

A prescription-screening tool to optimise pharmacotherapy for hospitalised neonates

		els of evidence	Gra	des of recommendation
	1++	High quality meta-analyses, systematic reviews of RCTs, or RCTs with a very low risk of bias	A	At least one meta-analysis, systematic review, or RCT rated as 1++ and directly applicable to the target
	1+	Well conducted meta-analyses, systematic reviews of RCTs, or RCTs with a low risk of bias	A	population OR A systematic review of RCTs or a body of evidence
	1-	Meta-analyses, systematic reviews or RCTs, or RCTs with a high risk of bias		
GOR scale	2++	High quality systematic reviews of case-control or cohort studies or High quality case-control or cohort studies with a very low risk of confounding, bias, or chance and a high probability that the relationship is causal	В	A body of evidence including studies rated as 2++ directly applicable to the target population and demonstrating overall consistency of results OR Extrapolated evidence from studies rated as 1++ or 1+
	2+	Well conducted case-control or cohort studies with a low risk of confounding, bias, or chance and a moderate probability that the relationship is causal	С	A body of evidence including studies rated as 2+ directly applicable to the target population and demonstrating overall consistency of results or Extrapolated evidence from studies rated as 2++
	2-	Case-control or cohort studies with a high risk of confounding, bias, or chance and a significant risk that the relationship is not causal		
	3	Non-analytic studies, eg case reports, case series	D	Evidence level 3 or 4 OR
	4	Expert opinion		Evidence level 3 of 4 of C

Item design

	Main statement
Start / stop /	Complementary informations
other	Grade of recommendations:
	References
Age/weight	Category/Subcategory
	Drua



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	Start reducing the percentage of parenteral nutrition as quickly as possible by the introduction of enteral nutrition until enteral nutrition	
8	finally replaces completely PN in order to minimise any side-effects from exposure to PN.	4
9	Start continuous parenteral glucose administration in preterm infants needing parenteral nutrition.	5
10	Start amino acid supply in the first day of life in preterm infant needing parenteral nutrition.	6
	Start continuous lipid emulsion infusion within the first 24-48 hours of life in preterm infant needing parenteral nutrition,	7
	Do not administer parenteral lipid emulsion at a dose higher than 3-4 g/kg/day in neonates.	7
	Check that a sufficient quantity of linoleic acid is administered in all neonates on parenteral nutrition.	8
	Start electrolytes supplementation with parenteral nutrition after onset of diuresis.	8
	Start vitamins and trace elements supplementation in neonates receiving parenteral nutrition.	ç
	Start vitamin D supplementation from the first days of life in all neonates.	ç
_	DIOLOGY	10
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	Start prostaglandin E1 (alprostadil) as an initial continuous intravenous infusion at 0.01 mcg/kg/min, until a definitive diagnosis is made in	10
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	Reassess the indication of prostaglandin E1 (PGE1) treatment.	10
	Stop ibuprofen, indomethacin and paracetamol in patients with duct dependent congenital heart disease.	10
		11
ale	nt Ductus Arteriosus (PDA) Consider pharmacological closure of confirmed patent ductus arteriosus (PDA) in preterm neonates after 2 weeks of life, with ibuprofen as	- 11
20	first-line treatment.	11
21	Reassess the indication of ibuprofen, indomethacin and paracetamol in preterm neonates <2 weeks of life with confirmed or unconfirmed	11
22	patent ductus arteriosus (PDA).	40
	Reassess the indication of ibuprofen, indomethacin and paracetamol in term neonates with patent ductus arteriosus (PDA).	12
	Do not use paracetamol as first-line treatment for patent ductus arteriosus (PDA) closure. Consider a switch to ibuprofen.	12
	tension	13
24	Do not use volume expansion as first line treatment in VLBW infants (BW <1500g) with hypotension.	13
25	Consider a conservative approach (permissive hypotension) for the managment of VLBW infants (BW <1500g) if the clinical examination is	13
	satisfactory in the face of apparent hypotension	
	ATOLOGY	14
\ner		14
26	Do not use routinely erythropoietin to limit exposure to blood transfusions in preterm neonates. The indication of treatment should be	14
	reassessed.	
	Start iron supplement of 2-3 mg/kg/day in all preterm infants fed human milk once full oral feeds have been achieved.	14
	ulation disorders	15
	Start oral Vitamin K in neonates breastfed by a mother treated with phenprocoumone.	15
	Check in all neonates that a complete Vitamin K prophylaxis has been given at birth.	15
_	mbocytopenia and Platelet Dysfunction	16
30	Consider platelets transfusion even in the absence of bleeding in all neonates with a platelet count of <30x109/L.	16
24	Consider platelets transfusion in neonates with a platelet count of 30-49x109/L and minor bleeding or those at risk for major bleeding.	45
31	Consider platelets transfusion in neonates with a platelet count of 50-49x 109/L and million bleeding of those at risk for major bleeding.	17
32	Consider platelets transfusion in neonates with a platelet of count 50-99x109/L only if bleeding is present.	17
33	Do not transfuse neonates with mild thrombocytopenia (platelet count 100-150x109/L) even if bleeding.	18
	Start intravenous immunoglobulin (IVIG) only in case of severe thrombocytopenia (platelet count of <50x109/L) or if bleeding persists	
34	despite compatible platelets transfusion or in combination with unmatched platelets transfusion in neonates with neonatal allo-immune	18
	thrombocytopenia (NAIT).	
	Start intravenous immunoglobulin (IVIG) as first line treatment in neonates with neonatal auto-immune thrombocytopenia and born to	
35	mothers who have idiopathic thrombocytopenic purpura (ITP), when platelet count is of <30x109/L or clinical bleeding is present.	19
	spasms and Thromboembolism	20
36	Start unfractionned heparin or low molecular weight heparin in neonates with a first event venous thromboembolism for at least 5 days.	20
	Start alteplase or urokinase only in case of major vessel occlusion causing critical compromise of organs or limbs in infants with venous	
37	thromboembolism.	20
	unombodinumini.	





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oneu		21
	ımothorax	21
38	Do not use routine supplemental oxygen use in infants with spontaneous pneumothorax.	21
	ea of Prematurity	22
	Start caffeine citrate in patients with apnea of prematurity (loading dose 20 mg/kg; maintenance dose 5mg/kg/day). Dose may be	00
- 4u i	increased to 10 mg/kg/day if apnea persists.	22
	Reassess the need for caffeine citrate treatment.	23
	Reassess the indication of anti-gastroesophageal reflux therapy in neonates with apnea.	23
	chopulmonary Dysplasia (BPD)	24
	Do not use dexamethasone in the prevention or the treatment of bronchopulmonary dysplasia.	24
	Do not use loop diuretics for prevention of BPD in preterm neonates.	25
	Do not use thiazid diuretics for prevention of BPD in preterm neonates. Use them judiciously for treatment of BPD in preterm neonates.	25
	piratory Distress Syndrome (= Hyaline Membrane Disease)	26
	Start surfactant therapy in infants born <26 weeks of gestational age who need FiO2 >0.30.	26
	Start surfactant therapy in infants born ≥26 weeks of gestational age who need FiO2 >0.40.	26
	onium Aspiration Syndrome	27
	Consider inhaled nitric oxide (iNO) in neonates with hypoxic respiratory failure due to MAS.	27
	Reassess the indication for antibiotics in patients with MAS alone.	27
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	istent Pulmonary Hypertension of the Newborn (PPHN)	28
	Start inhaled nitric oxide (iNO) in neonates who have severe PPHN.	28
	Do not use sildenafil as initial therapy for PPHN.	28
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	e Kidney Injury (AKI)	29
	Do not use nephrotoxic drugs in neonates if possible, especially in preterm infants.	29
	Stop all nephrotoxic drugs when possible in neonates with AKI (stage 1-3).	30
	Consider dosage adjustement for drugs highly excreted by renal elimination in neonates with AKI (stage 1-3). When needed, refer to a	
	specialist.	30
	TROENTEROLOGY	31
	ct hyperbilirubinemia (Conjugated Hyperbilirubinemia)	31
	Decrease or stop intravenous soybean-based lipid emulsion in neonates with marked progressive cholestasis associated with parenteral	
าวา แ	nutrition.	31
	Administer adequate protein intake of 2 to 3 g/kg/day to neonates with direct hyperbilirubinemia.	32
	Start fat-soluble vitamins (ADEK) in neonates with cholestasis.	32
	Consider ursodeoxycholic acid (UDCA) in neonate with direct hyperbilirubinemia.	33
	ect hyperbilirubinemia (Unconjugated Hyperbilirubinemia)	34
	Administer intravenous immunoglobulin (IVIG) to neonates with a positive direct Coombs test and severe hyperbilirubinemia, or to those	J4
	progressing to severe hyperbilirubinemia despite initial treatment.	34
	otizing Enterocolitis (NEC)	35
	Start probiotics in preterm neonates at high risk of developing NEC.	35
	Start probletics in preterm rechates at high risk of developing NEC. Stop all enteral medications in neonates suspected to have NEC.	36
	Do not use enteral antibiotics for the prevention of NEC.	36
	Start broad spectrum antibiotic promptly after blood cultures have been drawn in neonates with any stage of NEC.	36
	rointestinal Bleeding from the Upper Tract	
	-	37
64	Check that a Vitamin K prophylaxis was administered postdelivery in neonates with upper gastro-intestinal bleeding, to guide diagnostic.	37
Gast	roesophageal Reflux	38
- 1	Consider proton pump inhibitors or H2-blockers only in neonates with severe cases of acid gastroesophageal reflux disease (GERD),	20
	when non-pharmacological measures (including milk thickeners) have failed.	38





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Seiz	ures	40
67	Start phenobarbital as the first line agent in neonates with either EEG diagnosed or clinically apparent seizures when prolonged or frequent.	40
68	Consider phenytoin or a benzodiazepine or lidocaine in neonates with persistant seizures, despite adequate phenobarbital treatment.	41
69	Stop antiepileptic drugs if seizure-free for >72 hours in neonates with normal neurological examination and/or normal electroencephalography.	41
70	Consider pyridoxine only in neonates with recurrent seizures with no obvious cause.	41
	, SEDATION & NEONATAL ABSTINENCE SYNDROME	42
	Analgesia & Sedation	42
	Start pain management in neonates with non-pharmacological techniques (incl. Sucrose) if aproppriate.	42
72	Start paracetamol in neonates who are still in pain despite adequate non-pharmacological interventions.	42
73	Do not use nonsteroidal antiinflamatory agents (NSAID) as analgesics.	43
74	Start morphine as first line treatment for pain relief in neonates who are still in pain despite adequate non-pharmacological techniques and paracetamol treatment.	43
75	Start opioids as first line treatment for postoperative analgesia, and use them as long as pain assessment scales deem necessary.	44
76	Reassess the indication of morphine or fentanyl in chronically ventilated preterm neonates without pain.	45
77	Do not use ketamine treatment for routine management of pain.	45
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78	Consider non-pharmacological interventions for the initial management of all infants suspected of having or at risk of developing NAS. This may mitigate the need for medication.	46
79	Start morphine as the first line pharmacological treatment for NAS when opioids are used by the mother and supportive measures failed.	46
80	Start weaning of morphine as soon as Modified Finnegan scores are <8 for 24 to 48 hours in neonates with NAS.	47
81	Do not use morphine in neonates with NAS when the drugs used by the mother are non-opioids.	47
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82	Start empirical antibiotic treatment with high dose amoxicilline and gentamicin in neonates with diagnosed or strongly suspected meningitis.	48
83	Check results of cerebro-spinal fluid (CSF) culture as soon as they are available in order to reassess the need for treatment or the choice of antibiotics in neonates with suspected meningitis treated with empirical antibiotics.	48
84	Do not use corticosteroids for the treatment of neonates with suspected or confirmed bacterial meningitis. Reassess the corticosteroid indication.	49
Seps		50
85	Do not use empirical antibiotic therapy for asymptomatic neonates with a single risk factor of infection (incl. mother with suspected chorioamnionitis or unexplained premature delivery).	51
86	Start empirical antibiotic treatment after blood cultures have been drawn in all newborn infants with suggestive signs of neonatal infection.	51
87	Reassess the need for antibiotics after 48 hours in neonates treated empirically with antibiotics for suspected sepsis.	52
88	Do not use cephalosporins as first-line treatment in infant with suspected neonatal infection, because of the high risk of developing resistance. Use is restricted to special cases	52
89	Do not use intravenous immunoglobulin in the treatment of suspected or proven neonatal sepsis.	53
	Do not use Vancomycin as prophylaxis against sepsis in preterm neonates.	53
Нера		54
91	Administer an initial dose of hepatitis-B vaccine within 12 hours of birth in infants born to HBsAg-positive mothers, including infants weighing <2000g. Administer Hepatitis-B immune globulins (HBIG) 200 IU concurrently but at a different anatomic site.	54
92	Do not use early hepatitis-B vaccine in infants born to mothers whose HBsAg and HBeAg status is negative but with positive anti-HBs status (prior infection or at risk of infection).	54
Hum	an Immunodeficiency Virus (HIV)	55
	Start HIV prophylaxis with zidovudine as close to birth as possible for at least 4 weeks or consider tritherapy in neonates born to HIV-	
93	infected mothers who did not follow proper antenatal treatment or whose viremia are detectable.	55
94	Start tritherapy immediately in the neonate aged <72 hr if the mother is diagnosed postpartum with HIV infection.	55





