Flecainide Newborn use only

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Alert	Use in consultation with a Paediatric Cardiologist.	
	Contraindicated in infants with reduced myocardial contractility.	
	Use caution in patients with congenital heart disease—increased potential for pro-arrhythmic effects.	
Indication	Intravenous flecainide needs close cardiorespiratory monitoring Treatment of paroxysmal supraventricular tachycardia, paroxysmal atrial fibrillation/flutter and life-	
mulcation	threatening ventricular dysrhythmias as a second-line agent where tachycardia has been resistant to first-	
	line agents.	
Action	Decreases intracardiac conduction for all parts of the heart, with the greatest effect in the His-Purkinje	
	system. It acts by blocking fast sodium channels. As a type Ic agent, it slows cardiac conduction and	
	decreases contractility.	
Drug type	Type Ic antiarrhythmic agent.	
Trade name	Flecainide Sandoz Tablets; Flecatab Tablets; Tambocor solution for injection, Tambocor Tablets	
Presentation	IV:	
	10 mg/mL injection.	
	Oral:	
	20 mg/mL suspension compounded by pharmacy.	
_	50 mg, 100 mg tablets.	
Dose	Oral:	
	Starting dose: 1 mg/kg/dose 8 or 12 hourly. Increase by 1 mg/kg/dose as necessary to achieve maintenance of sinus rhythm up to the maximum dose.	
	IV:	
	2 mg/kg over at least 10 minutes.	
Dose adjustment	No information.	
Maximum dose	8 mg/kg/day	
Total cumulative		
dose		
Route	Oral [preferred route] or IV	
Preparation (for	Draw up 1mL (10mg of flecainide) and add 9mL of glucose 5% to make a final volume of 10 mL with a	
IV administration)	concentration of 1mg/mL.	
	It can also be administered undiluted.	
Administration	Oral:	
	Administer between milk feeds. Do not administer with milk. Milk decreases absorption of the drug.	
	IV:	
	Infusion over at least 10 minutes. Patient needs to be monitored very closely with the potential for an	
	acute deterioration.	
Monitoring	Initiate treatment in hospital with ECG monitoring in consultation with paediatric cardiologist.	
U	When intravenous route is used, continuous ECG monitoring is mandatory.	
	Perform ECG when the dosage is increased – monitor QRS duration and dysrhythmia.	
	Therapeutic trough concentrations are not routinely required (200–1000 microgram/L).	
Contraindications	Cardiogenic shock.	
	Hypersensitivity to flecainide.	
	Significant renal impairment (creatinine clearance < 50 mL/min).	
D	Reduced left ventricular ejection fraction.	
Precautions	Use with caution in patients with congenital heart disease or conduction system disease (right bundle	
	branch block, with left hemiblock and without pacemaker; second- or third-degree atrioventricular block, without pacemaker; sick sinus syndrome [bradycardia-tachycardia syndrome]).	
	Milk decreases oral flecainide absorption. Consider decreasing oral dose or dose monitoring if change of	
	milk diet.	
	Dosing adjustments are required in infants with renal impairment because 10% to 50% of a flecainide dose	
	is excreted in the urine.	
	Use with caution in significant hepatic impairment.	
Drug interactions	Use of any of the drugs prolonging QT interval (cisapride, amiodarone, clarithromycin, chloral hydrate,	
	ciprofloxacin, erythromycin, octreotide, sodium phosphate, vasopressin, ketoconazole, fluconazole,	
	hydrochlorothiazide, azithromycin, propranolol, digoxin, verapamil) with flecainide can lead to significant	
	increase in QT interval.	

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Adverse reactions	Adults:
	Common
	Cardiovascular: Palpitations (6.1%); Gastrointestinal: Nausea (up to 10%); Neurological: Dizziness (18.9% to 30%), Headache (4.5% to 9.6%); Ophthalmological: Blurred vision (10% to 38%), Photopsia (up to 30%); Respiratory: Dyspnoea (up to 10.3%); Other: Fatigue (7.7%).
	Serious
	Cardiac arrest, cardiac dysrhythmia, cardiogenic shock, disorder of pacing function, electrocardiogram
	abnormalities, heart block, heart failure (new onset or worsening [up to 25.7%]), prolonged QT interval,
	sinus node dysfunction (1% to less than 3%), syncope (1% to less than 3%), torsades de pointes, ventricular fibrillation, ventricular tachycardia.
	Children:
	Dizziness, blurred vision and headache have been reported in children.
Compatibility	5% glucose
Incompatibility	Incompatible with alkaline and chloride-containing solutions.
Stability	Diluted solution stable for 24 hours at 25°C.
	Oral suspension compounded by Pharmacy stable for up to 60 days.
Storage	
Excipients	Silicified microcrystalline cellulose, croscarmellose sodium, maize starch, magnesium stearate.
Special comments	
Evidence	Efficacy and safety:
	A review of published cases and subsequent reports found flecainide appeared to be safe (no deaths with
	usual oral dosing; < 1% incidence of serious proarrhythmia) and effective (73–100 % control, depending on
	mechanism) in children with supraventricular tachycardia. [1-4] (LOE IV GOR B) However, concerns
	regarding safety exist in patients with structural heart disease and cardiomyopathy. The Cardiac
	Arrhythmia and Suppression Trial (adults with AMI) demonstrated increased mortality in patients who
	received flecainide.[3-5] A report of young patients (4 days to 26 years) administered flecainide for
	treatment of SVT (n = 369) or VT (n = 103) found efficacy 71.4%, proarrhythmic response 7.4%, cardiac
	arrest 2.3% and died during treatment 2.1%. Cardiac arrest and deaths occurred predominantly among
	patients with underlying heart disease, particularly among patients receiving flecainide for
	supraventricular tachycardia (8.3%).[3] A report in children (n = 229) with congenital heart disease or
	cardiomyopathy, incidence of cardiac arrest in patients receiving flecainide was 3.0% with a mortality of
	4.3%, with no difference in cardiac arrest or mortality rate when compared to patients who received other
	antiarrhythmics.[4]
	Guidelines: For SVT, flecainide is effective as a first-line agent in infants, but typically used as a second-line
	agent because of its arrhythmogenic potential. It has been used in infants with re-entrant supraventricular tachycardia including Wolff- Parkinson-White syndrome, focal atrial tachycardia and permanent junctional reciprocating tachycardia (case reports). Has the potential for proarrhythmia in patients with congenital heart disease. Caution is advised when used in patients with congenital heart disease or conduction
	system disease. Milk feeds may decrease absorption. Concentration monitoring may assist in guiding therapy. Contraindicated if creatinine clearance <50 mL/min or reduced Left Ventricular Ejection Fraction.[6] (LOE IV GOR B)
	Bharmacokinotice:
	Pharmacokinetics: Flecainide is cleared via hepatic biotransformation and renal excretion. Infants < 1 year of age had a mean
	t ₂ of 11–12 hour; children aged 1 to 12 years had a t ₂ of 8 hours. Dosing schedules based on mg/m ² correlated better with plasma flecainide concentrations than did dosing based on mg/kg.[8, 9] Oral
	bioavailability in adults reported to be 78–100%.
Practice points	
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VERSION/NUMBER	DATE
Original 1.0	02/03/2017
Current 2.0	12/01/2021
REVIEW	12/01/2026

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