Digoxin Newborn use only

Alert	Digoxin has a narrow therapeutic index, check the dose carefully. Lanoxin adult injection is 10 times more concentrated than Lanoxin infant injection. Check product selection carefully. Rapid IV injection may cause hypertension and reduced coronary flow. Lanoxin Paediatric Elixir contains ethanol of approximately 84 mg/mL, equivalent to 10.6% absolute volume. The long-term effects of prolonged exposure to ethanol content from medicines have not been studied.				
Indication	Supraventricular tachycardia [atrioventricular reciprocating tachycardia or atrioventricular nodal re- entrant tachycardia, excluding Wolff-Parkinson-White]. Atrial fibrillation and atrial flutter. Heart failure [add-on treatment in infants with reduced ejection fraction if not otherwise contraindicated].				
Action	Slows heart rate and reduces AV nodal conduction by an increase in vagal tone and a reduction in sympathetic activity. A Na ⁺ /K ⁺ -ATPase inhibitor which increases the force of myocardial contraction by increasing the release and availability of stored intracellular calcium				
Drug type	Cardiac glycoside				
Trade name	Lanovin PG Sigmavin PG La	novin Sign	navin Lanovin Paediatri	Elivir Lanovin Infant 9	olution for Infusion
frade fiame	Lanoxin Solution for Infusion	novin, Sigi			
Presentation	OBAL:	1			
resentation	Lanoxin PG. Sigmaxin PG 62.	.5 microgra	am tablet		
	Lanoxin. Sigmaxin 250 micro	ogram table	et		
	Lanoxin Paediatric Elixir 50 r	nicrogram	/mL (contains propylene	glycol: approximately	52 mg/mL and
	ethanol: approximately: 84	mg/mL, eq	uivalent to 10.6% absolu	ite volume)	3.
	INTRAVENOUS:	0, , ,			
	Lanoxin Infant Solution for I	nfusion 50	microgram/2mL		
	Lanoxin Solution for Infusior	n 500 micro	ogram/2mL. CAUTION: C	ONCENTRATED produc	t
	Both contain ethanol, propy	lene glyco	, citric acid and sodium	phosphate.	
Dose				1	
	Term ≥37⁺⁰ weeks	Route	Frequency	Number of doses	Dose microgram/kg/dose
	Loading	Oral	8 hourly	3 doses	10
		IV*	8 hourly	3 doses	7.5
	Maintenance [#]	Oral	daily	daily	8 (up to 12 [#])
	8 hours after last loading dose	IV*	daily	daily	6 (up to 9*)
		1		T	1
	Preterm ≤36 ⁺⁶ weeks	Route	Frequency	Number of doses	Dose microgram/kg/dose
	Loading	Oral	8 hourly	3 doses	10
		IV*	8 hourly	3 doses	7.5
	Maintenance [#]	Oral	daily	daily	5-7.5 (up to 12*)
	8 hours after last loading dose	IV*	daily	daily	3.8-5.6 (up to 9*)
			1		
	Infants	Route	Frequency	Number of doses	Dose
	Infants 2-24 months	Route	Frequency	Number of doses	Dose microgram/kg/dose
	Infants 2-24 months Loading	Route Oral	Frequency 8 hourly	Number of doses	Dose microgram/kg/dose
	Infants 2-24 months Loading	Route Oral IV*	Frequency 8 hourly 8 hourly	Number of doses 2-3 doses 2-3 doses	Dose microgram/kg/dose 10 7.5 8.10
	Infants 2-24 months Loading Maintenance [#]	Route Oral IV* Oral	Frequency 8 hourly 8 hourly Daily or 2 divided doses	Number of doses 2-3 doses 2-3 doses Daily or 2 divided doses	Dose microgram/kg/dose 10 7.5 8-10
	Infants 2-24 months Loading Maintenance [#] 8 hours after last loading dose	Route Oral IV* Oral IV*	Frequency 8 hourly 8 hourly Daily or 2 divided doses Daily or 2 divided	Number of doses 2-3 doses 2-3 doses Daily or 2 divided doses Daily or 2 divided doses	Dose microgram/kg/dose 10 7.5 8-10 6-7.5
	Infants 2-24 months Loading Maintenance [#] 8 hours after last loading dose *IV dose: 75% of oral dose	Route Oral IV* Oral IV*	Frequency 8 hourly 8 hourly Daily or 2 divided doses Daily or 2 divided doses	Number of doses 2-3 doses 2-3 doses Daily or 2 divided doses Daily or 2 divided doses	Dose microgram/kg/dose 10 7.5 8-10 6-7.5
