## Sildenafil Newborn Use Only

are needed. honary Hypertension of the Neonate (PPHN): ctory to inhaled nitric oxide (iNO) and other conventional therapies or e who are persistently unable to be weaned off inhaled nitric oxide or uations where inhaled nitric oxide and high frequency ventilation are not available nary hypertension secondary to respiratory, cardiac or chest wall disease. bhodiesterase type 5 (PDE5) inhibitor. PDE5 is found in the smooth muscle of the culature, where it is responsible for the degradation of cyclic guanosine te (cGMP). cGMP produces smooth muscle relaxation. Sildenafil increases cGMP ary vascular smooth muscle cells resulting in relaxation. In patients with pulmonary this can lead to selective vasodilatation of the pulmonary vascular bed and, to a
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vasodilatation in the systemic circulation.
rase type 5 (PDE5) inhibitor.
r prepared
ction containing <b>10 mg/12.5 mL</b> = 0.8 mg/mL of sildenafil
<i>r</i> -prepared oral suspension
g/kg administered <b>over THREE hours</b> followed by:
1.6 mg/kg/day (0.067 mg/kg/hour) as a continuous infusion for up to 7 days.
I mg/kg/dose given 6 to 8 hourly and titrate up to 2 mg/kg/dose according to
p to maximum of 3 mg/kg/dose given 6 hourly.
ossible occurrence of sudden clinical deterioration during withdrawal of sildenafil, a
eduction should be considered when stopping sildenafil.
tion IV infusion (weight > 2.5 kg)
/kg (2 mg/kg of sildenafil) solution and make up to 15 mL using glucose 5%
sodium chloride 0.9%.
or <b>3 hours</b> (loading dose of 0.4 mg/kg) <b>followed by</b> 0.5 mL/h (0.067 mg/kg/h)
ation IV Infusion (weight $\leq 2.5$ kg) (by (2.26 ms (line foil)) solution and use here to 15 mb using shares 5%
/kg (3.36 mg /kg of sildenafil) solution and make up to 15 mL using glucose 5% sodium chloride 0.9%.
n for <b>3 hours</b> (loading dose of 0.4 mg/kg) <b>followed by</b> 0.3 mL/h (0.067 mg/kg/h)
pre drawing up the dose. Give via intragastric tube, preferably with feed to
f gastrointestinal irritation. If baby is not on enteral feeds or breast milk is not
dose via intragastric tube and flush with 0.5 mL water for injection.
od pressure and oxygenation.
atic function.
toring with echocardiogram.

Neonatal Medicines Formulary Consensus GroupSildenafilPage 1 of 5This RHW document is a modification of Neomed version. Dosage schedules remain the same. However, information on the<br/>commercial preparations not used at RHW might have been deleted. The risk rating might have been modified as per the local<br/>health district policy.

