
Contraindications	Arrhythmia, tachyarrhythmia and phaeochromocytoma.
Precautions	Hypovolaemia- Ensure adequate circulating blood volume prior to commencement.
	May increase pulmonary hypertension.
Drug interactions	Glyceryl trinitrate, nitroprusside and calcium channel blockers: May cause hypotension
	Digitalis glycosides: May increase the risk of cardiac arrhythmias.
	Phenytoin: May result in dose dependent, sudden hypotension and bradycardia.
Adverse	Ectopic beats, tachycardia and arrhythmia.
reactions	Systemic and pulmonary hypertension, especially at higher doses.
reactions	Reversible suppression of prolactin and thyrotropin secretion.
	Tissue necrosis at infusion site with extravasation, uraemia, mydriasis
Compatibility	
Compatibility	Fluids: Glucose 5%, glucose 10%, glucose in sodium chloride 0.9%, glucose 5% in Hartmann's, Hartmann's,
	mannitol 20%, sodium chloride 0.9%
	Y-site: Amino acid solutions,* amifostine, amiodarone, anidulafungin, atracurium, aztreonam, bivalirudin,
	caffeine citrate, caspofungin, ceftaroline fosamil, ciprofloxacin, cisatracurium, dexmedetomidine,
	dobutamine, esmolol, ethanol, fluconazole, foscarnet, glyceryl trinitrate, granisetron, haloperidol lactate,
	heparin sodium, hydrocortisone sodium succinate, labetalol, lignocaine, linezolid, methylprednisolone
	sodium succinate, metronidazole, midazolam, milrinone, morphine sulfate, mycophenolate mofetil,
	noradrenaline, pancuronium, pethidine, piperacillin-tazobactam (EDTA-free), potassium chloride,
	ranitidine, remifentanil, sodium nitroprusside, streptokinase, tigecycline, tirofiban, vecuronium, verapamil,
	zidovudine.
	*ANMF medical group consensus: TPN compatibility is complex. There is limited information on
	pharmaceutical compatibility of dopamine with neonatal PN formulations. Please refer to Micromedex IV
	compatibility section for further information.
Incompatibility	Fluids: Sodium bicarbonate and other alkaline solutions.
	Y-site: Aciclovir, alteplase, ampicillin, azathioprine, cephazolin, chloramphenicol, diazoxide, esomeprazole,
	ganciclovir, ibuprofen, indomethacin, insulin (short-acting), phenytoin, sodium bicarbonate, thiopentone.
Stability	Ampoule: Store below 30°C. Protect from light.
	Diluted solution: Stable for 24 hours below 25°C.
Storage	Store below 25°C
	Protect from light.
	Discard remainder after use.
Excipients	sodium metabisulfite

