### **Newborn use only**

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Alert	ORAL ADMINISTRATION ONLY					
	The first dose of rotavirus vaccine should be given to infants between 6 and 14 weeks chronological					
	age (prior to turning 15 weeks chronological age) and the second dose by 24 weeks of age (prior to					
	turning 25 weeks of age).					
	The interval between dose 1 and 2 should not be less than 4 weeks.					
	In Australia and New Zealand, regular look up for any online updates by the Immunisation Handbook					
	is recommended.					
Indication	Primary immunisation against rotavirus gastroenteritis.					
Action	Live attenuated human rotavirus vaccine that induces protective immunity against the G1P (8) strain					
	and some other non-G1 prevalent strains of rotavirus.					
Drug Type	Vaccine.					
Trade Name	Rotarix					
Presentation	1.5 mL oral suspension in an oral applicator with plunger stopper or in a squeezable tube.					
Dose	1.5 mL orally.					
	<b>Primary schedule:</b> 2-dose course administered with 2- and 4-month immunisations i.e., dose 1 can be					
	administered at 6 to 14 weeks of age and dose 2 can be administered at 14 to 24 weeks of age.					
	<b>NOTE:</b> Dosage interval between first and second doses must be greater than 4 weeks.					
	The first sound in the first and second descent must be greater than the citis					
	Schedule Age limit for Age limit for Minimum interval					
	first dose second dose between doses					
	2 oral doses (1.5 mL/dose) 6–14 weeks 14–24 weeks 4 weeks					
	NOTE: If most of the oral rotavirus vaccine has been regurgitated or vomited within minutes of					
	administration, a single repeat dose can be administered during the same immunisation encounter. If					
	an infant regurgitates or vomits only a small part of a vaccine dose, it is not necessary to repeat the					
	dose.					
	Catch-up schedule: If an infant has NOT had a dose of any rotavirus vaccine AND is ≥ 15 weeks then					
	that infant is NOT ELIGIBLE to commence any rotavirus vaccination dose. <sup>1</sup>					
	Preterm infants: Vaccine is administered at a chronologic age (without correction for prematurity)					
	similar to term infants, if the infant is clinically stable. <sup>1</sup>					
	<b>Hospitalised infants:</b> If standard infection control precautions are maintained and the infant is					
	medically stable, vaccination should not be delayed, particularly if the delay would result in an infant					
	being beyond the upper age limit for vaccination. <sup>1</sup>					
	Systemic corticosteroid therapy: Rotavirus vaccine is not contraindicated in neonates on inhaled or					
	systemic corticosteroids if they are otherwise medically stable. <sup>1</sup>					
	<b>Exposure to anti-CD20 therapy in utero:</b> Rotarix should be withheld in infants whose mothers were					
	taking anti-CD20 therapy (including rituximab) during pregnancy. Rotarix can be given to infants					
	whose mothers were taking other biologic immunosuppressants during pregnancy. <sup>1</sup>					
	/					
/	Other live vaccines: Rotavirus vaccine can be given at any time before or after the routine infant					
	immunisations and at any time before or after BCG vaccine. The recommendation for administering					
	live vaccines either at the same time or after an interval of four weeks only applies to injectable live					
	viral vaccines and, therefore, not to BCG or to the oral rotavirus vaccines. <sup>2</sup>					
Dose adjustment						
Maximum Dose	Limited data on the safety of administering higher than the recommended dose.					
Route	Oral or via gastric tube					
Preparation	The vaccine is ready to use; no reconstitution or dilution is required.					
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	The vaccine is presented as a clear and colourless liquid. It should be inspected visually for any foreign particulate matter and/or abnormal physical appearance. In the event of either being observed, the
	vaccine is not suitable for use.
	Oral: Administer entire applicator or dosing tube content on inside of cheek with child in reclining
	position.
	Gastric tube: For infants who can't take the vaccine orally, it can be administered via a gastric tube;
	flush with air to clear the tube.
	Can be given with or without feeds.
	Record details of vaccination in patient's Personal Health Record ('Blue Book'), Australian
	Immunisation Register and medication chart.
	Other vaccines can be given at the same time (refer to Drug interactions section).
NA - with a wine -	Discard the empty oral applicator and tip cap according to local regulations.
Monitoring	Symptoms suggestive of intussusception such as severe abdominal pain or distress, persistent
	vomiting, bloody stools, palpable abdominal mass, abdominal bloating and/or high fever.