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ContraindicationsHypersensitivity to sildenafilNot to be used in patients taking organic nitrates of any form e.g. glyceryl trinitrate, it	
i not to be used in datients taking organic nitrates of any form e.g. givcervi trinitrate. I	sosorbide
mononitrate, sodium nitroprusside	
<b>Precautions</b> Use with caution in neonates with sepsis or uncontrolled hypotension.	
Sildenafil clearance (in adults) is reduced in hepatic and severe renal impairment.	
Drug Interactions Sildenafil metabolism is principally mediated by the cytochrome P450 (CYP) isoforms	3A4 (major
route) and 2C9 (minor route). Inhibitors of these isoenzymes may reduce sildenafil clo	earance and
inducers of these isoenzymes may increase sildenafil clearance. Thus, erythromycin a	nd
fluconazole may increase concentrations of sildenafil by reducing hepatic clearance a	nd rifampicin
may decrease concentrations by inducing its hepatic metabolism.	
Avoid concomitant use of sildenafil with: Alprostadil (prostaglandin E1), other antihy	pertensives
and vasodilators, as they may have their effects potentiated by sildenafil.	
Adverse Reactions Most concerning short-term adverse effects: Worsening oxygenation and systemic hy	-
Epistaxis, respiratory symptoms (cough and nasal congestion), diarrhoea and vomitin	-
gastroesophageal reflux and abdominal pain, headaches, tremors, erections, facial flu	-
dizziness, irritability and (rarely) fever, skin disorders, pain in limbs and oedema have	
reported in children on sildenafil. The Sildenafil in Treatment-Naïve Children, Aged 1-	
With Pulmonary Arterial Hypertension long-term extension (STARTS-2) trial showed v	
in children receiving high doses of sildenafil as monotherapy. <sup>2</sup> A recent study conduct	-
and colleagues, found there was a statistically significant increase in adverse drug rea	
frequency in children receiving higher-than-recommended doses. However, it was no with a lower survival rate. <sup>14</sup>	associated
Sildenafil has the potential to adversely affect vision. <sup>13</sup>	
Impaired liver function tests.	
May increase the risk of severe retinopathy of prematurity if used in extremely preter	rm neonates
<b>Compatibility</b> Glucose 5%, sodium chloride 0.9%.	
Incompatibility No data – where possible administer via dedicated line.	
Stability       IV – infusion should be changed every 24 hours.         Oral suspension – as per pharmacy advice.	
StorageIV – unopened vials at room temperature (20–25°C).Oral suspension – refrigerate, do not freeze	
Special Comments         In paediatric patients with pulmonary arterial hypertension, an increased mortality right	sk was
associated with long-term (> 2 year) use. The mortality risk of long-term use in neona	
unknown.	
Evidence summary Neonates with pulmonary hypertension	
Shah et al <sup>9</sup> performed a systematic review on sildenafil compared with placebo or oth	ner
pulmonary vasodilators, irrespective of dose, route and duration of administration, ir	n neonates
with PPHN. Three eligible trials that enrolled 77 infants were identified. The methodo	ological
quality of the studies indicated low-moderate risk of bias. All studies were performed	in resource-
limited settings where iNO and high frequency ventilation were not available at the ti	-
There was a significant reduction in mortality in the sildenafil group (typical RR 0.20, s	
to 0.57; typical RD -0.38, 95% CI -0.60 to -0.16; Number needed to treat to benefit 3,	
6). Physiological parameters of oxygenation (oxygenation index, PaO <sub>2</sub> ) suggested a st	-
improvement after the first dose of sildenafil. No clinically important side effects wer	
Sildenafil in the treatment of PPHN has significant potential especially in resource lim	ited settings
(LOEI, GOR B).	+ 2016: 0
The European Paediatric Pulmonary Vascular Disease Network's consensus statement of PDHN and PH in PDD, especially if iNC	
sildenafil should be considered for treatment of PPHN and PH in BPD, especially if iNC	
available (LOE IIa GOR B). Intravenous sildenafil may be considered for treatment of F PPHN, in critically ill patients, especially in those with an unsatisfactory response to in	-
GOR B). <sup>4,5</sup>	
Paediatric pulmonary hypertension	