	Infants 2-24 months Loading Maintenance [#] 8 hours after last loading dose *IV dose: 75% of oral dose #Maintenance dose may inco cardiologist.	Route Oral IV* Oral IV*	Frequency 8 hourly 8 hourly Daily or 2 divided doses Daily or 2 divided doses rding to therapeutic drug	Number of doses 2-3 doses 2-3 doses Daily or 2 divided doses Daily or 2 divided doses monitoring and in cor	Dose microgram/kg/dose 10 7.5 8-10 6-7.5
	Infants 2-24 months Loading Maintenance [#] 8 hours after last loading dose *IV dose: 75% of oral dose #Maintenance dose may incl cardiologist. Doses should be titrated to a When switching from oral to	Route Oral IV* Oral IV* rease acco the lowest	Frequency 8 hourly 8 hourly Daily or 2 divided doses Daily or 2 divided doses rding to therapeutic drug dose needed to achieve y, reduce the digoxin do	Number of doses 2-3 doses 2-3 doses Daily or 2 divided doses Daily or 2 divided doses g monitoring and in cor effect. sage by 20–25% as in ta	Dose microgram/kg/dose 10 7.5 8-10 6-7.5 sultation with
Dose adjustment	Infants 2-24 months Loading Maintenance [#] 8 hours after last loading dose *IV dose: 75% of oral dose #Maintenance dose may incl cardiologist. Doses should be titrated to the When switching from oral to Renal impairment: Predomi impairment	Route Oral IV* Oral IV* rease acco the lowest OIV therap nantly ren	Frequency 8 hourly 8 hourly Daily or 2 divided doses Daily or 2 divided doses rding to therapeutic drug dose needed to achieve y, reduce the digoxin do ally cleared (about 70%)	Number of doses 2-3 doses 2-3 doses Daily or 2 divided doses Daily or 2 divided doses g monitoring and in cor effect. sage by 20–25% as in tag reduce dose by at leas	Dose microgram/kg/dose 10 7.5 8-10 6-7.5 hsultation with able above.

Digoxin Newborn use only

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Maximum dose	250 microgram daily	
Total cumulative		
dose		
Route	Oral	
	Intravenous	
Preparation	IV	
	CHECK PRODUCT SELECTION CAREFULLY. Dilution only applies to Lanoxin Infant Injection.	
	Lanoxin Infant Injection:	
	Add 2mL (50 microgram) of digoxin to 8 mL of sodium chloride 0.9% or glucose 5% to make a 5 microgram/mL	
	solution.	
Administration	ORAL: May be taken with or without food. ³² However, administer consistently at the same time with	
	respect to meals to avoid day to day variation. ³³	
	IV: Give over at least 10 minutes.	
	IM: Do not give IM (unpredictable absorption, local irritation).	
Monitoring	Check renal function and electrolyte concentrations before starting digoxin.	
	For intravenous infusion, continuous cardiac monitoring is recommended. It may not be necessary when IV	
	injection is used to temporarily replace oral dosing in a patient stabilised on digoxin. Check local guidelines.	
	The onset of effect is approximately 5 to 10 minutes, with a maximum effect being achieved after 2 hours.	
	Take drug levels at least 6 hours after the dose is given.	
	For oral treatment without loading dose, steady state is reached after about 7 days if renal function is	
	normal (half-life is 36 hours); this may be prolonged in renal impairment.	
	The therapeutic range for those with atrial tachyarrhythmias is 0.5 to 2 microgram/L (0.6 to 2.6 nmol/L) as	
	toxicity is more common at digoxin concentrations >2 microgram/L. However, toxic effects can occur at	
	lower concentrations, particularly in the elderly or in those with electrolyte disturbance, hypoxia or	
	hypothyroidism. GI symptoms (e.g. nausea, anorexia) may precede cardiac symptoms (e.g. arrhythmias).	
	Heart failure: Consider maintaining lower concentrations of 0.5 to 0.8 microgram/L (0.6 to 1 nmol/L) in	
	patients with heart failure who are in sinus rhythm.	
