Ranibizumab (Lucentis)

RHW Newborn use only

Alert	Used off-label in infants with retinopathy of prematurity (ROP) and must only be used for intravitreal injection by onbthalmologist only after fully informed parental consent
Indication	Aggressive posterior POP. Zone 1 Type 1 POP. Posterior Zone 2 Type 1 POP and as adjunct to failed
indication	laser treatment of ROP or where laser is not possible due to media opacity.
Action	Antineovascularisation agent. Binds to and inhibits vascular endothelial growth factor A (VEGF-A).
Drug type	Recombinant humanized IgG1 monoclonal antibody.
Trade name	
Presentation	Lucentis vial intravitreal injection 2.3 mg/0.23 ml
Fresentation	Lucentis viai intravitreal injection 2.5 mg/0.25 me
	Lucentis pre-filled syringe intravitreal injection is available but not recommended as the barrel only
	marks the adult dose of 0.5 mg and any lesser dose cannot be identified
Dose	0.12 - 0.2 mg (refer to special comments)
Dose	Dose can be repeated in 28 days if required
Dose adjustment	Not applicable
Maximum dose	
Total cumulative	
dose	
Route	Intravitreal
Preparation	Lucentis vial 2.3 mg/0.23 mL (1)
	 Disinfect the rubber stopper of the vial with appropriate antiseptic swab.
	• Attach a 5 micrometre filter needle (18G) to a 1 mL syringe using an aseptic non-touch technique.
	 Push the blunt filter needle until the needle touches the bottom edge of the vial.
	Withdraw all liquid from the vial.
	Ensure that the plunger rod is drawn back sufficiently when emptying the vial in order to
	completely empty the filter needle.
	Disconnect the syringe from the blunt filter needle.
	• Aseptically and firmly attach an injection needle (33G needle preferred) onto the syringe.
	• Expel the air from the syringe and adjust the dose to the 0.02 mL mark on the syringe.
Administration	Procedure should be performed by a suitably qualified ophthalmologist with experience with ROP
	and intravitreal injection in neonates using aseptic technique.
	Obtain informed parental/carer consent.
	• Sedate the patient as required under the supervision of neonatologist.
	• If the infant is on CPAP, presence of a CPAP mask is not compatible with an adequately isolated
	surgical field but Hudson prongs and the connecter "reversed" in direction should allow
	satisfactory draping and taping. Other option is to use high flow humidified nasal cannula (HHFNC)
	as an alternative if considered appropriate support.
	Proceduralist to scrub and wear sterile gloves.
	• Dedicated nurse assistant to be present. All staff providing care for the infant are recommended to
	wear surgical masks.
	• Use topical povidone lodine 5% as the skin and conjunctival sac preparation. Aqueous
	chlorhexidine 0.05% to 0.1% may be used in infants with hypersensitivity to povidone iodine. Wipe
	off any excess solution from the lids/skin immediately to prevent skin irritation. The conjunctival
	sac should be thoroughly irrigated with normal saline immediately after the injection.
	• Use a small fenestrated sterile drape and stick the edges down with sterile steristrips to isolate the
	surgical field.
	• Use frequent sterile topical amethocaine 0.5% to provide topical anaesthesia. A sterile cotton bud
	soaked with amethocaine 0.5% can be used to impregnate the injection site and give compression
	to lower intraocular pressure prior to injection. There is no requirement to give a subconjunctival
	injection of xylocaine as this creates chemosis and interferes with marking the injection site.
	Stabilise globe with 0.12 Bonn ophthalmic microforcep.
	• Use Castroviejo ophthalmic caliper to measure and mark the location of the injection site, which is
	1.5 mm posterior to the limbus, in the inferotemporal quadrant.