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95	Start respiratory syncytial virus prophylaxis with palivizumab in neonates with severe bronchopulmonary dysplasia (BPD).	56
96	Start respiratory syncytial virus prophylaxis with palivizumab in neonates with haemodynamically significant congenital heart disease AND other associated risk factors.	56
97	Dot not use respiratory syncytial virus prophylaxis with palivizumab routinely in preterm neonates.	57
98	Do not use palivizumab for the treatment of respiratory syncytial virus (RSV) infection. Stop the treatment, even if it was given before the infection.	57
Toxo	plasmosis	58
99	Administer a combination of pyrimethamine-sulfadiazine-folinic acid during the first year of life to neonates in whom a diagnosis of congenital toxoplasmosis is confirmed or probable.	58
100	Do not use spiramycin in neonates. Stop treatment and screen for potential QT interval prolongation.	58
	megalovirus (CMV)	59
101	Start antiviral treatment as soon as virologic testing is confirmed and within the first 30 days of life in symptomatic cytomegalovirus (CMV)	
101	infected newborns with central nervous system involvement or if life-threatening.	59
102	Stop antiviral treatment in neonates with asymptomatic cytomegalovirus infection	60
Herp	es Simplex Virus (HSV)	61
103	Start aciclovir IV in neonates with Herpes Simplex Virus (HSV) disease, regardless of maternal history or pending laboratory confirmation or exclusion of HSV.	61
104	Start a topical antiviral treatment in combination with aciclovir IV in neonates with Herpes Simplex Virus (HSV) disease with ocular involvment.	62
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	Start Varicella-Zoster Immune Globulin (VZIG) 125IU IM as soon as exposure is known and within a 72 hour period, independent of	
105	maternal history of varicella, in neonates born at <28 weeks of gestational age or who weighed <1000g at birth who have been significantly exposed to Varicella-Zoster Virus (VZV).	63
106	Start Varicella-Zoster Immune Globulin (VZIG) 125IU IM in neonates ≥28w GA or ≥1000g BW who have been significantly exposed postnatally to Varicella-Zoster Virus (VZV), only if born to mother who has no or unkown history of varicella.	63
107	Start Varicella-Zoster Immune Globulin (VZIG) 125IU IM as soon as possible, after birth or with onset of maternal illness, in term or late preterm neonates whose mother had varicella disease 5 days prior to or 2 days after delivery.	64
108	Start aciclovir IV in neonates who develop systemic symptoms or severe cutaneous Varicella-Zoster disease, or who are at high risk of infection.	64
109	Stop Varicella-Zoster immunoglobulin (VZIG) if neonatal chickenpox has developed.	65
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110	Do not use prophylactic antibiotic treatment in neonates at high risk of chlamydial infection (born to mothers who have untreated chlamydia).	66
111	Start erythromycin orally for 14 days in neonates with chlamydial conjunctivitis or with suspected or confirmed chlamydial pneumonia.	66
112	Start azithromycin as second line treatment when erythromycin is not avaliable in neonates with chlamydial conjunctivitis or with suspected or confirmed chlamydial pneumonia.	67
113	Stop topical antibiotics for the treatment of chlamydial conjunctivitis in neonates.	67
	prrhea	68
114115	Administer 1 dose of ceftriaxone IV or IM in all neonates born to mothers who have untreated gonorrhea. Administer 1 dose of ceftriaxone IV or IM in neonates with suspected or confirmed gonococcal ophtalmia neonatorum or other localized	68 68
116	gonococcal infection. Stop topical antibiotics in neonates with suspected or confirmed gonococcal ophtalmia neonatorum.	69
	Start ceftriaxone IV or IM in neonates with disseminated gonococcal infection.	69
ме tг 118	sicillin-Resistant Staphylococcal Aureus (MRSA) Infections Start vancomycin IV until bacteremia is excluded for localized Methicillin-resistant Staphylococcus aureus (MRSA) disease in preterm or	70 70
C!	very low-birthweight neonates or in more-extensive forms of the disease involving multiple sites in full-term neonates.	74
Sypl		71
119	Administer benzylpenicillin G IV, OR procaine penicillin to neonates with confirmed or presumed congenital syphilis, or born to syphilis infected mothers who have not been treated with penicillin at least four weeks prior delivery. Administer 1 does of horzathing penicillin G IM in penators with permal examination, born to syphilis infected methors who have been	71
120	Administer 1 dose of benzathine penicillin G IM in neonates with normal examination, born to syphilis infected mothers who have been adequately treated during pregnancy more than 4 weeks prior to delivery.	72
	plasma Urealyticum Infection	73
121	Reassess the use of macrolides or other antibiotics for the treatment of Ureaplasma urealyticum in neonates.	73





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122	Start empiric antibiotics after urine samples and cultures are collected in neonates with fever when urinary tract infection is suspected.	74
123	Consider antibiotic prophylaxis after an urinary tract infection (UTI) only in neonates with grade IV-V vesico-ureteric reflux.	75
	ussis	76
124	Start azithromycin oral daily for 5 days in neonates with suspected or confirmed pertussis infection, or in those in close contact with confirmed and contagious cases of pertussis.	76
Tub	erculosis	77
125	Start isoniazid prophylaxis orally in neonates born to mothers with tuberculosis (TB), or those in close contact with people with smear-positive pulmonary or laryngeal TB who have not had at least 2 weeks of anti-tuberculosis treatment.	77
126	Start anti-tuberculosis treatment in neonates with congenital tuberculosis or postnatal tuberculosis primary pulmonary disease.	77
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Meta	abolic Bone Disorder (MBD)	78
127	Administer calcium, phosphate and vitamin D in preterm infants <32 weeks of gestational age or <1500g or infants at risk of metabolic bone disorders.	78
128	Administer the maximal recommended doses of calcium, phosphate and vitamin D to prevent fractures in neonates with biochemical features of metabolic bone disease.	79
129	Stop steroids and furosemide as soon as possible in neonates at risk of metabolic bone disorder.	79
Thy	oid Disorders (OR Hypothyroidism)	80
130	Start levothyroxine (L-T4) immediately in neonates with thyroid function test (TFT) that results in either a free T4 (FT4) concentration below norms for age or a venous TSH concentration > 20 mIU/L.	w 80
Нур	erglycemia	81
131	Decrease glucose intake if necessary and decrease or stop drugs that worsen hyperglycemia, in neonates with hyperglycemia.	81
	Start insulin only in patients with persistent hyperglycemia when other methods of glucose control have failed.	82
	Do not provide high glucose infusion rates to prevent hypoglycemia in neonates receiving parenteral nutrition.	82
	Do not use early insulin therapy in neonates at risk of hyperglycemia.	83
Нур	oglycemia	84
135	Start IV glucose infusion in asymptomatic neonates with serum glucose level of <2.6 mmol/L if increased enteral caloric intake is not effective.	84
	Start IV glucose infusion immediately in symptomatic neonates with glucose levels <2.6 mmol/L.	85
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	y & Breast-Feeding	86
	Check the possible milk transfer of drugs taken by mothers to breastfed neonates, and monitor for potential adverse drug effects.	86
	g-drug Interactions	87
	Check changes in drug effect when initiating strong inhibitors or inducers of the cytochrome P450 and/or P-glycoprotein.	87
Vari		88
139	Do not use ceftriaxone in neonates who are being, or who have recently been given any IV fluids that contain calcium (such as TPN or Ringer Lactate)	88
140	Do not use trimethoprime - sulfamethoxazole in neonates.	88
141	Check excipients contained in prescribed drug formulations administered orally or parenterally since they can be harmful and responsible for adverse events in neonates, due to immature metabolism.	89





01. BASIC MANAGEMENT Body Care

Item 1

Do not use routinely topical ointments in preterm neonates.

There is no evidence that the use of emollient therapy prevents invasive infection or death in preterm infants in high-income countries.

Grade of recommendations: FRN

Stop

Cleminson J, McGuire W. Topical emollient for prevention of infection in preterm infants: a systematic review. *Lancet.* 2016;(1). doi:10.1002/14651858.CD001150.pub3.

Erdemir A, Kahramaner Z, Yuksel Y, et al. The effect of topical ointment on neonatal sepsis in preterm infants. *J Matern neonatal Med.* 2015;28(1):33-36. doi:10.3109/14767058.2014.900037.

Raboni R, Patrizi A, Cocchi G, Faldella G, Raone B. Comparison of two different neonatal skin care practices and their influence on transepidermal water loss in healthy newborns within first 10 days of life. *Minerva Pediatr.* 2014;66(October):369-374.

Campbell JR, Zaccaria E, Baker CJ. Systemic candidiasis in extremely low birth weight infants receiving topical petrolatum ointment for skin care: a case-control study. *Pediatrics*. 2000;105(5):1041-1045. doi:10.1542/peds.105.5.1041.

< 37w GA

Basic Management/Body Care

Ointments

Item 2

Stop the use of antiseptics <u>for the daily care</u> of the uncomplicated umbilical cord in <u>healthy hospitalized term</u> neonates.

In high-income settings, there is limited research which has not shown an advantage of antibiotics or antiseptics over simply keeping the cord clean. Antimicrobial agent may actually delay the time to cord separation. On the contrary, there is high-quality evidence that chlorhexidine skin or cord care in the community setting results in reduction of the incidence of omphalitis and neonatal mortality.

Stop

Grade of recommendations: FRN

Medves JM, O'Brien BAC. Cleaning solutions and bacterial colonization in promoting healing and early separation of the umbilical cord in healthy newborns. *Can J Public Heal.* 1997;88(6):380-382. doi:9458563.

Sinha A, Sazawal S, Pradhan A, et al. Chlorhexidine skin or cord care for prevention of mortality and infections in neonates. *Cochrane database Syst Rev.* 2015;3(3):CD007835. doi:10.1002/14651858.CD007835.pub2.

Zupan J, Garner P, Aaa O. Topical umbilical cord care at birth. *Cochrane database Syst Rev.* 2004;(3). doi:10.1002/14651858.CD001057.

Imdad A, Bautista RMM, Senen KAA, et al. Umbilical cord antiseptics for preventing sepsis and death among newborns. *Cochrane database Syst Rev.* 2013;(5). doi:10.1002/14651858.CD008635.pub2.

All

Basic Management/Body Care
Antiseptics





Vaccination

Item 3

Administer a dose of DTPa-IPV/Hib ± HBV and of pneumococcal vaccine at 60, 90 and 120 days of postnatal life to all hospitalized preterm neonates.

Hospitalized preterm infants should receive immunisation against diphtheria, tetanus, pertussis (whooping cough), polio, Haemophilus influenzae type b and pneumococcal at 2, 3 and 4 months of postnatal age. Use the combined vaccine. Cardiorespiratory function should be monitored in unstable preterm infants for 48 hours following immunisation. In preterm neonates <33 weeks of gestational age who will be discharged before their 60th day of postnatal age, the first dose of DTPa-IPV/Hib and of pneumococcal vaccine should be advanced. In this case, the immunisation can be administered from the 50th day of postnatal life, followed by booster vaccination 1 and 2 month later.

Start

Grade of recommendations: National Guidelines

Département de l'enfant et de l'adolescent - HUG. Cahier de l'interne. *Hôpitaux Univ Genève*. 2015;(Décembre).

Office fédéral de la santé publique, Commission fédérale pour les vaccinations, Société suisse de néonatologie, Société suisse de pédiatrie. Vaccinations des enfants nés prématurément. *Directives et recommandations.* 2009.

Office fédéral de la santé publique, Commission fédérale pour les vaccinations. Plan de vaccination suisse 2016. *Directives et recommandations*. 2016.

<37w GA

Basic Management/Vaccination

Item 4

Recommend BCG vaccine at discharge to neonates at high risk of tuberculosis exposure in the first year of life.

BCG vaccine is recommended only to newborns at risk of developing disseminated tuberculosis. Infants at risk are those who come from AND will return definitely to regions with high prevalence of tuberculosis (Africa, Asia, Latine America, Eastern Europe) before the age of 1 year of postnatal life. Short stays (vacation) in those regions are not an indication for immunisation.

Start

Grade of recommendations: National Guidelines

Office fédéral de la santé publique, Commission fédérale pour les vaccinations, Société suisse de néonatologie, Société suisse de pédiatrie. Vaccinations des enfants nés prématurément. Directives et recommandations. 2009.

Office fédéral de la santé publique, Commission fédérale pour les vaccinations. Plan de vaccination suisse 2016. *Directives et recommandations*. 2016.