	Therapeutic drug monitoring for digoxin should be performed using an assay free from interference with	
	digoxin-like immunoreactive factors, spironolactone, canrenoate, digoxin metabolites and steroids.	
Contraindications	Contraindicated in second- or third-degree heart block (without pacemaker), SVT involving accessory	
	pathway (Wolff-Parkinson-White syndrome), ventricular tachycardia and ventricular fibrillation,	
	hypertrophic obstructive cardiomyopathy, cor pulmonale (acute and chronic) or constrictive pericarditis.	
Precautions	In acute myocardial infarction, ischaemic heart disease or myocarditis, digoxin increases risk of	
	arrhythmias.	
	Use digoxin cautiously in sick sinus syndrome (risk of severe bradycardia or sinoatrial block).	
	Digoxin may worsen cardiac function in severe aortic stenosis because it increases the force of myocardial	
	contraction.	
	Digoxin increases risk of arrhythmias after DC cardioversion; withhold digoxin for 1–2 days before	
	cardioversion or use lowest effective energy.	
	Hyperthyroidism—may decrease digoxin concentration and increase sympathetic tone; monitor digoxin	
	concentration and alter dose when required or combine with another agent; dosage adjustment may be	
	required when condition is corrected.	
	Hypothyroidism—may increase digoxin concentration; monitor digoxin concentration and alter dose as	
	required; dosage adjustment may be required when condition is corrected.	
	Hypokalaemia, nypomagnesaemia, nypercalcaemia, acidosis, nypoxia—may increase sensitivity to digoxin	
Dura internetiene	(especially hypokalaemia); symptoms of toxicity may occur at lower digoxin concentrations.	
Drug interactions	reatment with drugs that slow cardiac conduction, cause bradycardia or arrhythmias may potentiate the	
	cardiac adverse effects of digoxin; use combinations carefully and monitor cardiac function.	
	reatment with drugs that inhibit or induce P-glycoprotein (ABCB1) may increase the risk of adverse	
	effects or decrease digoxin's efficacy.	
	Use of discuir and agoin increases risk of bradycardia and AV block - additive effect.	
	Use of algorin and amiodarone increases risk of dysrnythmias and torsade de pointes as amiodarone	
	blocks P-glycoprotein (ABCB1). Forsade de pointes might by facilitated by bradycardia caused by digoxin.	
	Use of digoxin and azoles, clarithromycin and some HIV-protease inhibitors increases risk of dysrhythmias	
	by inhibition of P-glycoprotein (ABCB1).	

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	Use of digoxin and non-dihydropyridine calcium channel blockers increases risk of bradycardia, asystole and sinus arrest by inhibition of P-glycoprotein (ABCB1) and their synergistic effect on the heart. Use of digoxin and loop or thiazide diuretics, amphotericin B, corticosteroids increase risk of dysrhythmias as hypokalaemia potentiates digoxin toxicity. Use of digoxin and IV calcium increases risk of dysrhythmias as hypercalcemia increases effect of cardiac glycosides. Use of digoxin and propafenone increases risk of dysrhythmia probably by inhibition of P-glycoprotein (ABCB1) by propafenone. P-glycoprotein (ABCB1)-inducers: Carbamazepine; phenytoin; rifampicin; St John's wort; tipranavir.
	P-glycoprotein (ABCB1)-inhibitors: Amiodarone, azithromycin, carvedilol, ciclosporin, clarithromycin, cobicistat, daclatasvir, erythromycin, everolimus, glecaprevir with pibrentasvir, isavuconazole, itraconazole, ketoconazole, lapatinib, ledipasvir, ritonavir, ticagrelor, tolvaptan, vandetanib, velpatasvir, vemurafenib, venetoclax, verapamil.
Adverse	Digoxin may worsen arrhythmias (proarrhythmic effect).
reactions	Digoxin has a narrow therapeutic range; adverse effects are related to its plasma concentration and very few occur at < 0.8 microgram/L (1 pmol/L)
	Digovin usually has an effect on the ECG and may result in prolonged PR interval ST depression or T wave
	inversion (these changes do not necessarily indicate digoxin toxicity or myocardial ischaemia)
	In children, arrhythmias (including sinus bradycardia) are the earliest and most frequent indicators that
	digoxin dosage is too high.
	Common (>1%): Anorexia, nausea, vomiting, diarrhoea, visual disturbances (e.g. blurred vision),
	drowsiness, dizziness, headache, rash, bradycardia, arrhythmia.