	 Compress globe for 20 seconds prior to injection with a sterile cotton bud. This lowers intraocular pressure by displacing aqueous and safeguards against the likelihood of CRA occlusion created by the pressure rise that accompanies intraocular injection. Slowly inject ranibizumab into vitreous cavity using 30 or 33g needle. Needle entry point is 1.5 mm posterior to the limbus, in the inferotemporal quadrant and enter 3-4 mm into the vitreous cavity parallel to the visual axis so as to avoid the relatively larger and more spherical neonatal crystalline lens. Perform indirect ophthalmoscope to ensure drug visible within vitreous cavity, lens is clear and central retinal artery (CRA) is perfusing. Apply gentle ocular massage if precarious and perform anterior chamber paracentesis (with 27g needle) if CRA obstructed due to increased intraocular pressure. At the discretion of ophthalmologist, either preservative free lubricant or chloramphenicol eye drops may be applied at the end of procedure and chloramphenicol eye drops may be continued three times a day for 3 days. Ophthalmologist to review within 24 hours or sooner if excessive eyelid swelling to exclude endophthalmitis.
Monitoring	Watch for any eye swelling/bleeding
Montoling	Monitor vital signs (eg. BP, heart rate, respiratory rate) throughout the procedure
	Monitor for signs and symptoms of infection or ocular inflammation
Contraindications	Hypersensitivity to the active substance or to any of the excinients
contraindications	Active or suspected ocular or periocular infections
	Active intraocular inflammation
Procentions	Pro existing arterial thromboombolic condition – a multidisciplinary team desision is required on a case
Frecautions	by case basic to access the possible impact of systemic abcorption and systemic side offacts
Drug interactions	Not applicable
Drug Interactions	Not applicable.
Adverse	Adverse effects reported in aduits treated with anti-vege for macular degeneration.
reactions	Ocular Intection
	Endensthalmitic
	Endophilialmilis Retiral detachment, retiral tears
	Retinal detachment, retinal tears
	Increased Intraocular pressure
	Corneal injuries/inflammation
	Lens opacities/cataract
	Arterial thromboembolic events
	Neonatal data are lacking.
Compatibility	Not applicable.
Incompatibility	Not applicable.
Stability	Stable in the sealed tray at 25°C for 24 hours
Storage	Vial: Store at 2°C to 8°C (refrigerate - do not freeze). Protect from light.
Excipients	Trehalose dihydrate, histidine hydrochloride monohydrate, histidine, polysorbate 20.
Special	New South Wales Paediatric Ophthalmology – ROP treating group consensus: Ranibizumab
comments	0.2mg/0.02mL intravitreal injection is the preferred option given the negligible systemic suppression of
	VEGF compared to bevacizumab. This dose equates to 40% the adult dose. 0.02mL is a workable dosage
	volume and is a simple draw up of 0.02mL from the supplied ampoule. Smaller volumes are technically
	difficult to reliably measure in a 1 cc syringe and amounts less than this volume are uncertain in the
	amount actually delivered in the eye. Lower dose of 0.12 mg has been reported to be efficacious in
	CARE-ROP trial, but for the reasons mentioned above, it is a small volume to work with.
Evidence	Background
	Vascular endothelial growth factor (VEGF) - A key regulator of angiogenesis in foetal life. In the
	normally developing retina, VEGF leads to the development of blood vessels from the optic nerve to
	the periphery. In preterm infants with disrupted angiogenesis, however, the expression and levels of
	VEGF differ markedly in the two different phases. While the levels are suppressed in the vaso-
	obliterative phase, there is an overproduction/expression of VEGF, leading to abnormal vascular
	proliferation in the vaso-proliferative phase.(2)

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Type 1 ROP: Retinal findings defined as type 1 ROP are: (1) Zone I ROP: any stage with plus disease; (2) Zone I ROP: stage 3 - no plus disease, (3) Zone II ROP: stage 2 or stage 3 with plus disease.(2) **Efficacy**