Controlodications	Parents should be advised to seek medical advice promptly where these symptoms are evident.
Contraindications	Anaphylaxis following a previous dose of rotavirus vaccine.
	Anaphylaxis following any vaccine component.
	Previous history of intussusception or a congenital abnormality that may predispose to
	intussusception. Fatal intussusception after the second dose has been reported in infants with a
	history of intussusception after the first dose.
	Severe Immunocompromised status including maternal anti-CD20 therapy during pregnancy Severe Combined Immunodeficiency (SCID).
	Do not administer to (i) infants older than 24 weeks of age as safety has not been demonstrated,
Procautions	
Precautions	
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Drug Interactions	
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ĺ	vaccines DTPa-HBV-IPV/Hib, pneumococcal conjugate vaccine and meningococcal serogroup C
	conjugate vaccine. The studies demonstrated that the immune responses and the safety profiles of
Precautions  Drug Interactions	particularly in relation to risk of intussusception, (iii) infants with malformation of the gastrointestina tract that could predispose them to intussusception, (iii) hereditary fructose intolerance, glucose/galactose malabsorption or sucrase-isomaltase insufficiency.  If infant is > 14 weeks and inadvertently receives 1st dose of rotavirus vaccine, reassure parents and discuss minimally increased risk of intussusception. Provide information on symptoms/signs of intussusception. If infant is < 25 weeks (upper limit for dose 2 of rotavirus vaccine), and minimum interval of 4 weeks between vaccine doses can be achieved, give a second dose of rotavirus vaccine.  Use with caution in infants with underlying conditions predisposing to severe rotavirus gastroenterit (including metabolic disorders or chronic gastrointestinal disease e.g., Hirschsprung's disease, malabsorption syndrome or short gut syndrome).  Severe acute gastroenteritis (e.g. necrotising enterocolitis)  Significant acute illness or temperature greater than 38°C (postpone vaccine until neonatologist approves).  Use with caution in immunosuppressed infants (the theoretical risk for vaccine virus-associated disease is considered likely to be less than their risk from being exposed to disease from natural infection).  Infants with a moderate to severe illness should be vaccinated after recovery. In addition to the factors mentioned above, this avoids superimposing potential adverse events related to vaccination on any concurrent illness.  Minor infections, without fever or systemic upset, are not reasons to postpone vaccination.  Viral shedding in stools, particularly after the first dose, could pose a risk of transmission of virus to immunocompromised close contacts. Good hygiene practices and contact precautions MUST be observed at ALL times (i.e. washing hands regularly, especially after changing nappies).  Co-administration studies have demonstrated that rotavirus vaccine can be given concomitantly wit any of the following vaccines: Diphtheria tetanus

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Adverse	Diarrhoea, vomiting and hematochezia.
Reactions	Intussusception—inform parents of the rare risk of intussusception and how to be alert for signs and
	symptoms.
	Irritability, flatulence, abdominal pain, dermatitis, Idiopathic thrombocytopenic purpura,
	bronchopneumonia, Kawasaki disease, hypotonic-hyporesponsive episode have also been reported
	after vaccination.
Overdess	Any suspected vaccine related adverse reactions should be reported to Therapeutic Goods Authority.
Overdose	AUSTRALIA: Contact the Poisons Information Centre on 13 11 26 for information on the management
	of overdose  NEW ZEALAND: Contact the National Poisons Centre on 0800 764 766 for information on the
	management of overdose.
Composibility	
Compatibility	Other vaccines can be given concomitantly.
Incompatibility	No information.
Stability	
Storage	Store between 2 and 8°C. Do NOT freeze as this reduces potency. Protect from light. Storage above or
	below the recommended temperature may decrease potency.
Excipients	sucrose, disodium adipate, Dulbecco's Modified Eagle Medium and
	sterile water
Special	RotaTeq and interchangeability of vaccine: As of July 2017, RotaTeq (pentavalent human-bovine
Comments	reassortant rotavirus vaccine) is not used in Australia, but it is available globally. RotaTeq is given as a
	3-dose course. Upper age limit for RotaTeq is prior to 33 weeks of age. An infant might have received
	1 or 2 doses of RotaTeq overseas prior to arrival in Australia. Where possible the completion of the
	course of rotavirus vaccine should be with the same vaccine from the same manufacturer. If either
	dose 1 or dose 2 of the rotavirus vaccine is given as RotaTeq (pentavalent human-bovine reassortant
	rotavirus vaccine) a third dose of either rotavirus vaccine should be given, provided the upper age
	limit and inter-vaccine interval are observed.