	Infrequent (0.1–1%): Depression, shortened QRS complex, atrial or ventricular extrasystoles, paroxysmal
	atrial tachycardia with AV block, ventricular tachycardia or fibrillation, heart block.
	Rare (<0.1%): Thrombocytopenia, seizures, confusion, psychosis, gynaecomastia (long-term use).
Compatibility	Fluids: Glucose 5%, Hartmann's, sodium chloride 0.9%, sodium chloride 0.45%. Not tested: glucose 10%.
	Y-site (30,32): Aciclovir, amikacin, aminophylline, amphotericin B lipid complex, ascorbic acid injection,
	atenolol, atracurium, atropine, azathioprine, azithromycin, aztreonam, calcium chloride, calcium
gluconate, capreomycin, cetalotin, cetazolin, cetotaxime, cetoxitin, cettazidime, ceftriaxone, cefu	
	sodium phosphate devmedetomidine dobutamine donamine dovucucline enalaprilat eninenbrine
	enoietin alfa, ervthromycin lactohionate, fentanyl, fluorouracil, folic acid (as sodium salt), furosemide
	ganciclovir, gentamicin, glycopyrrolate, heparin sodium, hydrocortisone sodium succinate, imipenem-
	cilastatin, indomethacin, isoproterenol, kanamycin, ketamine, labetolol, lidocaine, lincomycin, linezolid.
	lorazepam, magnesium sulfate, Meropenem, methylprednisolone sodum succinate, metronidazole,
	midazolam, milrinone, morphine sulfate, multiple vitamin injection, naloxone, netilmicin, nitroglycerin,
	nitroprusside sodium, norepinephrine, octreotide, pamidronate, penicillin G potassium, penicillin G
	sodium, pentobarbital, phenobarbital, piperacillin, piperacillin-tazobactam, potassium acetate, potassium
	chloride, propranolol, protamine, pyridoxine, ranitidine, remifentanil, rocuronium, sodium acetate, sodium
	bicarbonate, streptokinase, succinylcholine, suxamethonium, theophylline, thiamine, ticarcillin-
	clavulanate, tobramycin, tolazoline, urokinase, vancomycin, vasopressin, vecuronium, voriconazole.
Incompatibility	Fluids: No information
	V cite (20.22), Amindarana, amphatariain D chalactary sulfate complay, amphatariain D conventional
	selleidel amphotoricin B linesome caspofungin diazonam diazovide flusonazole phonytoin propofel
	sulfamethoxazole-trimethonrim Adrenaline (eninenbrine) amiodarone casnofungin fluconazole
	foscarnet, pentamidine, propofol
Stability	Infusion solution: Stable for up to 6 hours at 25° C.
Storage	Ampoule and oral elixir: Store below 25° C. Protect from light.
Excipients	Elixir: sucrose 30% w/v, sodium phosphate, citric acid, ethanol, propylene glycol, colouring (quinoline
	yellow Cl47005), methyl hydroxybenzoate, water
	IV: propylene glycol 41.5% w/v, ethanol, citric acid, sodium phosphate, water for injections
Special	Bioavailability of oral dose 60 to 85%.
comments	Half-life in infants 18 to 25 hours. 50 to 70% excreted in urine unchanged. Minimally metabolised by
	hepatic and intestinal enzymes to active and inactive metabolites.

	Onset of effect occurs 0.5–2 hours after initial oral dose of 500–750 micrograms and 5–30 minutes after initial IV dose of 400–600 micrograms; maximal effect occurs after 1–4 hours (IV) or 2–6 hours (oral). Regularly assess patients for digoxin toxicity (including resting heart rate); routine measurement of pulse
	rate before giving next dose of digoxin is not necessary.
	Assume that any arrhythmia that occurs in a child taking digoxin is due to the drug until proven otherwise.
	• Dose initially with one vial (40 mg diluted in 4 mL of water for injections) and repeat if symptoms
	persist or recur.
	• Full neutralisation dose of DigiFab is: Number of vials = serum digoxin concentration (nanogram/mL) x
	weight (kg) / 100 (rounded up to nearest vial). However, this is rarely indicated.