All

Basic Management/Vaccination





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Item 5	
	Check / administer Pertussis vaccination to close contacts of neonates.
Start	Preterm infants are at high risks of pertussis. Booster vaccination is recommended to the mother (unless immunized during pregnancy), the father (unless the last booster dates less than 10 years), the siblings (unless up-to-date), the grand-parents and/or all those who will be in close contact to the neonate before the age of 4 months (i.e. reception of 2 vaccine doses).
	Grade of recommendations: National Guidelines
	Office fédéral de la santé publique, Commission fédérale pour les vaccinations. Plan de vaccination suisse 2016. <i>Directives et recommandations</i> . 2016.
All	Basic Management/Vaccination
Item 6	
	Check status and recommend or administer vaccination to close contacts of
	neonates.
	Pertussis: booster vaccination is recommended to the mother (unless immunized during pregnancy), the father (unless the last booster dates less than 10 years), the siblings (unless up-to-date), the grand-parents and/or <u>all those who will be in close contact to the neonate before the age of 4 months</u> (i.e. reception of 2 vaccine doses).
	Haemophilus inflenzae type b: catch-up vaccination to brothers and sisters <5 years old.

Pneumococcal: catch-up vaccination to brothers and sisters <5 years old.

ROR: catch-up vaccination for all members of the family circle.

Varicella: catch-up vaccination for all members of the family circle.

Start

Grade of recommendations: National Guidelines

Office fédéral de la santé publique, Commission fédérale pour les vaccinations, Société suisse de néonatologie, Société suisse de pédiatrie. Vaccinations des enfants nés prématurément. Directives et recommandations. 2009.

Influenza: immunisation for all members of the family circle during the two first winters.

Office fédéral de la santé publique, Commission fédérale pour les vaccinations. Plan de vaccination suisse 2016. Directives et recommandations. 2016.

ΑII

Basic Management/Vaccination





Parenteral Nutrition (PN)

Item 7

Start parenteral nutrition shortly after birth in all preterm neonates when it is clear that enteral feeds will not be tolerated soon.

In the small preterm infant, starvation for just one day may be detrimental. Recommended volumes of parenteral nutrition:

<1500g BW (ml/kg/day):

Day 1: Fluid 80-90 ml/kg/day

Day 2: Fluid 100-110 ml/kg/day

Day 3: Fluid 120-130 ml/kg/day

Day 4: Fluid 130-150 ml/kg/day

Day 5: Fluid 140-160 ml/kg/day

Day 6: Fluid 160-180 ml/kg/day

>1500g BW (ml/kg/day):

Day 1: Fluid 60-80 ml/kg/day

Day 2: Fluid 80-100 ml/kg/day

Day 3: Fluid 100-120 ml/kg/day

Day 4: Fluid 120-150 ml/kg/day

Start Day 5: Fluid 140-160 ml/kg/day

Day 6: Fluid 140-160 ml/kg/day

Target parenteral energy intake (including protein) of stable patients may be roughly estimated as 110-120 kcal/kg for preterm infants. Energy intake should be adapted in patients with disease states that increase resting energy expenditure, such as pulmonary and cardiac disorders but should not be increased after uncomplicated surgery.

Grade of recommendations: International Guidelines

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

Moyses HE, Johnson MJ, Leaf AA, Cornelius VR. Early parenteral nutrition and growth outcomes in preterm infants: A systematic review and meta-analysis. *Am J Clin Nutr.* 2013;97(4):816-826. doi:10.3945/ajcn.112.042028.

Ehrenkranz RA. Early, Aggressive Nutritional Management for Very Low Birth Weight Infants: What Is the Evidence? *Semin Perinatol.* 2007;31(2):48-55. doi:10.1053/j.semperi.2007.02.001.

<37w GA Basic Management/Nutrition

Item 8

Start reducing the percentage of parenteral nutrition as quickly as possible by the introduction of enteral nutrition until enteral nutrition finally replaces completely PN in order to minimise any side-effects from exposure to PN.

Stop

Aim to reach full enteral feeding by about two weeks in babies weighing <1000 g at birth and by about one week in babies weighing 1000–1500g as clinically feasible.

Grade of recommendations: National Guidelines

Fusch C, Bauer K, Böhles HJ, et al. Neonatology/Paediatrics - Guidelines on Parenteral Nutrition, Chapter 13. *Ger Med Sci.* 2009;7:1-23. doi:10.3205/000074.

Dutta S, Singh B, Chessell L, et al. Guidelines for Feeding Very Low Birth Weight Infants. *Nutrients*. 2015;7:423-442. doi:10.3390/nu7010423.

<37w GA

Basic Management/Nutrition

Parenteral nutrition





Item 9

Start continuous parenteral glucose administration in preterm infants needing parenteral nutrition.

An early start of parenteral glucose together with amino acids from the very first day onwards contributes to preventing hyperglycemia in premature infants. The recommended starting dose of glucose is 4-8 mg/kg/min (5.8-11.5 g/kg/day). Recommended parenteral glucose supply in parenteral nutrition:

Neonates up to 3 kg: Day 1: 7 mg/kg/min (10 g/kg/day), Day 2: 9.7 mg/kg/min (14 g/kg/day), Day 3: 11.1 mg/kg/min (16 g/kg/day), Day 4: 12.5 mg/kg/min (18 g/kg/day). Glucose intake should usually cover 60–75% of non-protein calories. These recommendations need to be adapted to the clinical situation to oral and/or enteral energy intake and to the required weight gain for normal or catch up growth. It is important, especially when prescribing PN for infants, to accurately evaluate the carbohydrate load provided by concurrent infusion therapy. An excessively high carbohydrate intake can result in net lipogenesis with hepatic fat deposition and steatosis of the liver.

Start

Grade of recommendations: International Guidelines

Fusch C, Bauer K, Böhles HJ, et al. Neonatology/Paediatrics - Guidelines on Parenteral Nutrition, Chapter 13. *Ger Med Sci.* 2009;7:1-23. doi:10.3205/000074.

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

Cai W. CSPEN guidelines for nutrition support in neonates. *Asia Pac J Clin Nutr.* 2013;22(4):655-663. doi:10.6133/apjcn.2013.22.4.21.

<37w GA

Basic Management/Nutrition

Parenteral nutrition





Item 10

Start amino acid supply in the first day of life in preterm infant needing parenteral nutrition.

Start with 1.5-2 g/kg/day and increase up to 3.5-4 g/kg/day. Amino acid imbalances can result in toxic organ damage and may be involved in the development of PN-associated cholestasis. Achieving an adequate energy to protein ratio is as important as providing adequate energy intake. Recommended non-protein energy to protein ratio depends on neonate age and weight and varies between 25 and 40 kcal/g of protein (≈150-250 kcal/g of nitrogen). If energy intake is insufficient, protein is used as an energy source, and the nitrogen balance becomes less positive. Increasing the caloric intake will spare the protein loss and improve nitrogen retention, but with limited protein intake, the protein retention reaches a plateau, and the energy excess is used solely for fat deposition.

Grade of recommendations: B

Start

Cai W. CSPEN guidelines for nutrition support in neonates. *Asia Pac J Clin Nutr.* 2013;22(4):655-663. doi:10.6133/apjcn.2013.22.4.21.

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

Fusch C, Bauer K, Böhles HJ, et al. Neonatology/Paediatrics - Guidelines on Parenteral Nutrition, Chapter 13. *Ger Med Sci.* 2009;7:1-23. doi:10.3205/000074.

Ziegler EE. Meeting the nutritional needs of the low-birth-weight infant. *Ann Nutr Metab.* 2011;58(SUPPL. 1):8-18. doi:10.1159/000323381.

Agostoni C, Buonocore G, Carnielli VP, et al. Enteral nutrient supply for preterm infants: commentary from the European Society of Paediatric Gastroenterology, Hepatology and Nutrition Committee on Nutrition. *J Pediatr Gastroenterol Nutr.* 2010;50(1):85-91. doi:10.1097/MPG.0b013e3181adaee0.

Rigo J, Senterre J. Nutritional needs of premature infants: Current Issues. *J Pediatr.* 2006;149(5):80-88. doi:10.1016/j.jpeds.2006.06.057.

Tudehope D, Vento M, Bhutta Z, Pachi P. Nutritional requirements and feeding recommendations for small for gestational age infants. *J Pediatr.* 2013;162(3 SUPPL.):S81-S89. doi:10.1016/j.jpeds.2012.11.057.

<37w GA

Basic Management/Nutrition

Parenteral nutrition





Item 11

Start continuous lipid emulsion infusion within the first 24-48 hours of life in preterm infant needing parenteral nutrition,

The initiation of lipids within the first 2 days of life in VLBW infants appears to be safe and well tolerated; however, beneficial effects on growth could not be shown for this treatment. Lipid intake should usually provide 25–40% of non-protein calories in fully parenterally fed patients. The recommended starting dose of lipid emulsions is 1-2 g/kg/day and is increased by 0.5-1.0 g/kg/day, up to 3 g/kg/day. No difference have been shown between the different lipid emulsion formulations. Reduction of the dosage of lipid emulsions should be considered if serum or plasma triglyceride concentrations during infusion exceed 250 mg/dL. In critically ill or infected patients receiving lipid emulsions, more frequent monitoring of plasma triglyceride concentration and dose adjustment in case of hyperlipidaemia are recommended. Lipid emulsions should be protected by validated light-protected tubing during phototherapy to decrease the formation of hydroperoxides.

Start

Grade of recommendations: B

Vlaardingerbroek H, Veldhorst MAB, Spronk S, et al. Parenteral lipid administration to very-low-birth-weight infants - Early introduction of lipids and use of new lipid emulsions: A systematic review and meta-analysis. *Am J Clin Nutr.* 2012;96(2):255-268. doi:10.3945/ajcn.112.040717.

Drenckpohl D, McConnell C, Gaffney S, et al. Randomized Trial of Very Low Birth Weight Infants Receiving Higher Rates of Infusion of Intravenous Fat Emulsions During the First Week of Life. *Pediatrics.* 2008;122(4):743-751. doi:10.1542/peds.2007-2282.

Cai W. CSPEN guidelines for nutrition support in neonates. *Asia Pac J Clin Nutr.* 2013;22(4):655-663. doi:10.6133/apjcn.2013.22.4.21.

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

Hojsak I, Colomb V, Braegger C, et al. ESPGHAN Committee on Nutrition Position Paper. Intravenous Lipid Emulsions and Risk of Hepatotoxicity in Infants and Children: a Systematic Review and Meta-analysis. *J Pediatr Gastroenterol Nutr.* 2016;62(5):776-792. doi:10.1097/MPG.000000000001121.

<37w GA

Basic Management/Nutrition
Parenteral nutrition

Item 12

Do not administer parenteral lipid emulsion at a dose higher than 3-4 g/kg/day in neonates.

Maximum lipid oxidation of 4 g/kg/day is reached in full-term neonates with a glucose intake below 18 g/kg/day. An increase in the concentration of plasma triglycerides is to be expected if the infusion speed of the lipid emulsions exceeds the speed of hydrolysis of the triglycerides.

Other

Grade of recommendations: B

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

Fusch C, Bauer K, Böhles HJ, et al. Neonatology/Paediatrics - Guidelines on Parenteral Nutrition, Chapter 13. *Ger Med Sci.* 2009;7:1-23. doi:10.3205/000074.

All

Basic Management/Nutrition

Parenteral nutrition





Item 13

Check that a sufficient quantity of linoleic acid is administered in all neonates on parenteral nutrition.

In order to prevent Essential Fatty Acids (EFA) deficiency a minimum linoleic acid intake of 0.25 g/kg per day should be given to preterm infants and 0.1 g/kg per day to term neonates. Linoleic acid is contained in vegetal oils: soy oil (54%) and olive oil (10%). The appoximate linoleic acid content in existing lipid solutions on the swiss market are:

Lipofundin®: 29% of total lipids Lipoplus®: 24% of total lipids Omegaven®: 4% of total lipids SMOFlipid®: 19% of total lipids.

Start

Grade of recommendations: D

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

Fusch C, Bauer K, Böhles HJ, et al. Neonatology/Paediatrics - Guidelines on Parenteral Nutrition, Chapter 13. *Ger Med Sci.* 2009;7:1-23. doi:10.3205/000074.

Vlaardingerbroek H, Veldhorst MAB, Spronk S, et al. Parenteral lipid administration to very-low-birth-weight infants - Early introduction of lipids and use of new lipid emulsions: A systematic review and meta-analysis. *Am J Clin Nutr.* 2012;96(2):255-268. doi:10.3945/ajcn.112.040717.

All Basic Management/Nutrition
Parenteral nutrition

Item 14

Start electrolytes supplementation with parenteral nutrition after onset of diuresis.

Recommended starting dose is sodium 2-3 mmol/kg/day, potassium 1-2 mmol/kg/day, calcium 0.6-0.8 mmol/kg/day, phosphates 1.0-1.2 mmol/kg/day and magnesium 0.3-0.4 mmol/kg/day.

Start Grade of recommendations: D

Cai W. CSPEN guidelines for nutrition support in neonates. *Asia Pac J Clin Nutr.* 2013;22(4):655-663. doi:10.6133/apjcn.2013.22.4.21.

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

All Basic Management/Nutrition

Parenteral nutrition





Item 15

Start vitamins and trace elements supplementation in neonates receiving parenteral nutrition. The optimum time to begin with trace element supplementation in premature infants <1500 g birth weight is not clear. It is proposed to start supplementation on the 5th day of life to coincide with an increase in body weight. Vitamin preparations should, if possible, be administered together with the lipid emulsion in order to limit light-induced lipid peroxidation and vitamin loss. Parenteral zinc supply is recommended in daily dosages of 450–500 mg/kg per day for premature infants. Start Grade of recommendations: D Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). J Pediatr Gastroenterol Nutr. 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4. Fusch C, Bauer K, Böhles HJ, et al. Neonatology/Paediatrics - Guidelines on Parenteral Nutrition, Chapter 13. Ger Med Sci. 2009;7:1-23. doi:10.3205/000074. Basic Management/Nutrition All Parenteral nutrition Item 16 Start vitamin D supplementation from the first days of life in all neonates. Recommended dose is 400 IU once daily during the first year of life and 600 IU once daily during the two next years. Start

Paediatrica. 2012;23(4):22-24.