Evidence	Both rotavirus vaccines have similar efficacy (around 70%) against rotavirus gastroenteritis. The
	efficacy against severe rotavirus gastroenteritis is higher and ranged from 85% to 100% in clinical trials
	in many different countries. <sup>1</sup>
	Preterm infants: Rotavirus vaccine appears safe and equally immunogenic in preterm infants compared to term infants. Vaccine is administered at a chronologic age (without correction for
	prematurity) similar to term infants, if the infant is clinically stable. <sup>1-7</sup>
	Hospitalised infants: If standard infection control precautions are maintained, administration of
	rotavirus vaccine to hospitalised infants, including hospitalised preterm infants, would be expected to
	carry a low risk for transmission of vaccine viruses. Furthermore, the rotavirus vaccine is highly
	attenuated and does not revert to a high virulence strain. Provided that the infant is medically stable,
	vaccination should not be delayed, particularly if the delay would result in an infant being beyond the
	upper age limit for vaccination. If a recently vaccinated infant is hospitalised for any reason, no
	precautions other than routine standard precautions need be taken to prevent the spread of vaccine
	virus in the hospital setting. <sup>1,2</sup>
	Vaccine recipients may have a 1–3% higher risk of developing diarrhoea or vomiting in the week after
	vaccine administration. The incidence of fever, irritability and other adverse events was similar in both
	vaccine and placebo recipients in clinical trials. 1,7-9
/	Vomiting and diarrhoea have not been noted as important adverse events in post-marketing
	surveillance of rotavirus vaccines.
	The increased risk of intussusception after rotavirus vaccination is estimated at approximately 6
	additional cases of intussusception among every 100,000 infants vaccinated, or 14 additional cases
	per year in Australia. The overall benefits of preventing gastroenteritis from rotavirus are much
	greater than the small risk of intussusception. <sup>1</sup>
	Fatal intussusception after the second dose has been reported in infants with a history of
	intussusception after the first dose. <sup>1</sup>

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	Case reports indicate prolonged vaccine virus-associated gastrointestinal disease after rotavirus				
	vaccination in infants with Severe Combined Immunodeficiency (SCID). As these infants are unlikely to				
	generate a protective immune response to the vaccine and because of the potential harm, rotavirus				
	vaccines are contraindicated for infants with SCID. <sup>1</sup>				
Practice points					
References	1. Australian Immunisation Handbook, Australian Government Department of Health and Aged Care. Accessed on 25/05/2025.				
	2. Greenbook. United Kingdom Immunisation schedule. Immunisation against infectious disease.				
	Update. Rotavirus. Chapter 27b. Accessed on 25 January 2018.				
	3. Armstrong C. AAP updates on guidelines on rotavirus vaccination. Am Fam Physician 2010;81(4):552-553				
	4. Product information:				
	https://www.ebs.tga.gov.au/ebs/picmi/picmirepository.nsf/pdf?OpenAgent&id=CP-2010-PI-06401-3 Accessed 08/05/2025.				
	5. Pereira P, Vetter V, Standaert B, Benninghoff B. Fifteen years of experience with the oral live-				
	attenuated human rotavirus vaccine: reflections on lessons learned. Expert Rev Vaccines. 2020 Aug;19(8):755-769.				
	6. Fathima P, Jones MA, Moore HC, et al. Impact of Rotavirus Vaccines on Gastroenteritis Hospitalizations in Western Australia: A Time-series Analysis. J Epidemiol. 2021 Aug 5;31(8):480-486.				
	7. Costantino C, Conforto A, Bonaccorso N, et al. Safety of Rotavirus Vaccination in Preterm Infants				
	Admitted in Neonatal Intensive Care Units in Sicily, Italy: A Multicenter Observational Study.  Vaccines (Basel). 2023 Mar 23;11(4):718.				
	8. Van Dongen JAP, Rouers EDM, Schuurman R, et al. Rotavirus Vaccine Safety and Effectiveness in				
	Infants with High-Risk Medical Conditions. Pediatrics. 2021 Dec 1;148(6): e2021051901.				
	<ol> <li>Wu Z, Li Q, Liu Y, L H, Mo Z, et al. Efficacy, safety and immunogenicity of hexavalent rotavirus vaccine in Chinese infants. Virol Sin. 2022 Oct;37(5):724-730.</li> </ol>				

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