Anti-VEGF for type 1 ROP: Sankar et al 2018, in Cochrane systematic review, evaluated the efficacy or safety of anti-VEGF agents compared with laser/cryotherapy in type 1 ROP. (2) Six randomised or quasi-randomised controlled trials involving 383 infants were included. Four trials compared intravitreal bevacizumab monotherapy with conventional laser therapy (3-7). One trial compared ranibizumab monotherapy with laser therapy (8) and one study compared intravitreal pegaptanib plus conventional laser therapy with laser and cryotherapy (9). When used as monotherapy, bevacizumab/ranibizumab did not reduce the risk of retinal detachment, pre-discharge mortality, corneal opacity requiring corneal transplant, or lens opacity requiring cataract removal. The risk of retreatment also did not differ between groups. Subgroup analysis showed a significant reduction in the risk of recurrence in infants with zone I ROP (RR 0.15, 95% CI 0.04 to 0.62), but an increased risk of recurrence in infants with zone II ROP (RR 2.53, 95% CI 1.01 to 6.32). There was a significant increase in the risk of recurrence of ROP in eyes that received bevacizumab (RR 5.36, 95% CI 1.22 to 23.50; RD 0.10, 95% CI 0.03 to 0.17). Infants who received intravitreal bevacizumab had a significantly lower risk of refractive errors at 30 months of age. No trial included in this meta-analysis reported neurodevelopmental outcomes.(2)

Li et al 2018, in their meta-analysis, compared the efficacy of anti-VEGF and laser treatments in type-1 and threshold ROP.(10) This study included 4 RCTs and 6 comparative non-randomised studies (CNS) involving 1158 patients. Retreatment incidence was significantly increased in anti-VEGF (OR 2.52; 95% CI 1.37 to 4.66; P = 0.003) compared to the laser treatment. Retreatment incidence was 6.8-21.4% and 1.4-14% in Anti-VEGF and laser groups respectively. Average time interval between initial treatment and retreatment was 7.5 weeks (95% CI 2.00, 17.08 weeks). The longest retreatment time was 17 weeks (postmenstrual age not more than 57 weeks). While the retreatment incidence was higher, anti-VEGF treatment was safer, with a relatively reduced incidence (OR 0.29; 95% CI 0.10 to 0.82; P = 0.02) of eye complications (corneal opacity, cataract, preretinal or intravitreal haemorrhage and retinal detachment). There was less myopia in comparison to laser therapy (WMD 3.03D; 95% CI 1.48 to 4.59; p=0.0001).(10)

A descriptive review by American Academy of Ophthalmology in 2018 analysed 5 RCTs and 7 comparative non-randomised case series found that intravitreal anti-VEGF therapy is as effective as laser photocoagulation for achieving regression of acute ROP. But, ROP recurrence rate was higher, indicating a vigilant and extended follow-up. (11)

Anti-VEGF preparations and doses for ROP: Of 14 studies (RCTs and comparative non-randomised studies), (4-9, 12-19), 12 studies (5 of them RCTs) evaluated bevacizumab, 2 studies evaluated ranibizumab and 1 study trialled pegaptanib. RCTs evaluating bevacizumab used 0.5 mg to 1.25 mg (Beat-ROP trial and Karkhaneh et al – 0.625 mg in 0.025 mL; Lepore et al – 0.5 mg in 0.02 mL; O'Keefe et al and Moran et al – 1.25 mg in 0.1 mL). Zhang et al in their RCT used 0.3 mg in 0.03 mL of ranibizumab.

Author	Study	Anti-VEGF	Dose
CARE-ROP trial (20)	RCT	Ranibizumab	0.12 mg versus 0.2 mg
Beat-ROP trial 2011 (6)	RCT	Bevacizumab	0.625 mg in 0.025 mL
Karkhaneh 2016 (4)	RCT	Bevacizumab	0.625 mg in 0.025 mL
Lepore 2014 (5)	RCT	Bevacizumab	0.5 mg in 0.02 mL
O'Keefe 2016 (7)	RCT	Bevacizumab	1.25 mg in 0.05 mL
Moran 2014 (16)	RCT	Bevacizumab	1.25 mg in 0.1 mL
Harder 2013 (13)	Case series	Bevacizumab	0.375 mg – 0.625 mg
Isaac 2015 (15)	Case series	Bevacizumab	0.625 mg in 0.025 mL
Hwang 2015 (14)	Case series	Bevacizumab	0.625 mg in 0.025 mL
Mueller 2016 (17)		Bevacizumab	0.625 mg in 0.025 mL
Lee 2010 (19)	Case series	Bevacizumab	0.5 mg in 0.02 mL
		plus laser	