All Basic Management/Nutrition
Parenteral nutrition

L'Allemand D, Neuhaus TJ, Janner M, et al. Recommandations de l' Office fédéral de la santé publique concernant l'apport en vitamine D en Suisse – quelle signification pour le pédiatre?

Grade of recommendations: National Guidelines





2. CARDIOLOGY **Congenital Heart Disease** Item 17 Start prostaglandin E1 (alprostadil) as an initial continuous intravenous infusion at 0.01 mcg/kg/min, until a definitive diagnosis is made in an infant suspected of having ductus-dependant heart disease. Start Grade of recommendations: A Donofrio MT, Moon-Grady AJ, Hornberger LK, et al. Diagnosis and treatment of fetal cardiac disease: a scientific statement from the American Heart Association. Circulation . 2014;129(21):2183-2242. doi:10.1161/01.cir.0000437597.44550.5d. Eichenwald EC, Kim MS, Weisman LE. Overview of cyanosis in the newborn. UpToDate.com. http://www.uptodate.com/contents/overview-of-cyanosis-in-the-newborn. Published 2014. Accessed March 10, 2016. Cardiology/Congenital Heart Disease ΑII Item 18 Reassess the indication of prostaglandin E1 (PGE1) treatment. Infants with Non Duct Dependent Lesions (Ventricular septal defect and others such as Atrioventricular canal defect) do not require PGE1 infusion. Grade of recommendations: Institutional Guidelines Stop Clinical Practice Committee. Antenatally Diagnosed Major Congenital Heart Disease Management at Delivery and in NICU. Newborn Services Clinical Guideline - Auckland District Health Board. http://www.adhb.govt.nz/newborn/Guidelines/Cardiac/AntenatallyDiagnosedCHD.htm. Published 2013. Accessed March 3, 2016. Cardiology/Congenital Heart Disease ΑII Prostaglandin E1 (PGE1) Item 19 Stop ibuprofen, indomethacin and paracetamol in patients with duct dependent congenital heart disease. Ibuprofen, indomethacin or paracetamol must not be used in patients with congenital heart disease in whom patency of the ductus arteriosus is necessary for satisfactory pulmonary or systemic blood flow (e.g. pulmonary atresia, severe tetralogy of Fallot, severe coarctation of the aorta). Stop Grade of recommendations: Manufacturer Ovation Pharmaceuticals. Product Information: Neoprofen(R) IV injection. Micromedex - Truven Health Analytics Inc. http://www.micromedexsolutions.com/. Published 2006. Accessed September 15, 2016. Fresenius Kabi USA. Product Information: Indomethacin IV injection. Micromedex - Truven Health Analytics Inc. http://www.micromedexsolutions.com/. Published 2014. Accessed September 15, 2016.



ΑII



Cardiology/Congenital Heart Disease Ibuprofen, indomethacin, paracetamol

Patent Ductus Arteriosus (PDA)

Confirmed patent ductus arteriorus (PDA)

Substantial ductal shunting may be associated with an increased ratio of left atrial to aortic Definition root dimensions ≥1.5:1, ductal diameter ≥1.5 mm, left ventricular volume and pressure loading, and reversal of diastolic flow in the descending aorta or in cerebral or renal arteries. Benitz WE. Patent Ductus Arteriosus in Preterm Infants. Pediatrics. 2016;137(1):1-6. Item 20 Consider pharmacological closure of confirmed patent ductus arteriosus (PDA) in preterm neonates after 2 weeks of life, with ibuprofen as first-line treatment. Ibuprofen is as effective as indomethacin in closing a PDA and currently appears to be the drug of choice. Ibuprofen reduces the risk of NEC and transient renal insufficiency. Recommended dose is 10 mg/kg as the initial dose followed by 5 mg/kg 24 and 48 hours Start later. When possible, choose the enteral route for the administration of ibuprofen. **Grade of recommendations: FRN** Ohlsson A, Walia R, Shah Sachin S. Ibuprofen for the treatment of patent ductus arteriosus in preterm or low birth weight (or both) infants. Cochrane Database Syst Rev. 2015;(2). doi:10.1002/14651858.CD003481.pub6. Benitz WE. Patent Ductus Arteriosus in Preterm Infants. Pediatrics. 2016;137(1):1-6. <37w GA or Cardiology/PDA <2500g BW Item 21 Reassess the indication of ibuprofen, indomethacin and paracetamol in preterm neonates <2 weeks of life with confirmed or unconfirmed patent ductus arteriosus (PDA). The cumulative evidence supports the conclusion that early (in the first 2 weeks after birth), routine (as prophylaxis or for infants with echocardiographic confirmation of ductal patency Stop with or without clinical signs) treatment to close the ductus arteriosus does not improve long-term outcomes for preterm infants. **Grade of recommendations: National Guidelines** Benitz WE. Patent Ductus Arteriosus in Preterm Infants. Pediatrics. 2016;137(1):1-6.



<37w GA



Cardiology/PDA

Ibuprofen, indomethacin, paracetamol

Item 22

item 22	
	Reassess the indication of ibuprofen, indomethacin and paracetamol in term neonates with patent ductus arteriosus (PDA).
Stop	In term neonates, inhibitors of prostaglandin synthesis are not effective for PDA closure, and thus are not recommended.
1 0.0	Grade of recommendations: Textbook
	Grade of recommendations. Textbook
	Doyle T, Kavanaugh-McHugh A, Soslow J, Hill K. Management of patent ductus arteriosus. <i>UpToDate.com.</i> http://www.uptodate.com/contents/management-of-patent-ductus-arteriosus. Published 2016. Accessed March 14, 2016.
>37w GA	Cardiology/PDA
	Ibuprofen, indomethacin, paracetamol
Item 23	
	Do not use paracetamol as first-line treatment for patent ductus arteriosus (PDA) closure. Consider a switch to ibuprofen.
Stop	Paracetamol appears to be a promising new alternative to indomethacin and ibuprofen for the closure of a PDA with potentially fewer adverse effects. Additional studies testing this intervention with long-term follow-up are needed before paracetamol can be recommended as standard treatment for a PDA in preterm infants.
	Grade of recommendations: FRN
	Ohlsson A, Shah PS. Paracetamol (acetaminophen) for prevention or treatment of pain in newborns. Cochrane Database Syst Rev. 2015;(6). doi:10.1002/14651858.CD011219.pub2.
<34w GA or	Cardiology/PDA
<2500g	Paracetamol
	radotanto





Hypotension

Hypotension

Definition

Hypotension is defined as a mean blood pressure (MBP) <30 mmHg or, during the first 3 days of postnatal life, a MBP with a number lower than the infant's gestational age in weeks

Vargo L, Seri I. The management of hypotension in the very-low-birth-weight infant: guideline for practice. *National Association of Neonatal Nurses*. 2011.

Subhedar Nimish V, Shaw NJ. Dopamine versus dobutamine for hypotensive preterm infants. *Cochrane Database Syst Rev.* 2003;(3). doi:10.1002/14651858.CD001242.

Item 24

Do not use volume expansion as first line treatment in VLBW infants (BW <1500g) with hypotension.

Hypotension in VLBW infants should be treated on the basis of the etiology of the hypotension, when an etiology is known. In general, the early use of volume expansion with normal saline solution, fresh frozen plasma, albumin, plasma substitute, or blood in VLBW infants with hypotension is not recommended. Evidence that VLBW infants with hypotension benefit from volume expansion is insufficent, as is the evidence to determine what type of volume expansion should be used in VLBW infants. The majority of VLBW infants who are hypotensive are not hypovolemic and have normal circulating blood volume.

Grade of recommendations: National Guidelines

Dempsey, E. M. Challenges in Treating Low Blood Pressure in Preterm Infants. *Children.* 2015;2(2), 272–288. doi:10.3390/children2020272

Vargo L, Seri I. The management of hypotension in the very-low-birth-weight infant: guideline for practice. *National Association of Neonatal Nurses*. 2011.

<1500g BW

Stop

Cardiology/hypotension

Normal saline, fresh frozen plasma, albumin, plasma substitute, blood

Item 25

Consider a conservative approach (permissive hypotension) for the managment of VLBW infants (BW <1500g) if the clinical examination is satisfactory in the face of apparent hypotension

A careful clinical and biochemical assessment of a potentially hypotensive infant is an essential first step towards management. This should include: heart rate, capillary refill time, urine output, serum lactate concentration, pH, base excess and haemoglobin. If a pharmacological treatment is considered, dopamine can be a valid option for the sole treatment of hypotension. Hydrocortisone may be as effective as dopamine when used as a primary treatment for hypotension. Clinical trials are underway and could provide stronger recommendations in the near future to guide clinicians in the management of the hypotension of the VLBW neonate.

Start / Stop

Grade of recommendations: FRN

Dempsey, E. M. Challenges in Treating Low Blood Pressure in Preterm Infants. *Children.* 2015;2(2), 272–288. doi:10.3390/children2020272

Ibrahim H, Sinha IP, Subhedar NV. Corticosteroids for treating hypotension in preterm infants. *Cochrane Database Syst. Rev.* 2011;12. doi:10.1002/14651858.CD003662.pub4.

Subhedar Nimish V, Shaw NJ. Dopamine versus dobutamine for hypotensive preterm infants. *Cochrane Database Syst Rev.* 2003;(3). doi:10.1002/14651858.CD001242.

Paradisis M, Osborn DA. Adrenaline for prevention of morbidity and mortality in preterm infants with cardiovascular compromise. *Cochrane Database Syst Rev.* 2004;1. doi:10.1002/14651858.CD003958.pub2.

Vargo L, Seri I. The management of hypotension in the very-low-birth-weight infant: guideline for practice. *National Association of Neonatal Nurses.* 2011.

<1500g BW

Cardiology/hypotension





03. HEMATOLOGY

Anemia

Item 26

Do not use routinely erythropoietin to limit exposure to blood transfusions in preterm neonates. The indication of treatment should be reassessed.

Aside from research about its possible neuroprotective potential (EPO appears to be a promising drug in many conditions where neonatal brain injury occurs), there is little current justification for the use of erythropoietin in neonatal medicine, except in a few limited situations (for example to respect the views of parents who are Jehovah's witnesses).

Stop

Grade of recommendations: A

Ohlsson A, Sm A. Early erythropoietin for preventing red blood cell transfusion in preterm and / or low birth weight infants (Review). *Cochrane Database Syst Rev.* 2014;(4). doi:10.1002/14651858.CD004863.pub4.

Sm A, Ohlsson A. Late erythropoietin for preventing red blood cell transfusion in preterm and/or low birth weight infants. *Cochrane Database Syst Rev.* 2014;(4). doi:10.1002/14651858.CD004868.pub4.

Canadian Paediatric Society Fetus and Newborn Committee, Lemyre B, Sample M, Lacaze-Masmonteil T. Minimizing blood loss and the need for transfusions in very premature infants. *Paediatr Child Heal.* 2015;20(8):451-456.

Neonatal Formulary. Erythropoietin. Neonatal Formulary.

http://www.neonatalformulary.com/pdfs/commentary/ERYTHROPOIETIN-(commentary).pdf. Published 2014. Accessed July 12, 2016.

<37w GA

Hematology/Anemia OR Prevention/Anemia Erythropoietin (EPO)

Item 27

Start iron supplement of 2-3 mg/kg/day in all preterm infants fed human milk once full oral feeds have been achieved.

This is the amount of iron supplied by iron-fortified formulas and infant fed with preterm formula do not need supplementation. Preterm infants fed human milk should receive an iron supplement of 2-3 mg/kg/day starting once full enteral feed have been achieved and continued until the infant is weaned to iron-fortified formula or begins eating complementary foods that supply the 2 mg/kg of iron. An exception to this practice would include infants who have received an iron load from multiple transfusions of packed red blood cells. Term healthy infants have sufficient iron for at least the first 4 months of life and should not receive iron supplementation. Supplementation with 4-6 mg/kg/day can be considered in newborns who are iron deficient. The available data suggest that infants who receive iron supplementation have a slightly higher haemoglobin level, improved iron stores and a lower risk of developing iron deficiency anaemia when compared with those who are unsupplemented. However, it is unclear whether iron supplementation in preterm and low birth weight infants has long term benefits in terms of neurodevelopmental outcome and growth.

Start

Grade of recommendations: National Guidelines

Baker RD, Greer FR, American Academy of Pediatrics. Diagnosis and Prevention of Iron Deficiency and Iron-Deficiency Anemia in Infants and Young Children (0–3 Years of Age). *Pediatrics*. 2010;126(5):1040-1050. doi:10.1542/peds.2010-2576.