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STARTS-1 trial performed by Barst et al, <sup>1</sup> studied the effectiveness of oral sildenafil in children with pulmonary arterial hypertension. 238 children with a weight $\geq$ 8 kg were randomised to low-, medium-, or high-dose sildenafil or placebo orally 3 times daily for 16 weeks. The primary comparison was percent change from baseline in peak oxygen consumption for the 3 sildenafil doses combined versus placebo. Percent change in PV' O <sub>2</sub> for the 3 sildenafil doses combined was only marginally significant; however, PV' O <sub>2</sub> , functional class and haemodynamic improvements with medium and high doses suggest efficacy with these doses.
STARTS-2 was the extension of the STARTS-1 trial. <sup>2</sup> In STARTS-2, sildenafil-treated patients continued STARTS-1 dosing; placebo-treated patients were randomised to 1 of the 3 sildenafil dose groups. Patients requiring additional pulmonary arterial hypertension–specific therapy discontinued study treatment; survival follow-up was attempted. Hazard ratios for mortality were 3.95 (95% confidence interval, 1.46–10.65) for high versus low and 1.92 (95% confidence interval, 0.65–5.65) for medium versus low dose; however, multiple analyses raised uncertainty about the survival/dose relationship. In summary, although children randomised to higher compared with lower sildenafil doses had an unexplained increased mortality, all sildenafil dose groups displayed favourable survival for children with pulmonary arterial hypertension. Combined with STARTS-1 data, the overall profile favoured the medium dose.
<b>Preterm infants at risk of BPD</b> Konig et al <sup>6</sup> performed an RCT in preterm infants, < 28 weeks gestational age, if they were mechanically ventilated on day 7 of life. Infants were randomised to a 4-week course of either oral sildenafil (3 mg/kg/day) or placebo solution. Twenty infants were randomised, 10 received sildenafil and 10 received placebo. Sildenafil did not reduce length of invasive (median 688 versus 227 h) or non-invasive ventilation (median 1609 versus 1416 h). More infants in the sildenafil group required postnatal steroid treatment. One infant developed hypotension following sildenafil administration and was excluded after three doses. Conclusion: Sildenafil as an early treatment for preterm infants at risk of BPD is not beneficial.
<b>Prevention of rebound pulmonary HTN after weaning iNO</b> Namachivayam et al <sup>8</sup> performed an RCT in 30 ventilated infants and children from varying respiratory conditions including BPD (average age 0.4 y) and receiving 10 ppm iNO. They were randomised to either 0.4 mg/kg as a single oral dose of sildenafil or placebo 1 h before discontinuing iNO. Rebound occurred in 10/14 of the placebo group and 0 out of 15 in the sildenafil group. Four placebo patients couldn't be weaned off iNO, whereas all in the sildenafil group were successfully weaned (p = 0.042). A single oral dose of sildenafil may be considered for this particular scenario (LOE II, GOR C).
<b>Pre-op oral sildenafil for children with CHD prior to cardiopulmonary bypass</b> Vassalos et al, <sup>12</sup> in a randomised trial, compared the effects of oral sildenafil (0.5 mg/kg) and placebo, administered the day before cardiac surgery, in 24 children. Postoperatively, mean pulmonary vascular resistance and oxygenation index remained unchanged, whilst oxygen delivery and bi-ventricular systolic function were significantly reduced in the sildenafil group. In this trial, pre-operative sildenafil did not affect postoperative pulmonary vascular resistance. There was, however, a negative impact on ventricular function and oxygenation. Therefore, sildenafil is not recommended for this particular indication.
<b>Post-op sildenafil in infants after cardiac surgery</b> Stocker et al <sup>11</sup> performed an RCT in 16 ventilated infants early after closure of ventricular or atrioventricular septal defects. They were randomly assigned to one of two groups. Seven infants received iNO (20 ppm) first, with the addition of intravenous sildenafil (0.35 mg/kg over 20 min) after 20 min. Eight infants received sildenafil first, iNO was added after 20 min. Intravenous sildenafil augmented the pulmonary vasodilator effects of iNO in infants early after cardiac surgery. However, sildenafil produced systemic hypotension and impaired oxygenation, which was

<ul> <li>not improved by INO. Siltenafil is not recommended for this particular indication.</li> <li>The European Paediatric Pulmonary Vascular Disease Network Consensus 2016<sup>5</sup>: Beneficial haemodynamic effects of sildenafil have also been demonstrated in failing Fontan circulations.</li> <li>Sildenafil improved max. oxygen consumption IVO; max) and pulmonary biochafil therapy improved excise tolerance and ventilatory efficiency in Fontan patients.</li> <li>Pharmacokinetics</li> <li>Steinhorn et al<sup>10</sup> performed a multicentre, open-label, dose-esolation, pilot, pharmacokinetics study of 36 near-term and term neonates with echo-confirmed idiopathic PHN or secondary PHN with MAS, RDS, sepsis or pneumonia and O1: 15. The study included 8 sequential "step-up" dosing groups. Nost infants received a loading dose of sildenafil to bring the plasma concentration of sildenafil to a target, followed by a maintenance infusion for the remainder of the study. The duration of IV sildenafil was for at least 48 hours and up to 7 days. They found sildenafil to be well tolerated, particularly with a dosing regimen comprising a loading dose of 0.4 mg/kg delivered over 3 hours, followed by a maintenance infusion of 1.6 mg/kg/day.</li> <li>Sildenafil and retinopathy of prematurity</li> <li>Sildenafil and retinopathy of prematurity</li> <li>Sildenafil is 10 x more selective for PDE5 as compared to PDE6. PDE6 is found in the retina. If used in extremely preterm meonates, sildenafil may increase the risk of severe retinopathy of prematurity. A retrospective, case-controlled study. Garcia Ac, et al. A randomized, double-blind, placebo-controlled, dose-ranging study of oral sildenafil increase the risk of retrospective, case-target induce the side charance induces with pulmonary arterial hypertension. Expert consensus statement on the diagnosis and treatment of paediatric pulmonary hypertension. Expert consensus statement on the diagnosis and tretentment of paediatric pulmonary hypertension. Evergen Paedia</li></ul>				
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Neonatal Medicines Formulary Consensus GroupSildenafilPage 4 of 5This RHW document is a modification of Neomed version. Dosage schedules remain the same. However, information on the<br/>commercial preparations not used at RHW might have been deleted. The risk rating might have been modified as per the local<br/>health district policy.

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