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Walz 2016 (18)	Case series	Bevacizumah	Dose not reported	
Zhang 2016 (8)	RCT	Ranibizumab	0.3 mg in 0.03 mL	
Gunay 2017 (12)	Case series	Bevacizumab	0.625 mg in 0.025 mL	
		Ranibizumab	0.25 mg in 0.025 mL	
Summary on short term ef	fficacy of anti-VEG	F for primary treatn	nent of type 1 ROP: Anti-VEGF	is
probably as efficacious as l	aser treatment for	the acute type 1 RC	OP but the necessity for retreat	ment is
higher (up to 21.4%) in cor	nparison to laser th	nerapy (up to 14%).	Incidence of eye complications	and
myopia were less in anti-V	EGF group.			
Ranibizumab dose compa	risons: CARE-ROP s	tudy group perform	ied a multi-centre, double blin	d (20
randomised controlled trai	I to compare 2 dos	es of ranibizumab (0.12 mg vs 0.2 mg) in 19 infant	S (38
treatment Rescue therany	was defined as the	eeu ioi rescue tileid e need for either las	er or anti-VEGE within 28 days	of
initial treatment or laser tr	eatment at any tim	e Other outcomes	was retreatment with the same	e anti-
VEGF dose if ROP activity r	eappeared after 28	davs of initial treat	ment. When analysed per eve	. 94.4%
of eyes in the 0.12-mg grou	up and 92.9% in the	e 0.20-mg group rea	iched 24 weeks post-treatmen	t
without need for rescue th	erapy. Two (5.2%)	eyes required rescu	e therapy with full resolution.	
Recurrence of any ROP sta	ge was more preva	lent in the 0.20 mg	group. Two infants in each gro	up (8
eyes [21.1%]) had recurrer	ices that were seve	ere enough to warra	nt retreatment. Eleven eyes (5	5.0%)
had full physiologic vascula	arization in the 0.12	2-mg group, and onl	y 3 eyes (16.7%) achieved full	
vascularization in the 0.20-	mg group suggesti	ng that higher anti-	VEGF doses may impede physi	ologic
vascularization. Free plasm	a VEGF levels were	e measured before (baseline) and during the first s	ix weeks
after ranibizumab injection	1. Several VEGF leve	els were below dete	ection limit at baseline (i.e. bet	ore
injection in either group (2	ol	ed suppression of m	lean vege levels after rambizur	nad
injection in either group.(2	0)			
Ranibizumab in zone II RO	P: In a randomised	controlled trial. Zha	ang et al compared 0.3 mg of	
ranibizumab and laser ther	apy for zone II RO	P. A substantial prop	portion of infants developed	
recurrence of ROP after rai	nibizumab in comp	arison to laser thera	apy (52% versus 4%). (21)	
Topical antibiotics during	and following intra	vitreal injections:	A descriptive review of adult st	udies
did not find firm evidence	supporting benefit	for topical antibioti	c prophylaxis	
for post-injection endopht	halmitis and may c	arry harmful effects	with possibly higher endopht	nalmitis
rates, and increasing antip	Iotic resistance. Fir	m evidence is lackin	ig for neonates to recommend	or
refute the children practice.	(22)			
Topical antisepsis (povidor	e-iodine): A prosp	ective randomised e	evaluation in adults of topical a	ntibiotic
plus povidone-iodine versu	is povidone-iodine	alone showed that	patients undergoing regular	
intravitreal injections, the	rate of positive bac	cterial cultures was 8	8% in the group that	
received a three-day cours	e of pre-injection t	opical gatifloxacin ir	n addition to povidone–	
iodine immediately prior to	o injection, compar	ed with just 4% in t	he group that received povido	ne-
iodine alone (p = 0.32).(23)				
Safety				
Concerns remain regarding	the potential long	term local and syst	emic adverse effects of anti-Vi	EGF.
Local eye complications:	here was no signifi	cant difference in th	ne incidence of corneal opacity	with
the incidence of esteract in	was not round in a	z studies (4, 8) and c	one study did not find any diffe	erence in
haemorrhage were report	ad in 2 studies and	did not find these of	α	
Systemic absorption and s		VEGE is an importa-	nt neurodevelonmental growt	h factor
in the early newborn perio	d. Wu et al. in a pr	ospective cohort stu	idy, measured serum VEGF lev	els for
up to 12 weeks after intrav	vitreal ranibizumab	(0.25 mg) or bevaci	izumab (0.625 mg) in infants w	ith ROP.
Serum VEGF level significat	ntly decreased betw	ween baseline and u	up to 8 weeks in bevacizumab	group (P
= 0.007). There was no sigr	nificant difference i	in the serum VEGF le	evel between baseline and up	to 8
weeks in ranibizumab grou	ip (P = 0.212).(24)			