Evidence	Efficacy
	Heart failure: Digoxin has traditionally been used in the setting of atrial fibrillation and advanced heart failure. In a systematic review of the offects on total mortality in patients with systelic heart failure, digoxin
	did not reduce all-cause and heart failure mortality but did reduce heart failure symptoms and
	readmissions for heart failure by 32% (OR 0.68, 95% CI 0.61–0.75, P <0.00001). Benefits appeared greater
	in patients with severely reduced ejection fraction (≤25%) or NYHA III–IV functional class. Post-hoc
	subgroup analyses by serum digoxin concentrations (SDC) found patients within the range 0.5–0.8 ng/mL
	had their risk of all-cause mortality reduced by 20% (HR 0.80, 95% Cl 0.68–0.94, P = 0.005). Increased
	arrhythmic complications have been identified in patients with SDC concentrations \geq 1.2 ng/mL. If used in the context of any renal impairment, digovin requires very careful dose and level monitoring to prevent
	toxicity.[1, 2]
	In a systematic review of RCTs of digoxin therapy for cor pulmonale in adult patients, 4 studies with only 76
	patients were included and found overall there was no statistically significant improvement in RVEF,
	exercise capacity, NYHA class, heart failure score or body weight.[3]
	However, there are no RCTs comparing digoxin versus placebo or other drug therapy in infants with heart failure. Digoxin has been a component of standard treatment in several trials of other drug therapy in
	paediatric populations with heart failure in the context of congenital heart disease [4-7] and dilated
	cardiomyopathy [8, 9]. One of these trials, Buchhorn et al 2001 in an RCI of propranoiol and standard
	and left-to-right shunts reported propranolol treatment but not digoxin and diuretics alone reduced
	clinical symptoms of heart failure.
	Recommendation: The Pediatric Cardiac Intensive Care Society 2014 Consensus Statement reported that
	digoxin is not currently used as a first-line therapy in the management of heart failure. Digoxin has a class
	left ventricular election fraction unless otherwise contraindicated. The current recommendations are
	based on results from the Digitalis Investigation Group study that showed no mortality benefit over
	placebo, but did document a reduction in overall hospitalizations and heart failure-related
	hospitalizations). Careful attention to dosing and concomitant renal dysfunction must be considered when
	using digoxin. Serum levels of 0.5–0.9 ng/mL are typically targeted for optimal benefit. Digoxin should be
	theranies that may alter digoxin levels including amiodarone and/or beta blockers [10] [I OF III-2 GOR D]
	Treatment of symptomatic patent ductus arteriosus (PDA): A single RCT reported 15 preterm infants
	weighing \leq 1,500 gm at birth who had a symptomatic PDA were treated according to a medical
	management protocol (fluid restriction, digoxin and trusemide) versus 10 treated with early surgical
	digoxin for management of symptomatic PDA is unclear. [I OF II GOR D]
	Management of supraventricular tachycardia in children: [11]
	Haemodynamically unstable: Cardioversion is the definitive intervention to terminate SVT in children who
	are haemodynamically unstable. Adenosine may be given while preparing to cardiovert if the drug is
	while preparing for cardioversion or drug therapy, but cardioversion should not be delayed to administer
	vagal manoeuvres. Cardioversion — direct current cardioversion at 0.5 to 2.0 J/kg should be performed.

	Haemodynamically stable: Antiarrhythmic therapy — if the vagal manoeuvre does not convert SVT that is haemodynamically stable to normal rhythm, an intravenous (IV) catheter should be placed for the administration of antiarrhythmic drugs. Adenosine is the drug of choice for acute management of SVT; procainamide and amiodarone are sometimes given for tachycardia that is refractory to adenosine. For SVT that is refractory to adenosine, choices for IV antiarrhythmic therapy include procainamide and amiodarone. Digoxin is not usually used because of the delay in achieving therapeutic levels and the narrow therapeutic margin with the risk of serious toxicity. In addition, digoxin should not be given if WPW syndrome is suspected, since it may potentiate accessory pathway conduction.
	Sanatini et al 2012 [12] in a RCT of 61 infants <4 months with SVT (atrioventricular reciprocating tachycardia or atrioventricular nodal re-entrant tachycardia excluding Wolff-Parkinson-White) compared digoxin (loading dose 30 microgram/kg/day, maintenance 10.5 microgram/kg/day) versus propranolol (0.5 mg/kg as a single dose then 1.0 mg/kg/dose 8-hourly). SVT recurred in 19% of patients on digoxin and 31% of patients on propranolol (P = 0.25). No first recurrence occurred after 110 days of treatment. The 6-month recurrence-free status was 79% for patients on digoxin and 67% for patients on propranolol (P = 0.34), and there were no first recurrences in either group between 6 and 12 months. There were no deaths and no serious adverse events related to study medication.