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Special comments	Discard admixtures exhibiting colour change.
Evidence	Efficacy
	Hypotension
	In a random effects meta-analysis of 7 trials (n=286) in preterm infants Dopamine was found to significantly
	increase mean and systolic arterial blood pressure. For the increase in blood pressure, Dopamine was
	associated with a significantly greater overall efficacy than Dobutamine, colloid or hydrocortisone alone.
	Dopamine was also associated with increased cerebral blood flow with a greater increase in hypotensive
	than in normotensive preterm infants. (1-2) However, in these meta-analyses, no differences were found
	among regimens regarding survival and other secondary clinical outcomes. A considerable inter-individual
	variability in blood pressure response has been reported in the included studies. (3) (LOE I, GOR B)
	In systematic review of 28 RCTs and 12 different comparisons of 6 commonly used vasopressors in adult
	patients, Gamper et at found insufficient evidence to recommend any one of the vasopressors over others
	in the assessed doses. The choice of a specific vasopressor may therefore be individualized and left to the
	discretion of the treating physicians. (13) (LOE I, GOR B)
	Dose escalation: Comparative data to guide the dose escalation strategy is very limited. Randomised
	control trials comparing efficacy of inotropes in neonatal patients increased dopamine dose after allowing a
	variable period of 10 -30 minutes for optimal effect. (9-12)
	Septic shock
	In a RCT Baske et al compared Dopamine (10–20 μg/kg/min) or Adrenaline (0.2–0.4 μg/kg/min) as a first- line vasoactive medication in 40 neonates for successful reversal of fluid-refractory septic shock. The mean
	gestational age of participants at birth was 30 weeks and their mean postnatal age at treatment was 6 days.
	Reversal of shock was defined as achievement of systolic and diastolic blood pressure > fifth centile,
	capillary refill time < 3 seconds and a left ventricular output \geq 150 mL/kg/min. The proportion of neonates
	achieving reversal of shock by 45 min, haemodynamic stability anytime during therapy and all-cause
	mortality by 28 days were comparable in the two groups. Moreover, the two groups had comparable
	lactate clearance, duration of vasoactive therapy and incidence of intraventricular haemorrhage,
	bronchopulmonary dysplasia, necrotising enterocolitis and retinopathy of prematurity. In the subgroup of
	extremely low birthweight infants (n=18), Adrenaline was more efficient in achieving hemodynamic stability
	but there were no differences in the other outcomes. (10) A systematic review of two pediatric and one
	neonatal RCT comprising of 220 participants with septic shock also reported comparable efficacy of
	Dopamine and Adrenaline for the treatment of septic shock. (14)
	Good quality data from randomised control trials or prospective studies for comparing Dopamine and
	Noradrenaline for management of septic shock in neonates are lacking. In a retrospective cohort study,
	Nissimov et al investigated the clinical outcomes of extremely preterm neonates who received either
	Dopamine (n=113) or Noradrenaline (n=43) as a first line agent for management of septic shock in two
	different epochs. Dopamine was administered at a dose of 5 -20 mcg/kg/min and noradrenaline at 0.05-0.4
	mcg/kg/min. Infants who received Noradrenaline had a lower episode related mortality (OR 0.55; 95% Cl 0.32-0.92) new neurological injuny (OR 0.32: 95% Cl 0.13-0.82) and subsequent NEC/sensis (OR 0.34, 95%
4	0.33-0.92), new neurological injury (OR 0.32; 95% CI 0.13-0.82) and subsequent NEC/sepsis (OR 0.34, 95% CI 0.18 - 0.65). (15) A meta-analysis of 11 RCTs in adult patients which compared Dopamine and
	Noradrenaline for septic shock showed no statistically significant difference in the mean arterial pressure
	but favourable effect of Noradrenaline on heart rate, cardiac index and urine output. The Noradrenaline
· ·	group had 11% reduction in absolute risk of all-cause mortality at 28 days. (11) Baseline severity of illness
	and development of arrhythmias during treatment were significant predictors of mortality. (16, 17)
	Effect on pulmonary arterial pressure
	In a small cohort of 18 preterm infants with a mean gestational age of 28 weeks and postnatal age 4 days
	Dopamine was used for treatment of hypotension. Transthoracic cardiac ultrasound was used to assess
	pressure gradient through the patent ductus arteriosus (PDA) and estimate mean pulmonary arterial
4	pressure. Authors noted increase in both systemic and pulmonary arterial pressures after a mean
	Dopamine dose of 13 mcg/kg/min was reached. The mean systemic blood pressure increased by 41% and
	the mean pulmonary arterial pressure increased by 43%. The pulmonary to systemic mean arterial pressure
	(PAP/SAP) ratio increased in 50% infants and in 18% infants unidirectional left to right shunt across the PDA
	became bidirectional due to increased PAP/SAP ratio. (9)
	Dopamine to prevent renal dysfunction in indomethacin-treated preterm newborn infants
	Dopamine improved urine output (2.5 vs 1.8 ml/kg/hour) but there was no evidence of effect on serum
	creatinine, incidence of oliguria (urine output < 1ml/kg/hour) or frequency of failure to close the ductus
	arteriosus. (5) (LOE I, GOR B) Moreover, evidence from well-performed clinical studies to support the