Mills RJ, **Davies MW**. Enteral iron supplementation in preterm and low birth weight infants. *Cochrane Database Syst Rev.* 2012;(3):CD005095. doi:10.1002/14651858.CD005095.pub2.

Canadian Paediatric Society Fetus and Newborn Committee, Lemyre B, Sample M, Lacaze-Masmonteil T. Minimizing blood loss and the need for transfusions in very premature infants. *Paediatr Child Heal.* 2015;20(8):451-456.

<37w GA

Hematology/Anemia OR Prevention/Anemia





Coagulation disorders Item 28 Start oral Vitamin K in neonates breastfed by a mother treated with phenprocoumone. Breastfed infants from mothers treated with phenprocoumone should receive oral vitamin K (phytomenadione) 1mg once a week. This doesn't apply if the mother is treated with acenocoumarol because of the short elimination half life of this drug. Start **Grade of recommendations: National Guidelines** Schubiger G, Laubscher B, Bänziger O, Société Suisse de Néonatologie, Commission de nutrition de la Société suisse de pédiatrie, Société suisse de gynécologie et obstétrique. Prophylaxie à la vitamine K chez le nouveau-né : nouvelles recommandations. Swiss Soc Neonatol. 2003. ΑII Hematology/Coagulation disorders

Item 29	
	Check in all neonates that a complete Vitamin K prophylaxis has been given at birth.
Start	Adequate prophylaxis depends on clinical context and gestational age: healthy neonates >34 weeks of gestational age; >2000g birthweight: 4 hours after birth: 2 mg of oral phytomenadione 4 days after birth: 2 mg of oral phytomenadione 4 weeks after birth: 2 mg of oral phytomenadione ill neonates / preterms with infusion / nil by mouth neonates: 4 hours after birth: 0.5 mg of IV/IM phytomenadione 4 weeks after birth: 2 mg of oral phytomenadione
	Grade of recommendations: National Guidelines Schubiger G, Laubscher B, Bänziger O, Société Suisse de Néonatologie, Commission de
	nutrition de la Société suisse de pédiatrie, Société suisse de gynécologie et obstétrique. Prophylaxie à la vitamine K chez le nouveau-né : nouvelles recommandations. Swiss Soc Neonatol. 2003.
All	Hematology/Coagulation disorders OR Prevention/Coagulation disorders





Thrombocytopenia and Platelet Dysfunction

Risk factors for major bleeding in infant with thrombocytopenia:

- <1000g and <7 days
- Clinically unstable (e.g. fluctuating BP)

- Definition Previous major bleeding (e.g. Grade 3-4 IVH, pulmonary haemorrhage)
 - Current minor bleeding
 - Concurrent coagulopathy
 - Requiring surgery or exchange transfusion

Roberts I, Murray N. Neonatal thrombocytopenia: causes and management. Arch Dis Child - Fetal Neonatal Ed. 2003;88(5):359F-364. doi:10.1136/fn.88.5.F359.

Clinical Practice Committee. Neonatal Thrombocytopenia. Newborn Services Clinical Guideline -Auckland District Health Board.

http://www.adhb.govt.nz/newborn/Guidelines/Blood/Platelets/NeonatalThrombocytopenia.htm. Published 2016. Accessed July 15, 2016.

Item 30

Consider platelets transfusion even in the absence of bleeding in all neonates with a platelet count of <30x109/L.

The recommended volume of platelets to be transfused is 10-20 mL/kg. Transfuse over 30 minutes and monitor platelet count one hour post transfusion.

In neonates receiving platelet transfusion, the administered platelet type should be:

- Human platelet antigen (HPA) compatible platelets for neonates with neonatal alloimmune thrombocytopenia (NAIT), except in emergency situations in order to avoid delays to platelet transfusion; in that case, consider IV immunoglobulin (IVIG) (1-2 g/kg) in combination with the unmatched platelets.
- Blood group-compatible, cytomegalovirus (CMV) negative for all other infants.
- Irradiation of platelets is not routinely required but consider for babies with definite or suspected immunodeficiency or those who have undergone intrauterine transfusions.

Start

Grade of recommendations: Institutional Guidelines

Roberts I, Murray N. Neonatal thrombocytopenia: causes and management. Arch Dis Child - Fetal Neonatal Ed. 2003;88(5):359F-364. doi:10.1136/fn.88.5.F359.

Clinical Practice Committee. Neonatal Thrombocytopenia. Newborn Services Clinical Guideline -Auckland District Health Board.

http://www.adhb.govt.nz/newborn/Guidelines/Blood/Platelets/NeonatalThrombocytopenia.htm. Published 2016. Accessed July 15, 2016.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

ΑII

Hematology/Thrombocytopenia and Platelet Dysfunction





Item 31

Consider platelets transfusion in neonates with a platelet count of 30-49x10⁹/L and minor bleeding or those at risk for major bleeding.

The recommended volume of platelets to be transfused is 10-20 mL/kg. Transfuse over 30 minutes and monitor platelet count one hour post transfusion.

In neonates receiving platelet transfusion, the administered platelet type should be:

- Human platelet antigen (HPA) compatible platelets for neonates with neonatal alloimmune thrombocytopenia (NAIT), except in emergency situations in order to avoid delays to platelet transfusion; in that case, consider IV immunoglobulin (IVIG) (1-2 g/kg) in combination with the unmatched platelets.
- Blood group-compatible, cytomegalovirus (CMV) negative for all other infants.
- Irradiation of platelets is not routinely required but consider for babies with definite or suspected immunodeficiency or those who have undergone intrauterine transfusions.

Start

Grade of recommendations: Institutional Guidelines

Roberts I, Murray N. Neonatal thrombocytopenia: causes and management. *Arch Dis Child - Fetal Neonatal Ed.* 2003;88(5):359F-364. doi:10.1136/fn.88.5.F359.

Clinical Practice Committee. Neonatal Thrombocytopenia. Newborn Services Clinical Guideline - Auckland District Health Board.

http://www.adhb.govt.nz/newborn/Guidelines/Blood/Platelets/NeonatalThrombocytopenia.htm. Published 2016. Accessed July 15, 2016.

Health.vic. Thrombocytopenia in neonates. Victoria State Government.

https://www2.health.vic.gov.au:443/hospitals?and?health?services/patient?care/perinatal?reproductive/neonatal?ehandbook/conditions/thrombocytopenia. Published 2015. Accessed July 18, 2016.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

ΑII

Hematology/Thrombocytopenia and Platelet Dysfunction

Item 32

Consider platelets transfusion in neonates with a platelet of count 50-99x10⁹/L only if bleeding is present.

The recommended volume of platelets to be transfused is 10-20 mL/kg. Transfuse over 30 minutes and monitor platelet count one hour post transfusion.

In neonates receiving platelet transfusion, the administered platelet type should be:

- Human platelet antigen (HPA) compatible platelets for neonates with neonatal alloimmune thrombocytopenia (NAIT), except in emergency situations in order to avoid delays to platelet transfusion; in that case, consider IV immunoglobulin (IVIG) (1-2 g/kg) in combination with the unmatched platelets.
- Blood group-compatible, cytomegalovirus (CMV) negative for all other infants.
- Irradiation of platelets is not routinely required but consider for babies with definite or suspected immunodeficiency or those who have undergone intrauterine transfusions.

Start

Grade of recommendations: Institutional Guidelines

Roberts I, Murray N. Neonatal thrombocytopenia: causes and management. *Arch Dis Child - Fetal Neonatal Ed.* 2003;88(5):359F-364. doi:10.1136/fn.88.5.F359.

Clinical Practice Committee. Neonatal Thrombocytopenia. Newborn Services Clinical Guideline - Auckland District Health Board.

http://www.adhb.govt.nz/newborn/Guidelines/Blood/Platelets/NeonatalThrombocytopenia.htm. Published 2016. Accessed July 15, 2016.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

All

Hematology/Thrombocytopenia and Platelet Dysfunction





Item 33

Do not transfuse neonates with mild thrombocytopenia (platelet count 100-150x10⁹/L) even if bleeding.

Grade of recommendations: Review

Stop

Roberts I, Murray N. Neonatal thrombocytopenia: causes and management. *Arch Dis Child - Fetal Neonatal Ed.* 2003;88(5):359F-364. doi:10.1136/fn.88.5.F359.

Clinical Practice Committee. Neonatal Thrombocytopenia. Newborn Services Clinical Guideline - Auckland District Health Board.

http://www.adhb.govt.nz/newborn/Guidelines/Blood/Platelets/NeonatalThrombocytopenia.htm. Published 2016. Accessed July 15, 2016.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

ΑII

Hematology/Thrombocytopenia and Platelet Dysfunction

Item 34

Start intravenous immunoglobulin (IVIG) only in case of severe thrombocytopenia (platelet count of <50x109/L) or if bleeding persists despite compatible platelets transfusion or in combination with unmatched platelets transfusion in neonates with neonatal <u>allo</u>-immune thrombocytopenia (NAIT).

Recommended dose of IVIG is 1 g/kg, which can be repeated 24h after if thrombocytopenia persists.

Start / Stop

Grade of recommendations: Institutional Guidelines

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

Peterson J, McFarland J, Curtis BR, Aster R. Neonatal alloimmune thrombocytopenia: pathogenesis, diagnosis and management. *Br J Haematol.* 2013;161(1):3-14. doi:10.1111/bjh.12235.

Neonatal Formulary. Immunoglobulin. *Neonatal Formulary*. http://www.neonatalformulary.com/pdfs/commentary/IMMUNOGLOBULIN-(commentary).pdf. Published 2014. Accessed July 19, 2016.

ΑII

Hematology/Thrombocytopenia and Platelet Dysfunction





Item 35

Start intravenous immunoglobulin (IVIG) as first line treatment in neonates with neonatal auto-immune thrombocytopenia and born to mothers who have idiopathic thrombocytopenic purpura (ITP), when platelet count is of <30x109/L or clinical bleeding is present. Recommended dose of IVIG is 1g/kg. Platelets transfusions are less likely to be effective and should be used as an adjuvant treatment for those who exhibit severe bleeding. Start **Grade of recommendations: Institutional Guidelines** The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, Wong W, Glader B. Approach to the newborn who has thrombocytopenia. Neoreviews. 2004;5(10). doi:10.1542/neo.5-10-e444. ΑII Hematology/Thrombocytopenia and Platelet Dysfunction





Vasospasms and Thromboembolism

Item 36

Start

Start unfractionned heparin or low molecular weight heparin in neonates with a first event venous thromboembolism for at least 5 days.

Unfractionned heparin: 75 units/kg IV over 10 minutes, followed by 28 units/kg per hour continuous infusion. Four hours after initiating therapy, measure aPTT, then adjust dose to achieve an aPTT that corresponds to an anti-factor Xa level of 0.35 to 0.7 (this is usually equivalent to an aPTT of 60 to 85 seconds). Treatment should be limited to 10 to 14 days. Some experts recommend switching to low molecular weight heparin after 3 to 5 days. For renal vein thrombosis requiring treatment, 6 weeks to 3 months of heparin/low molecular weight heparin therapy is recommended.

Low molecular weight heparin: Eg: Enoxaparine:

Term infants: initial, 1.7 mg/kg per dose subcutaneous every 12 hours.

Preterm infants: initial, 2 mg/kg per dose subcutaneous every 12 hours.

Adjust dosage to maintain anti-factor Xa level between 0.5 and 1.0 unit/mL. It will usually take several days to attain levels in the target range.

Dosage requirements to maintain target anti-factor Xa levels in preterm infants are quite variable, ranging from 0.8 to 3 mg/kg every 12 hours.

Grade of recommendations: National Guidelines

Monagle P, Chan AKC, Goldenberg NA, et al. Antithrombotic therapy in neonates and children: Antithrombotic therapy and prevention of thrombosis, 9th ed: American college of chest physicians evidence-based clinical practice guidelines. *Chest.* 2012;141(2 SUPPL.):e737S–e801S. doi:10.1378/chest.11-2308.

Neofax. Heparin. *Micromedex*. http://neofax.micromedexsolutions.com/. Published 2016. Accessed July 21, 2016.

Neofax. Enoxaparin. *Micromedex.* http://neofax.micromedexsolutions.com/. Published 2016. Accessed July 21, 2016.

ΑII

Hematology/Vasospasms and Thromboembolism

Item 37

Start alteplase or urokinase only in case of major vessel occlusion causing critical compromise of organs or limbs in infants with venous thromboembolism.

If thrombolysis is required, tissue plasminogen activator (tPA) (alteplase) or urokinase can be used, and plasminogen (fresh frozen plasma [FFP]) administration is suggested prior to start therapy. Alteplase recommended doses for dissolution of intravascular thrombi: 200 mcg/kg per hour (0.2 mg/kg per hour). Duration of therapy is 6 to 48 hours. If administering directly into the thrombus, dose may be increased after 6 hours to a maximum of 500 mcg/kg per hour. If localized bleeding occurs, stop infusion for 1 hour and restart using 100 mcg/kg per hour. Discontinue heparin several hours prior to initiation of therapy. Use urokinase as follows: try a dose of 5000 unit/kg an hour, and consider increasing the dose two- or even four-fold if blood flow does not improve within 8 hours.