	Hornik et al 2014 [13] in a retrospective cohort of infants with SVT from the Pediatrix Medical Group neonatal ICU database compared 342 infants exposed to digoxin versus 142 infants exposed to propranolol. The incidence rate of treatment failure was 6.7/1,000 infant-days of exposure to digoxin and 15.4/1,000 infant-days of exposure to propranolol. Treatment failure was higher on propranolol when compared with that on digoxin (adjusted hazard ratio, 1.97; 95% CI 1.05–3.71). Hypotension was more frequent during exposure to digoxin versus propranolol (39.4 vs 11.1/1,000 infant-days; p <0.001). There was no difference in frequency of other clinical adverse events.
	Bolin et al 2017 [14] reported a retrospective cohort of infants with SVT from the Pediatric Health Information System database admitted at ≤2 days of age with structurally normal hearts and treated with an antiarrhythmic medication. 2,657 neonates were identified with a median gestational age of 37 weeks (interquartile range 34 to 39). Digoxin and propranolol were most commonly prescribed; digoxin use steadily decreased to 23% of antiarrhythmic medication administrations over the study period, whereas propranolol increased to 77%. Multivariable comparisons revealed that the odds of mortality for neonates on propranolol were 0.32 times those on digoxin (95% CI 0.17 to 0.59; p <0.001). Propranolol for the neonate with SVT is associated with lower in-hospital mortality and hospital costs compared with digoxin.
	Recommendation: ANZCOR recommendation for pharmacological management of specific dysrhythmias in the paediatric advanced life support guideline is that, for SVT, adenosine is the drug of choice. Amiodarone may be used to treat haemodynamically stable or unstable SVT. Alternative drugs are procainamide, digoxin, a beta blocker or a calcium channel blocker. Calcium channel blockers should not be used to treat SVT in infants and should be avoided or used cautiously in children because they may induce hypotension and cardiac depression.[15]
	Atrial fibrillation — Atrial fibrillation is uncommon in children and most paediatric cases are associated with CHD, cardiomyopathy or Wolff-Parkinson-White syndrome.[16] The management of neonatal atrial fibrillation is unclear with use of digoxin and cardioversion reported.[17, 18] In adult populations, systematic review found when digoxin was compared with all control interventions there was no evidence of a difference in all-cause mortality (RR 0.82; Cl 0.02 to 31.2); serious adverse events (RR 1.65; Cl 0.24 to 11.5); quality of life; heart failure (RR 1.05 Cl 0.00 to 1141.8) or stroke (RR 2.27; Cl 0.00 to 7887.3). Digoxin was superior compared with placebo in reducing the heart rate, but inferior compared with beta blockers. Meta-analyses on acute heart rate control showed that digoxin was inferior compared with both calcium antagonists (MD 21.0 bpm; Cl -30.3 to 72.3) and with amiodarone (MD 14.7 bpm; Cl -0.58 to 30.0). Meta-analysis on acute conversion to sinus rhythm showed that digoxin compared with amiodarone reduced the probability of converting atrial fibrillation to sinus rhythm (RR 0.54; Cl 0.13 to 2.21).[19]
	Atrial flutter: Atrial flutter can occur in fetuses and neonates with structurally normal hearts. Comorbid conditions are not usually present; however, cases of atrial flutter associated with neonatal Coxsackie myocarditis and following maternal treatment with lithium have been reported. Neonatal atrial flutter

rarely reoccurs following cardioversion with or without medical treatment. In the newborn with atrial

flutter, initial therapy with digoxin has been the traditional approach. However, this has never been demonstrated to be any more efficacious than primary electrical cardioversion. [16] Casey et al reported a case series of 25 newborns with atrial flutter; 7 of 21 converted to sinus rhythm with digoxin therapy and electrical conversion resulted in sustained sinus rhythm in 9 of 16 patients (13 after failure of digoxin and 3 as the first treatment). Sinus rhythm was achieved in 23 patients and two died of complications of prematurity without resolution of atrial flutter.[20] Texter et al 2006 reported a case series of 50 infants with atrial flutter, 72% presented within the first 48 hours of life. Sinus rhythm was restored in 20 of 23 (87%) attempts at direct current cardioversion and 7 of 22 (32%) attempts at transoesophageal pacing; 7 required antiarrhythmic therapy. An additional arrhythmia, all supraventricular, appeared in 11 (22%) infants. The recurrence of atrial flutter developed in 6 infants all with an additional arrhythmia. Twelve received digoxin loading as first-line therapy. Sinus rhythm occurred in 4 infants within hours of beginning
