# Omeprazole

### Newborn use only

Alert Short and long-term safety data in infants are limited.	
Indication Treatment of gastroesophageal reflux disease (GORD).	
Prophylaxis in congenital tracheoesophageal fistula and oesophageal atresia (role unclea	
Action Proton pump inhibitor (PPI). Bind to the hydrogen/potassium ATPase enzyme system (pro	oton pump),
inhibiting both stimulated and basal acid secretion.	
Drug Type Proton Pump Inhibitor.	
Trade Name Oral tablet: Multiple brands available.	
Oral capsule: Multiple brands available.	
Oral suspension: Omeprazole (PediPPI) (powder for) oral suspension 2mg/mL (75mL) ava	ilable from
Symbion via special access scheme	
IV: Omeprazole Sandoz Powder for Injection.	
Presentation Oral: Available in 10mg and 20 mg. Available in capsules or enteric coated tablets.	
Oral suspension of 2 mg/mL, 5mg/mL or other strengths may be prepared in pharmacy.	Omeprazole
(PediPPI) 2mg/mL powder for oral suspension available via special access scheme	
IV: 40mg/vial of Omeprazole in dry powder form.	
Dose PO: 1-2.5 mg/kg/day in 1 to 2 divided doses.(1,2)	
IV: 0.5 mg/kg/dose 12-24 hourly (3,4,5,6)	
<b>Dose adjustment</b> Therapeutic hypothermia – No information.	
ECMO – No information.	
Renal impairment – No dose adjustment is required.	
Hepatic impairment – Dose reduction is recommended. However, no specific information	n available.
Maximum daily 2.5 mg/kg/day (1)	
dose	
Total cumulative	
dose	
Route PO, IV	
<b>Preparation PO:</b> Prepared by hospital pharmacy – No preparation is required.	
Powder for oral suspension: May require reconstitution.	
IV: Add 10 mL of sodium chloride 0.9% to 40 mg powder for reconstitution to make a con	centration of 4
mg/mL. Draw up 1 mL (4 mg) and add 9 mL of sodium chloride 0.9% to make a final volum	e of 10 mL with
a concentration of 0.4 mg/mL.	
Administration PO: Administer prior to meals. Shake the bottle well before administration.	
IV: Infuse over 30 minutes.	
Monitoring Serum magnesium, in patients on prolonged therapy or who use digoxin or drugs that magnesium.	ay cause
hypomagnesaemia (e.g. diuretics) concomitantly.	
Serum vitamin $B_{12}$ — every 1 to 2 years in patients on prolonged therapy.	
Contraindications Hypersensitivity to any component of the product.	
<b>Precautions</b> Short- and long-term safety data in infants are limited. There have been safety concerns	with long term
usage in adults. Current FDA's maximum recommended duration of therapy of PPIs is up	to 8 weeks.
<b>Drug Interactions</b> Concurrent use of fluconazole may result in increased plasma concentrations of omepraz	ole.
Concurrent use of iron may result in reduced non-heme iron bioavailability.	
Omeprazole is mainly metabolised via hepatic cytochrome P450 system (CYP2C19) and m	nay be
expected to interact with the metabolism of other drugs metabolised by this enzyme.	
Omeprazole may reduce phenytoin clearance – monitor phenytoin levels.	
Omeprazole produces a profound and sustained inhibition of gastric acid secretion. The a	bsorption of
compounds whose absorption depends on gastric pH (e.g. ketoconazole, itraconazole, er	lotinib etc.)
may decrease and the absorption of drugs such as digoxin can increase during treatment	with
omeprazole. Monitor digoxin levels.	
Adverse Increased risk of neonatal intestinal and pulmonary infections.	
Reactions Hypomagnesaemia.	
Compatibility Fluids: Glucose 5%, sodium chloride 0.9%	
i idias. Gideose 5/0, sodidii cilionde 0.3/0	
Y-site: Cisatracurium, Furosemide, Morphine sulfate, Temocillin	

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	Omeprazole is well tolerated clinically and with respect to laboratory tests. There are potential risks including increase of neonatal intestinal and pulmonary infections and occurrence of severe hypomagnesaemia.(1,11-15)
Practice points	
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