Start / stop

Grade of recommendations: National Guidelines

Monagle P, Chan AKC, Goldenberg NA, et al. Antithrombotic therapy in neonates and children: Antithrombotic therapy and prevention of thrombosis, 9th ed: American college of chest physicians evidence-based clinical practice guidelines. *Chest.* 2012;141(2 SUPPL.):e737S–e801S. doi:10.1378/chest.11-2308.

ΑII

Hematology/Vasospasms and Thromboembolism





4. PNEUMOLOGY

Oxygen saturation targets

The lowest oxygen saturation level recommended to commence oxygen therapy:

<36 weeks GA: commence oxygen when saturations fall below 90% in ambiant air ≥36 weeks GA: commence oxygen when saturations fall below 93% in ambiant air</p>

Definition Target oxyg

Target oxygen saturations and alarm limits for babies needing supplemental oxygen:

<36 weeks GA: 90-94%

≥36 weeks GA: 93-97% (except infants with Persistent Pulmonary Hypertention of the Newborn: ≥95%).

Austin N, Newborn Clinical Network. Practice recommendation for Oxygen saturation targets for newborns cared for in neonatal units, New Zealand. *The Paeditric Society New Zealand & The National Child & Youth Clinical Network.* 2015.. 2015.

Pneumothorax

Definition

Primary pneumothorax: pneumothorax without any obvious lung diseases.

Spontaneous pneumothorax (SP) is a form of primary pneumothorax in neonates. It usually occurs in the absence of inciting risk factors at birth.

Shaireen H, Rabi Y, Metcalfe A, et al. Impact of oxygen concentration on time to resolution of spontaneous pneumothorax in term infants: a population based cohort study. *BMC Pediatr*. 2014;14(1):1-8. doi:10.1186/1471-2431-14-208.

Item 38

Do not use routine supplemental oxygen use in infants with spontaneous pneumothorax.

In infants with pneumothorax and respiratory distress, oxygen supplementation should be provided as needed to maintain adequate saturation. The rate of recovery for spontaneous pneumothoraces is not improved with oxygen supplementation or nitrogen washout (60 to 100% inspired O2 concentration) which expose infants to the risks of hyperoxia.

Stop

Grade of recommendations: Cohort Study

Shaireen H, Rabi Y, Metcalfe A, et al. Impact of oxygen concentration on time to resolution of spontaneous pneumothorax in term infants: a population based cohort study. *BMC Pediatr*. 2014;14(1):1-8. doi:10.1186/1471-2431-14-208.

Austin N, Newborn Clinical Network. Practice recommendation for Oxygen saturation targets for newborns cared for in neonatal units, New Zealand. *The Paeditric Society New Zealand & The National Child & Youth Clinical Network.* 2015.. 2015.

≥37w GA Pneumology/Pneumothorax
Oxygen, nitrogen





Apnea of Prematurity

Clinically significant apnea of prematurity

Definition

Apnea is a cessation of breathing for 20 seconds or longer or a shorter pause accompanied by bradycardia (<100 beats per minute), cyanosis, or pallor.

Eichenwald EC, American Academy of Pediatrics Committee on Fetus and Newborn. Apnea of prematurity. *Pediatrics*. 2016;137(1):1-7.

Item 39

Start caffeine citrate in patients with apnea of prematurity (loading dose 20 mg/kg; maintenance dose 5mg/kg/day). Dose may be increased to 10 mg/kg/day if apnea persists.

Caffeine citrate is a safe and effective treatment of apnea of prematurity and improves neurodevelopmental outcomes at 2 years of age. When caffeine is not available, use theophylline treatment at a 5-6 mg/kg loading dose and 2-6 mg/kg/day maintenance dose, divided every 8-12h. 2mg of caffeine citrate contains 1 mg of caffeine.

Start

Grade of recommendations: Randomized Controlled Trial

Eichenwald EC, American Academy of Pediatrics Committee on Fetus and Newborn. Apnea of prematurity. *Pediatrics*. 2016;137(1):1-7.

Schmidt B, Roberts RS, Davis P, et al. Caffeine Therapy for Apnea of Prematurity. *The New England Journal of Medicine*. 2006; 354(20):2112–2121.

Schmidt B, Roberts RS, Davis P, et al. Long-Term Effects of Caffeine Therapy for Apnea of Prematurity. *New England Journal of Medicine*. 2007;357(19):1893–1902. https://doi.org/10.1056/NEJMoa073679

Schmidt B, Anderson PJ, Doyle LW, et al. Survival Without Disability to Age 5 Years for Apnea of Prematurity. *Jama*. 2012;307(3):275–282. https://doi.org/10.1001/jama.2011.2024

Henderson-Smart David J, Steer Peter A. Caffeine versus theophylline for apnea in preterm infants. *Cochrane Database Syst Rev.* 2010;(1). doi:10.1002/14651858.CD000273.pub2.

<37w GA

Pneumology/Apnea





Item 40

item 40	
	Reassess the need for caffeine citrate treatment.
	Timely discontinuation of methylxanthines is advised to avoid unnecessary delays in discharge. A clinically significant apnea event–free period before discharge of 7 days is recommended after methylxanthine discontinuation, although a longer period may be suitable for infants born at less than 26 weeks' gestation.
Stop	
	Grade of recommendations: National Guidelines
	Eichenwald EC, American Academy of Pediatrics Committee on Fetus and Newborn. Apnea of prematurity. <i>Pediatrics.</i> 2016;137(1):1-7.
	Doyle J, Davidson D, Katz S, et al. Apnea of prematurity and caffeine pharmacokinetics: potential impact on hospital discharge. <i>J Perinat.</i> 2016;36:141-144. doi:10.1038/jp.2015.167.
<37w GA	Pneumology/Apnea
	Caffeine citrate
Item 41	
	Reassess the indication of anti-gastroesophageal reflux therapy in neonates with apnea.
Stop	Evidence suggests that gastroesophageal reflux (GER) is not associated with apnea of prematurity, and treatment of presumed or proven GER solely for the reduction in apnea events is not supported by currently available evidence.
	Grade of recommendations: National Guidelines
	Eichenwald EC, American Academy of Pediatrics Committee on Fetus and Newborn. Apnea of prematurity. <i>Pediatrics</i> . 2016;137(1):1-7.
<37w GA	Pneumology/Apnea
	Esomeprazole, Omeprazole





Bronchopulmonary Dysplasia (BPD)

Bronchopulmonary Dysplasia (BPD) Gestational Age < 32 wk ≥ 32 wk > 28d but < 56d postnatal age or 36 wk PMA or discharge to discharge to home, whichever home, whichever comes first Time point of assessment comes first Treatment with oxygen > 21% for at least 28 d plus Breathing room air by 56d Breathing room air at 36 wk Mild BPD PMA or discharge, whichever postnatal age or discharge, comes first whichever comes first Need for 30% oxygen at 56 d Need for 30% oxygen at 36 wk postnatal age or discharge, PMA or discharge, whichever Moderate BPD whichever comes comes first first Need for ≥30% oxygen and/or Need for ≥30% oxygen and/or positive pressure (PPV or positive pressure (PPV or Severe BPD NCPAP) at 36 wk PMA or NCPAP) at 56d postnatal age or discharge, whichever comes first discharge, whichever comes first

Definition

Definition of abbreviations: BPD = bronchopulmonary dysplasia; NCPAP = nasal continuous positive airway pressure; PMA = postmenstrual age; PPV = positive-pressure ventilation

BPD usually develops in neonates being treated with oxygen and positive pressure ventilation for respiratory failure, most commonly respiratory distress syndrome. Persistence of clinical features of respiratory disease (tachypnea, retractions, rales) are considered common to the broad description of BPD and have not been included in the diagnostic criteria describing the severity of BPD. Infants treated with oxygen > 21% and/or positive pressure for nonrespiratory disease (e.g., central apnea or diaphragmatic paralysis) do not have BPD unless they also develop parenchymal lung disease and exhibit clinical features of respiratory distress. A day of treatment with oxygen > 21% means that the infant received oxygne > 21% for more than 12 h on that day. Treatment with oxygen > 21% and/or positive pressure at 36 wk PMA, or at 56 d postnatal age or discharge, should not reflect an "acute" event, but should rather reflect the infant's usual daily therapy for several days preceding and following 36 wk PMA, 56 d postnatal age, or discharge.

Jobe A, Bancalari E. Bronchopulmonary Dysplasia. *Am J Respir Crit Care Med.* 2001;163(7):1723-1729. doi:10.1164/ajrccm.163.7.2011060.

Item 42

Do not use dexamethasone in the prevention or the treatment of bronchopulmonary dysplasia. Administering high-dose dexamethasone to prevent or treat chronic lung disease (CLD) is not recommended. The routine use of low-dose dexamethasone for all infants who require assisted ventilation after seven days of age to prevent or treat CLD is not recommended. Grade of recommendations: National Guidelines Jefferies A, Canadian Paediatric Society. Treat of prevent chronic lung disease in preterm infants. Paediatr Child Health. 2012;17(10):573. Pneumology/BPD AND Pneumology/Prevention of BPD OR Prevention/Prevention of BPD





Dexamethasone

Item 43

	Do not use loop diuretics for prevention of BPD in preterm neonates.
	Current evidence does not support the use of loop diuretics for prevention of BPD.
Stop	
Оюр	Grade of recommendations: Review
	Tropea K, Christou H. Current pharmacologic approaches for prevention and treatment of bronchopulmonary dysplasia. <i>Int J Pediatr.</i> 2012;2012:598606. doi:10.1155/2012/598606.
All	Pneumology/Prevention of BPD OR Prevention/Prevention of BPD
	Furosemide, torasemide
Item 44	
	Do not use thiazid diuretics for prevention of BPD in preterm neonates. Use them judiciously for treatment of BPD in preterm neonates.
Stop	No clear evidence is present for use of thiazide diuretics for the prevention or management of BPD. In patient with BPD, thiazide and spironolactone were shown to decreased oxygen requirement and improved lung function in the treatment group compared to placebo but failed to show any improvement in the survival rate, duration of oxygen requirement, or length of hospital stay.
	Grade of recommendations: Review
	Tropea K, Christou H. Current pharmacologic approaches for prevention and treatment of bronchopulmonary dysplasia. <i>Int J Pediatr.</i> 2012;2012:598606. doi:10.1155/2012/598606.
<37w GA	Pneumology/BPD AND Pneumology/Prevention of BPD OR Prevention/Prevention of BPD
	Hydrochlorthiazid, chlorthalidone, spironolactone





Respiratory Distress Syndrome (= Hyaline Membrane Disease)

Respiratory Distress Syndrome (RDS)

Definition

 PaO_2 <50 mmHg (<6.6 kPa) in room air, central cyanosis in room air or need for supplemental oxygen to maintain PaO_2 >50 mmHg (>6.6 kPa), as well as the classical chest X-ray appearances.

Sweet DG, Carnielli V, Greisen G, et al. European consensus guidelines on the management of neonatal respiratory distress syndrome in preterm infants-2013 update. *Neonatology*. 2013;103(4):353-368.

Item 45

Start surfactant therapy in infants born <26 weeks of gestational age who need FiO2 >0.30.

In infants born <26 weeks of gestational age who need FiO2 >0.30, porcine-derived surfactant poractant alpha therapy is recommended at a dose of 200 mg/kg. In infants with signs of persistant RDS and respiratory support, give 100 mg/kg 6-12 hours after the first dose and 12 hours after the second dose. Maximum dose is 400 mg/kg. Administering more than three doses of surfactant has not shown to have a benefit.

Start

Grade of recommendations: International Guidelines

Sakonidou S, Dhaliwal J. The management of neonatal respiratory distress syndrome in preterm infants (European Consensus Guidelines—2013 update). *Arch Dis Child - Educ Pract Ed.* 2015;100(5):257-259. doi:10.1136/archdischild-2014-306642.

Davis DJ, Barrington KJ, Canadian Paediatric Society. Recommendations for neonatal surfactant therapy. *Paediatr Child Health (Oxford)*. 2015;10(2):109-116.

<26w GA Pneumology/RDS

Item 46

Start surfactant therapy in infants born ≥26 weeks of gestational age who need FiO2 >0.40.

In infants born ≥26 weeks of gestational age who need FiO2 >0.40, porcine-derived surfactant poractant alpha therapy is recommended at a dose of 200mg/kg. In infant with signs of persistant RDS and respiratory support, give 100 mg/kg 6-12 hours after the first dose and 12 hours after the 2nd dose. Maximum dose is 400 mg/kg. Administering more than three doses of surfactant has not shown to have a benefit.

Start

Grade of recommendations: International Guidelines

Sakonidou S, Dhaliwal J. The management of neonatal respiratory distress syndrome in preterm infants (European Consensus Guidelines—2013 update). *Arch Dis Child - Educ Pract Ed.* 2015;100(5):257-259. doi:10.1136/archdischild-2014-306642.

Davis DJ, Barrington KJ, Canadian Paediatric Society. Recommendations for neonatal surfactant therapy. *Paediatr Child Health (Oxford)*. 2015;10(2):109-116.