the digoxin load; the remaining eight required additional intervention.[21]
Recommendation: In the newborn with atrial flutter, initial therapy with digoxin has been the traditional
approach. However, this has never been demonstrated to be any more efficacious than primary electrical
cardioversion.[16]
Safety
Jailing groups digovin is associated with a neutral effect on martality in randomized trials and a lower
rate of admissions to hospital across all study types.[22] However, in a meta-analysis of hospital adverse drug reactions (ADRs), the mean fatal ADR prevalence varied from 0.01% in paediatric patients to 0.44% in the elderly. Warfarin, aspirin, renin-angiotensin system inhibitors and digoxin accounted for 60% of fatal ADRs.[23]
Ventricular fibrillation following adenosine therapy for SVT in a neonate with concealed Wolff-Parkinson- White sundrome treated with digavin has been reported [24]
While syndrome treated with digoxin has been reported.[24]
(ABCB1) such as verapamil, amiodarone or macrolide antibiotics can enhance oral bioavailability of digoxin by decreasing its efflux from the enterocytes into the lumen of the intestine and decrease its active tubular
secretion into the urine in the kidney. As a result, plasma concentrations of digoxin may significantly
increase to toxic levels [see drug interactions]. Recommended window of therapeutic concentrations is quite narrow (0.8–2.0 ng/mL) and more recent recommendations suggest even lower and more narrow range (0.5–1.0 ng/mL) [25]
Increased arrhythmic complications have been identified in patients with serum digoxin concentrations ≥1.2 ng/mL. If used in the context of any renal impairment, digoxin requires very careful dose and level monitoring to prevent toxicity [1]
Hypokalaemia increases the incidence of arrhythmias and sudden cardiac death. The risk is increased in
natients with pre-existing heart disease and in those treated with digoxin
Although cases of digoxin poisoning are fewer than those involving calcium channel and beta blockers, the
mortality rate from digoxin is far greater.
Specific antidote therapy with digoxin-specific antibody fragments (digoxin-Fab) should be used if there are
arrhythmias associated with haemodynamic instability. Digoxin-Fab interferes with digoxin immunoassay
measurement and can lead to overestimation of plasma digoxin concentrations.[26]
Lanoxin Paediatric Elixir contains approximately 52 mg/mL of propylene glycol and 84 mg/mL of ethanol.
equivalent to 10.6% absolute volume (email correspondence with the manufacturer on 21 st March 2019).
Long-term effects of prolonged exposure to ethanol content are unknown.
Pharmacokinetics/pharmacodynamics
Digoxin is a cardiac glycoside. Digoxin's mechanism of action is related to both causing an increase in
parasympathetic tone as well as inhibition of the Na ⁺ /K ⁺ ATPase, which indirectly increases intracellular
calcium. Its onset of action is 5 to 60 minutes when given intravenously, with peak effect seen in 1 to 6
hours. When given orally, onset of action is 1 to 2 hours. with peak effect seen at 2 to 8 hours. The half-life
of digoxin varies by age, ranging from 61 to 170 hours in preterm neonates. from 35 to 45 hours in full-
term neonates and from 18 to 25 hours in infants. [27] Digoxin toxicity in neonates and infants can present
as significant bradycardia or cardiac arrhythmias. Digoxin is contraindicated in patients with WPW because
of its effect on the accessory pathway and the AV node causing predisposition for fatal arrhythmias.[28]
Monitoring

	Digoxin has 11 different methodologies reported Australia and New Zealand laboratories for therapeutic	
	drug monitoring (TDM). Digoxin immunoassays may have a problem with interference from digoxin-like	
	immunoreactive factors, spironolactone, canrenoate, digoxin metabolites and steroids.[29]	
Practice points	, , , , , , , , , , , , , , , , ,	
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