Safety

	Limited data suggest higher dose dopamine may reduce cardiac output. (8,18) (LOE II, GOR C) There is insufficient safety data in neonates for use at doses > 20 micrograms/kg/min. In a systematic review, Sassano-Higgins did not find statistically significant difference in adverse neurological outcome between dopamine, dobutamine, adrenaline, colloid or Hydrocortisone administration when used for hypotension. (2) In a secondary analysis of a prospectively enrolled cohort of 61 neonates, Solanki et al reported the effect of Dopamine (n=33) on cerebral autoregulation. The mean birth weight of the subjects was 849 g and the mean gestation was 26 weeks. In this study, significantly less epochs without Dopamine exposure were associated with impaired cerebral autoregulation compared with Dopamine exposure epochs (14.5% vs. 30.7%; p< 0.001). (19) However, presence of hypotension, gestational age at birth and postnatal age independently affect cerebral autoregulation and are important confounders. (20) Pharmacokinetics The cardiovascular and renal effects of dopamine result from its direct action on dopaminergic, serotonergic and alpha/ beta adrenoceptors. Its effects are dose dependent with some overlap in receptor activation as well as inter-patient variability in binding affinities at different doses. In general, at low doses Dopamine receptors are preferentially activated accounting for its renal effects and at doses > 2 micrograms/kg per minute, the beta and alpha adrenoceptors are stimulated mediating its cardiovascular effects. Steady-state plasma Dopamine concentrations and plasma clearance rates were observed within 20 minutes (dose range 1–8 microgram/kg/min). There is a linear correlation between infusion rate and plasma Dopamine concentration and plasma clearance of Dopamine in neonates is reported to be 385 ml/kg/minute with a large inter-individual variation. (22) Dopamine in neonates is heart rate. (21) The median plasma clearance of Dopamine in meantes is reported to be between 2-9 min in different stud
Practice points	
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routine use of low dose Dopamine for improving renal function in critically ill neonates is insufficient. (6)

Dopamine increases heart rate and has a higher propensity to develop cardiac arrythmias. (9, 16,17)

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Appendix	24. Merati	veTM Mi an, USA.	cromedex Available	Compleat: https://doi.org/10.1000/100000000000000000000000000000	//www.mi	patibility cromedex	(electronio	version).	Merative	, Ann Arbo	or,
	Table 1: Inf (suggested					centration	n 1 mg/m l		7		
	Rate (mL/hr)	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	1
	Weight (kg)				Approxi	mate mic	rogram/k	g/minute			
	0.5	3	7	10	13	17	20	23	27	30	33
	1	2	3	5	7	8	10	12	13	15	17
	1.5	1	2	3	4	6	7	8	9	10	11
	2	1	2	3	3	4	5	6	7	8	8
	2.5	1	1	2	3	3	4	5	5	6	7
	3	1	1	2	2	3	3	4	4	5	6
	3.5	<1	1	1	2	2	3	3	4	4	5
	4	<1	1	1	2	2	3	3	3	4	4
	4.5	<1	1	1	1	2	2	3	3	3	4
	5 Table 2: Inf (suggested			1 using dopa	1 amine con	2 centratior	2 n 2 mg/ml	2	3	3	3
	Rate (mL/hr)	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	1
	Weight (kg)	Weight (kg) Approximate microgram/kg/minute									
	0.5	7	13	20	27	33	40	47	53	60	67
	1	3	7	10	13	17	20	23	27	30	33
	1.5	2	4	7	9	11	13	16	18	20	22
	2	2	3	5	7	8	10	12	13	15	17
	2.5	1	3	4	5	7	8	9	11	12	13
	3	1	2	3	4	6	7	8	9	10	11
	3.5	1	2	3	4	5	6	7	8	9	10
	4	1	2	3	3	4	5	6	7	8	8
	4.5	1	1	2	3	4	4	5	6	7	7
	5	1	1	2	3	3	4	5	5	6	7

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Rate 0.1 (mL/hr)	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	
Weight (kg)			Approxi	mate mic	rogram/k	g/minute			
0.5 17	33	50	67	83	100	117	133	150	1
1 8	17	25	33	42	50	58	67	75	00
1.5 6	11	17	22	28	33	39	44	50	1
2 4	8	13	17	21	25	29	33	38	Z
2.5 3	7	10	13	17	20	23	27	30	
3 3	6	8	11	14	17	19	22	25	-
3.5 2	5	7	10	12	14	17	19	21	
4 2	4	6	8	10	13	15	17	19	
4.5 2	4	6	7	9	11	13	15	17	
5 2	3	5	7	8	10	12	13	15	-
	$rogram/kg/min) = \frac{Rate (mL/hr) \times Concentration (microgram/mL)}{Weight (kg) \times 60}$ hr) = $\frac{60 \times Dose (microgram/kg/min) \times Weight (kg)}{Concentration (microgram/mL)}$								

VERSION/NUMBER	DATE	
Original 1.0	26/05/2025	
REVIEW	26/05/2030	

This standard concentration formulary has been developed by the ANMF standard concentration working group. The working group (in alphabetical order): Mohammad Irfan Azeem, Susanah Brew, Cindy Chen, Michelle Jenkins, Kerrie Knox, Rebecca O'Grady

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