≥26w GA Pneumology/RDS





Meconium Aspiration Syndrome

D - 6::4:	Meconium Aspiration Syndrome (MAS)
Definition	Presence of respiratory distress and chest X-ray changes, not explained by other
	pathology, where there has been meconium stained amniotic fluid prior to delivery.
	Stenson BJ, Smith CL. Management of meconium aspiration syndrome. <i>Paediatr Child Heal (United Kingdom)</i> . 2012;22(12):532-535. doi:10.1016/j.paed.2012.08.015.
Item 47	Ringuomij. 2012,22(12):552-555. doi:10.1010/j.paed.2012.00.015.
item 47	Consider inhaled nitric oxide (iNO) in neonates with hypoxic respiratory
	· /
	failure due to MAS.
	Frank was in a sainten of the sainten to MAO infrate account of the sainten of th
	For hypoxic respiratory failure due to MAS, infants responded well to combined iNO and high frequency ventilation treatment in comparison to either iNO or high frequency
Start	ventilation.
	veritination.
	Grade of recommendations: Review
	0.000
	Swarnam K, Soraisham AS, Sivanandan S. Advances in the Management of Meconium Aspiration
	Syndrome. Int J Pediatr. 2012;2012:1-7. doi:10.1155/2012/359571.
All	Pneumology/MAS
Item 48	
	Reassess the indication for antibiotics in patients with MAS alone.
	Prophylactic use of antibiotics in meconium aspiration syndrome is not recommended
	unless there is an identified risk of infection.
Stop	
	Grade of recommendations: Review
	Swarnam K, Soraisham AS, Sivanandan S. Advances in the Management of Meconium Aspiration
• "	Syndrome. Int J Pediatr. 2012;2012:1-7. doi:10.1155/2012/359571.
All	Pneumology/MAS Antibiotics
Item 49	Anubious
116111 49	Administer a bolus instillation of surfactant in intubated infants with MAS
Start	
	requiring more than 50% oxygen.
	In infants with MAC symfostant administration may reduce the according of requiretem illness
	In infants with MAS, surfactant administration may reduce the severity of respiratory illness and decrease the number of infants with progressive respiratory failure requiring support
	with extracorporeal membrane oxygenation (ECMO) (A). At the time of review, more trials
	are needed to evaluate the place of diluted surfactant as lavage therapy in MAS, and no
	recommendation can be made (FRN).
	Grade of recommendations: A
	Davis DJ, Barrington KJ, Canadian Paediatric Society. Recommendations for neonatal surfactant
	therapy. Paediatr Child Health (Oxford) . 2015;10(2):109-116.
	El Shahed A, Dargaville PA, Ohlsson A, Soll R. Surfactant for meconium aspiration syndrome in
	term and late preterm infants. Cochrane Database Syst Rev. 2014;(12). doi:10.1002/14651858.CD002054.pub3.
	Hahn S, Choi HJ, Soll R, Dargaville PA. Lung lavage for meconium aspiration syndrome in newborn infants. <i>Cochrane Database Syst Rev.</i> 2013;(4):CD003486. doi:10.1002/14651858.CD003486.pub2.
All	Pneumology/MAS





Persistent Pulmonary Hypertension of the Newborn (PPHN)

Persistent Pulmonary Hypertension of the Newborn (PPHN)

Definition

Elevated pulmonary vascular resistance and right-left shunt through the ductus arteriosus and/or foramen ovale, and absence of congenital heart abnormalities, demonstrated by echocardiography.

Severe PPHN

PPHN with a oxygenation index ≥25.

Cabral JEB, Belik J. Persistent pulmonary hypertension of the newborn: Recent advances in pathophysiology and treatment. *J Pediatr (Rio J)*. 2013;89(3):226-242. doi:10.1016/j.jped.2012.11.009.

Adams JM, **Stark AR**. Persistent pulmonary hypertension of the newborn. *UpToDate.com*. http://www.uptodate.com/contents/persistent-pulmonary-hypertension-of-the-newborn. Published 2013. Accessed April 1, 2016.

Item 50

Start inhaled nitric oxide (iNO) in neonates who have severe PPHN.

Inhaled nitric oxide (iNO) is indicated to reduce the need for extracorporeal membrane oxygenation (ECMO) support in term and near-term infants with severe PPHN. Currently, the initial recommended concentration of iNO is 20 ppm. Using higher concentrations is not anymore effective, and is associated with a higher incidence of methemoglobinemia and formation of nitrogen dioxide (D). Once oxygenation improves, iNO can usually be weaned relatively rapidly to 5 ppm, and discontinued within 5 days. It should be weaned gradually in steps to the lowest dose possible for a period before discontinuation (D).

Start

Grade of recommendations: A

American Heart Association, American Thoracic Society, Abman SH, et al. Pediatric pulmonary hypertension. *Circulation*. 2015;132:2037-2099. doi:10.1161/CIR.000000000000329.

Cabral JEB, Belik J. Persistent pulmonary hypertension of the newborn: Recent advances in pathophysiology and treatment. *J Pediatr (Rio J)*. 2013;89(3):226-242. doi:10.1016/j.jped.2012.11.009.

All

Pneumology/PPHN

Item 51

Do not use sildenafil as initial therapy for PPHN.

Sildenafil is not recommended as initial therapy for PPHN when inhaled nitric oxyde is available. Sildenafil can be used as adjunctive therapy for infants with PPHN who are refractory to iNO or to attenuate rebound pulmonary hypertension after iNO withdrawal or to shorten the time to extubation.

Stop

Grade of recommendations: FRN

Shah PS, Ohlsson A. Sildenafil for pulmonary hypertension in neonates. *Cochrane database Syst Rev.* 2011;(8). doi:10.1002/14651858.CD005494.pub3.

American Heart Association, American Thoracic Society, Abman SH, et al. Pediatric pulmonary hypertension. *Circulation*. 2015;132:2037-2099. doi:10.1161/CIR.000000000000329.

All

Pneumology/PPHN





05. NEPHROLOGY Acute Kidney Injury (AKI)

Definition of acute kidney injury (AKI) Stage | Serum Creatinine (SCr) No change in SCr or rise <0.3 mg/dL 1.5-1.9 times reference SCr (lowest previous SCr value) ≥0.3 mg/dl (≥26.5 µmol/l) increase within 48h 2.0-2.9 times reference SCr Definition 3.0 times reference SCr OR Increase in serum creatinine to ≥4.0 mg/dL (≥353.6 3 OR SCr ≥2.5 mg/dL OR Initiation of dialysis Selewski DT, Charlton JR, Jetton JG, et al. Neonatal Acute Kidney Injury. Pediatrics. 2015;136(2):e463. doi:10.1542/peds.2014-3819.

Item 52

Do not use nephrotoxic drugs in neonates if possible, especially in preterm infants.

Nephron mass is lower in preterm infants since nephrogenesis is active until 36 weeks of gestational age and interruption of gestation results in a loss of total nephron number. Moreover, preterm infants are more vulnerable to acute kidney injury (AKI) with the potential loss of nephrons after birth. Nonsteroidal anti-inflammatory drugs (NSAIDs), including indomethacin and ibuprofen, paracetamol, aspirin, aminoglycosides (gentamicin, amikacin, tobramycin) and glycopeptide antibiotics (vancomycin, teicoplanin), betalactams (penicillins, cephalosporins), amphotericin B, antiviral agents (aciclovir), diuretics, proton pump inhibitors, and phenytoin can be nephrotoxic and cause AKI in neonates. This list is not exhaustive. When nephrotoxic agents must be started, monitor cystatin-C and/or serum creatinine before and after the initiation of treatment.

Stop

Grade of recommendations: Review

Rodieux F, Wilbaux M, van den Anker JN, Pfister M. Effect of Kidney Function on Drug Kinetics and Dosing in Neonates, Infants, and Children. *Clin Pharmacokinet*. 2015;54(12):1183-1204. doi:10.1007/s40262-015-0298-7.

Abitbol CL, Seeherunvong W, Galarza MG, et al. Neonatal kidney size and function in preterm infants: What is a true estimate of glomerular filtration rate? *J Pediatr.* 2014;164(5):1026-1031.e2. doi:10.1016/j.jpeds.2014.01.044.

Naughton CA. Drug-induced nephrotoxicity. *Am Fam Physician*. 2008;78(6):743-750. doi:10.1016/S0025-7125(05)70541-1.

All

Nephrology/Acute Kidney Injury (AKI) OR Prevention/Acute Kidney Injury (AKI)





Item 53

Stop	Stop all nephrotoxic drugs when possible in neonates with AKI (stage 1-3).
	Nonsteroidal anti-inflammatory drugs (NSAIDs), including indomethacin and ibuprofen, paracetamol, aspirin, aminoglycosides (gentamicin, amikacin, tobramycin) and glycopeptide antibiotics (vancomycin, teicoplanin), amphotericin B, antiviral agents (aciclovir), diuretics, and phenytoin can be nephrotoxic and cause AKI in neonates. This list is not exhaustive.
	Grade of recommendations: D
	Rodieux F, Wilbaux M, van den Anker JN, Pfister M. Effect of Kidney Function on Drug Kinetics and Dosing in Neonates, Infants, and Children. <i>Clin Pharmacokinet</i> . 2015;54(12):1183-1204. doi:10.1007/s40262-015-0298-7.
All	Nephrology/Acute Kidney Injury (AKI)
	Nephrotoxics
Item 54	
Other	Consider dosage adjustement for drugs highly excreted by renal elimination in neonates with AKI (stage 1-3). When needed, refer to a specialist.
	Grade of recommendations: D
	Grade of recommendations: D





6. GASTROENTEROLOGY

Direct hyperbilirubinemia (Conjugated Hyperbilirubinemia)

Definition

Direct hyperbilirubinemia

Direct bilirubinemia >17 μ mol/L if total bilirubin is <85.5 μ mol/L, or a value of direct bilirubin that represents >20% of the total bilirubin if total bilirubin is >85.5 μ mol/L.

Moyer V, Freese DK, Whitington PF, et al. Guideline for the evaluation of cholestatic jaundice in infants: recommendations of the North American Society for Pediatric Gastroenterology, Hepatology and Nutrition. *J Pediatr Gastroenterol Nutr.* 2004;39(2):115-128. doi:10.1097/00005176-200408000-00001.

Item 55

Decrease or stop intravenous soybean-based lipid emulsion in neonates with marked progressive cholestasis associated with parenteral nutrition.

Consider switching soybean-based lipid emulsion to fish oil-based lipid emulsions or emulsions with reduced omega-6 fatty acids and increased omega-3 fatty acids. Exemples of these type of lipid emulsions are Omegaven® (fish-oil), SMOFlipid® (soy-oil, medium-chain triglycerides, olive-oil, omega-3 fatty acids) and Lipoplus® (soy-oil, medium-chain triglycerides, omega-3 fatty acids).

Others

Grade of recommendations: International Guidelines

Dani C, Pratesi S, Raimondi F, Romagnoli C. Italian guidelines for the management and treatment of neonatal cholestasis. *Ital J Pediatr.* 2015;41(69):1-12. doi:10.1186/s13052-015-0178-7.

Lauriti G, Zani A, Aufieri R, et al. Incidence, Prevention, and Treatment of Parenteral Nutrition—Associated Cholestasis and Intestinal Failure—Associated Liver Disease in Infants and Children: A Systematic Review. *J Parenter Enter Nutr.* 2014;38(1):70-85. doi:10.1177/0148607113496280.

Koletzko B, Goulet O, Hunt J, et al. Guidelines on Paediatric Parenteral Nutrition of the European Society of Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) and the European Society for Clinical Nutrition and Metabolism (ESPEN), Supported by the European Society of Paediatric Research (ESPR). *J Pediatr Gastroenterol Nutr.* 2005;41(Suppl 2):S1-S4. doi:10.1097/01.mpg.0000181841.07090.f4.

All

Gastroenterology/Direct hyperbilirubinemia Total Parenteral Nutrition (TPN), lipid emulsion





Item 56

	Administer adequate protein intake of 2 to 3 g/kg/day to neonates with direct hyperbilirubinemia.
Start	
Otait	Grade of recommendations: Review
	Feldman AG, Sokol RJ. Neonatal Cholestasis. Neoreviews - <i>Am Acad Pediatr.</i> 2013;14(2). doi:10.1542/neo.14-2-e63.
All	Gastroenterology/Direct hyperbilirubinemia

Item 57

Start fat-soluble vitamins (ADEK) in neonates with cholestasis.

Prescribe fat-soluble vitamins during cholestasis and for 3 months following resolution of jaundice; doses will require daily monitoring. Follow your institution guidelines for dosage.

- Vitamin A: monitor serum vitamin A
- Vitamin D: Monitor bone biochemistry
- Vitamin E: monitor serum vitamin E
- Vitamin K: monitor PT and APTT

Start

Grade of recommendations: National Guidelines

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

Dani C, Pratesi S, Raimondi F, Romagnoli C. Italian guidelines for the management and treatment of neonatal cholestasis. *Ital J Pediatr.* 2015;41(69):1-12. doi:10.1186/s13052-015-0178-7.