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	Neurodevelopmental outcomes: A study from the Canadian Neonatal Network demonstrated 3.1
	times higher odds (95% Cl 1.2 to 8.4) of severe neurodevelopmental disabilities in preterm infants born
	before 29 weeks' gestation and treated with bevacizumab, after adjusting for key confounders like
	gestation, gender, maternal education, Score for Neonatal Acute Physiology-II (SNAP-II) score,
	bronchopulmonary dysplasia, sepsis, and severe brain injury. (25) However, this comparison was
	adjusted for many infant variables but not ROP severity, and there was a significantly greater
	proportion of patients with zone I disease in the bevacizumab group. A retrospective study published
	by the National Institute of Child Health and Human Development (NICHD) Neonatal Research Network
	involved 405 preterm infants <27 weeks gestation who were treated with either surgery or
	bevacizumab for ROP. Primary outcome was the composite of death or neurodevelopmental
	impairment. Composite primary outcome did not differ between the groups but the odds of death (aUR
	2.54 [95% Cl 1.42 to 4.55]; P = .002), a cognitive score <85 (aUR 1.78 [95% Cl 1.09 to 2.91]; P = .02), and a Crass Mater Europhic Classification Scole level >2 (aOP 1.78 [95% Cl 1.04 to 2.91]; P = .02), and
	a Gross Motor Functional Classification Scale level 22 (aUR 1.73 [95% CI 1.04 to 2.88]; $P = .04$) were
	significantly higher with bevacizumab (0.625 mg) and laser therapy for POP, compared with a
	birthweight, and gestational age matched control group of children who had received laser treatment
	for ROP. They found no difference in the mean cognitive language or motor scores on the RSID III test
	(27) Lien et al studied BSID scores at 24 months of age in 61 infants who had received either
	bevacizumab (0.625 mg) monotherapy, laser monotherapy, or a combination of bevacizumab and laser
	therapy (required for salvage therapy). The patients who required combination (salvage) therapy had a
	higher incidence of mental or psychomotor impairment, but there was no difference between the
	groups that had either modality as monotherapy.(28)
	Pharmacokinetics
	Pharmacokinetic data in adults with macular degeneration estimate that vitreous half-life of
	ranibizumab is about 9 days and on reaching the systemic circulation, ranibizumab has a short half-life
	of 2 hours. The systemic-to-vitreous exposure ratio for ranibizumab was estimated to be 1:90,000. The
	steady-state serum concentrations of total ranibizumab were at all times below the concentrations
	needed to reduce VEGF-A-induced endothelial cell proliferation in vitro by 50%.(29)
Practice points	
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