All

Gastroenterology/Direct hyperbilirubinemia





Item 58

Consider ursodeoxycholic acid (UDCA) in neonate with direct hyperbilirubinemia. Consider ursodeoxycholic acid (UDCA) treatment at the dosage of 20-30 mg/kg/day in divided doses until jaundice resolve. Ursodeoxycholic acid (UDCA) has been found to have beneficial effects on many forms of cholestasis, and is generally used as first-line therapy for pruritus due to cholestasis, parenteral nutrition-induced cholestasis, biliary atresia after surgical treatment, and α1-antitrypsin deficiency (C). Start **Grade of recommendations: National Guidelines** Dani C, Pratesi S, Raimondi F, Romagnoli C. Italian guidelines for the management and treatment of neonatal cholestasis. Ital J Pediatr. 2015;41(69):1-12. doi:10.1186/s13052-015-0178-7. The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016. ΑII Gastroenterology/Direct hyperbilirubinemia





	Indirect hyperbilirubinemia (Unconjugated Hyperbilirubinemia)	
Item 59		
	Administer intravenous immunoglobulin (IVIG) to neonates with a positive direct Coombs test and severe hyperbilirubinemia, or to those progressing to severe hyperbilirubinemia despite initial treatment.	
Start	IVIG should be administered, if not so already, in infants with isoimmunisation. Dose: 1 g/kg	
	Grade of recommendations: National Guidelines	
	Barrington KJ, Sankaran K, Canadian Paediatric Society Fetus and Newborn. Guidelines for detection, management and prevention of hyperbilirubinemia in term and late preterm. <i>Paediatr Child Heal.</i> 2007;12(Suppl B):1B-12B.	
All	Gastroenterology/Indirect hyperbilirubinemia	





Necrotizing Enterocolitis (NEC)

Necrotizing Enterocolitis Review of Bell's **Clinical Findings** Radiographic Gastrointestinal Bell's stages **Stages Findings** Findings Stage I Normal gas pattern Apnea and Gastric residuals, bradycardia, occult blood in stool, or mild ileus Suspect mild abdominal temperature instability distention Stage II A Apnea and lleus gas pattern Grossly bloody bradycardia, stools, prominent with one or more temperature dilated loops and abdominal instability focal pneumatosis distention, absent bowel sounds Definition Proven Stage II B Thrombocytopenia Widespread Abdominal wall and mild metabolic pneumatosis, edema with palpable ascites, portalloops and venous gas tenderness Stage III A Mixed acidosis, Prominent bowel Worsening wall oliguria, loops, worsening edema, erythema hypotension, ascites, no free air and induration coagulopathy Advanced Stage III B Shock, deterioration Pneumo-Perforated bowel in laboratory values peritoneum and vital signs

Walsh MC, Kliegman RM. Necrotizing enterocolitis: treatment based on staging criteria. *Pediatr Clin North Am.* 1986;33(1):179-201.

Bell MJ, Ternberg JL, Feigin RD, et al. Neonatal necrotizing enterocolitis. Therapeutic decisions based upon clinical staging. *Ann Surg* . 1978;187(1):1-7.

Item 60

Start probiotics in preterm neonates at high risk of developing NEC.

In patient at high risk of developing NEC (preterm < 32 weeks GA or <1500g), initiate a probiotic treatment, with a preparation combining Lactobacillus and Bifidobacterium species. Initiate at the time of the first feed until 36 weeks of gestational age or discharge. Use only probiotic drugs fulfilling pharmaceutical regulations.

Start

Grade of recommendations: Systematic Review / Meta-analysis

AlFaleh K, Anabrees J. Probiotics for prevention of necrotizing enterocolitis in preterm infants. *Cochrane database Syst Rev.* 2014;(4). doi:10.1002/14651858.CD005496.pub4.

ProPrems Study Group, Jacobs SE, Tobin JM, et al. Probiotic effects on late-onset sepsis in very preterm infants: a randomized controlled trial. *Pediatrics*. 2013;132(6):1055-1062. doi:10.1542/peds.2013-1339.

Deshpande GC, Rao SC, Keil AD, Patole SK. Evidence-based guidelines for use of probiotics in preterm neonates. *BMC Med.* 2011;9(1):92. doi:10.1186/1741-7015-9-92.

<32w GA and/or <1500g

Prevention/Prevention of NEC OR Gastroenterology/Necrotizing enterocolitis (NEC)





Item 61	
	Stop all enteral medications in neonates suspected to have NEC.
Stop	When an infant is suspected to have NEC (stage I), all enteral medications should be discontinued. Enteral route can be used again if investigation exclude NEC. In stage II-III NEC, enteral route must not be used for 7-14 days to allow gastrointestinal rest.
	Grade of recommendations: Review
	Sharma R, Hudak M. A clinical perspective of necrotizing enterocolitis: past, present, and future. <i>Clin Perinatol.</i> 2013;40(1):27-51. doi:10.1016/j.clp.2012.12.012.
All	Gastroenterology/Necrotizing enterocolitis (NEC) All oral treatments
Item 62	
	Do not use enteral antibiotics for the prevention of NEC.
Stop	Evidence suggests that enteral antibiotics reduce the incidence of NEC in low birth weight infants. However, concerns about adverse outcomes persist, particularly related to the development of resistant bacterial infection.
	Grade of recommendations: Systematic Review / Meta-analysis
	Bury RG, Tudehope D. Enteral antibiotics for preventing necrotizing enterocolitis in low birthweight or preterm infants. <i>Cochrane Database Syst Rev.</i> 2001;(1). doi:10.1002/14651858.CD000405.
<32w GA and/or <1500g	Prevention/Prevention of NEC OR Gastroenterology/Necrotizing enterocolitis (NEC)
Item 63	Antibiotics, immunoglobulin
item 63	Start broad spectrum antibiotic promptly after blood cultures have been
	drawn in neonates with any stage of NEC.
	After blood cultures have been drawn, prompt initiation of treatment with IV gentamicin and amoxicillin. If evolving to Bell's stage II-IV NEC, antibiotic treatment should be continued for 7 to 14 days. Follow your institution guideline for dosage.
Start	Grade of recommendations: Textbook
	Sharma R, Hudak M. A clinical perspective of necrotizing enterocolitis: past, present, and future. <i>Clin Perinatol.</i> 2013;40(1):27-51. doi:10.1016/j.clp.2012.12.012. Schanler R, Abrams SA, Kim MS. Management of necrotizing enterocolitis in newborns. <i>UpToDate.com.</i> http://www.uptodate.com/contents/management-of-necrotizing-enterocolitis-innewborns. Published 2015. Accessed April 25, 2016. Lin PW, Stoll BJ. Necrotising enterocolitis. <i>Lancet.</i> 2006;368:1271-1283. doi:10.1016/S0140-6736(06)69525-1.
All	Gastroenterology/Necrotizing enterocolitis (NEC)





	Gastrointestinal Bleeding from the Upper Tract	
Item 64		
Other	Check that a Vitamin K prophylaxis was administered postdelivery in neonates with upper gastro-intestinal bleeding, to guide diagnostic.	
	All neonates who have hematemesis should be screened for coagulopathy due to: - failure to administer prophylaxis postdelivery - maternal thrombocytopenic purpura - hemophilia - von Willebrand disease	
	Grade of recommendations: Review Boyle JT. Gastrointestinal bleeding in infants and children. <i>Pediatr Rev.</i> 2008;29(2):39-52. doi:10.1542/pir.29-2-39.	
All	Gastroenterology/Gastrointestinal Bleeding from the Upper Tract	





Gastroesophageal Reflux

Gastroesophageal Reflux (GER)

Definition

Gastroesophageal reflux is physiologic in the neonate. Only rarely does reflux become a "disease" (GERD).

Gastresophageal Reflux Disease (GERD)

Cause overt oesophagitis or is associated with other symptoms. This should be assessed by clinical judgment.

Health.vic. Gastro-oesophageal reflux (GOR) in neonates. *Victoria State Government*. https://www2.health.vic.gov.au/hospitals-and-health-services/patient-care/perinatal-reproductive/neonatal-ehandbook/conditions/gastro-oesophageal-reflux. Published 2016. Accessed lune 24, 2016.

Neonatal Formulary. Omeprazole Web Commentary. *Neonatal Formulary*. http://www.neonatalformulary.com/pdfs/commentary/OMEPRAZOLE-(commentary).pdf. Published 2014. Accessed June 24, 2016.

Item 65

Consider proton pump inhibitors or H2-blockers only in neonates with severe cases of acid gastroesophageal reflux disease (GERD), when non-pharmacological measures (including milk thickeners) have failed.

Do not offer acid-suppressing drugs, such as proton-pump inhibitors or H2-receptor antagonists, to treat overt regurgitation in infants with isolated symptoms. In severe cases of GERD, omeprazole should be administered orally at an initial dose of 0.7 mg/kg once a day. It can be raised to a maximum dose of 1.4 mg/kg once a day after 7-14 days if this does not inhibit gastric acid production. IV use: Give 0.5 mg/kg once a day over 5 min. Recommended dose of ranitidine is 1.5 mg/kg 3x/day for term infants and 0.5 mg/kg 3x/day for preterm infants. Treatment should be reassessed regularly.

Start / stop

Grade of recommendations: National Guidelines

National Institute For Health and Care Excellence. Managing gastro-oesophageal reflux and reflux disease in infants. *NICE Pathways.* https://pathways.nice.org.uk/pathways/dyspepsia-and-gastro-oesophageal-reflux-disease. Published 2016. Accessed June 24, 2016.

Neonatal Formulary. Omeprazole Web Commentary. *Neonatal Formulary*. http://www.neonatalformulary.com/pdfs/commentary/OMEPRAZOLE-(commentary).pdf. Published 2014. Accessed June 24, 2016.

Czinn SJ, Blanchard S. Gastroesophageal reflux disease in neonates and infants: when and how to treat. *Paediatr Drugs.* 2013;15:19-27. doi:10.1007/s40272-012-0004-2.

All

Gastroenterology/Gastroesophargeal reflux Proton pump inhibitors, H2-receptor antagonists





Item 66

Do not use metoclopramide, domperidone or erythromycin to treat gastroesophageal reflux or gastroesophageal reflux disease. Pro-kinetics such as metoclopramide and domperidone are not recommended for the treatment of GER due to lack of evidence and concerns regarding adverse effects. Erythromycin has limited benefit, may facilitate bacterial resistance and should not be routinely prescribed. **Grade of recommendations: National Guidelines** Stop Chakraborty M, Damodaran K, Barr S. Guidelines for the management of gastro-oesophageal reflux disease (GORD) in neonates. UHW Cardiff NICU. http://www.cardiffnicu.com/Portal/Nutrition/GORD.pdf. Published 2013. Accessed June 24, 2016. National Institute For Health and Care Excellence. Managing gastro-oesophageal reflux and reflux disease in infants. NICE Pathways. https://pathways.nice.org.uk/pathways/dyspepsia-and-gastrooesophageal-reflux-disease. Published 2016. Accessed June 24, 2016. The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016. All Gastroenterology/Gastroesophargeal reflux Metoclopramide, domperidone, erythromycin





7. NEUROLOGY

Seizures

A seizure is defined clinically as a paroxysmal alteration in neurologic function. This definition includes:

Definition

- 1. Epileptic seizures: phenomena associated with corresponding EEG seizure activity e.g. clonic seizures
- 2. Non-epileptic seizures: clinical seizures without corresponding EEG correlate e.g. subtle and generalized tonic seizures
- 3. EEG seizures: abnormal EEG activity with no clinical correlation.

WHO Collaborating Center For Training and Research in Newborn Care. Management of neonatal seizures. *Newbornwhocc.org.* http://www.newbornwhocc.org/2014_pdf/Neonatal seizures 2014.pdf. Published 2014. Accessed May 11, 2016.

Item 67

Start phenobarbital as the first line agent in neonates with either EEG diagnosed or clinically apparent seizures when prolonged or frequent.

Start

Phenobarbital should be used as the first line agent (at a loading dose of 20 mg/kg IV over 10-15 min and a maintenance dose of 2.5-5 mg/kg IV, IM or oral, once daily beginning 12-24h after loading dose) in neonates with either EEG diagnosed or clinically apparent seizures when prolonged (greater than 3 minutes), frequent (greater than 3 per hour). Phenobarbital is recommended as first-line treatment given its inclusion in the only RCT of first-line treatment of neonatal seizure, the fact that it is the most studied anti-epileptic medication in animals, and its historical precedence as the first-line antiepileptic drug for neonates. Use phenobarbital with caution since there is extremely limited evidence on its effect on long-term neonatal neurodevelopment.

Grade of recommendations: International Guidelines

WHO Collaborating Center For Training and Research in Newborn Care. Management of neonatal seizures. *Newbornwhocc.org.* http://www.newbornwhocc.org/2014_pdf/Neonatal seizures 2014.pdf. Published 2014. Accessed May 11, 2016.

Slaughter LA, Patel AD, Slaughter J. Pharmacological Treatment of Neonatal Seizures: A Systematic Review. *J Child Neurol.* 2013;28(3):351-364. doi:10.1177/0883073812470734.

All Neurology/Seizures





Item 68

item 68	
Start	Consider phenytoin or a benzodiazepine or lidocaine in neonates with
	persistant seizures, despite adequate phenobarbital treatment.
	In neonates who continue to have seizures despite administation of the maximal tolerated dose of phenobarbital, either phenytoin or a benzodiazepine or lidocaine may be used as the second-line agent for the control of seizures. The use of phenytoin or lidocaine requires cardiac monitoring facilities.
	Cuada of uncommondations: International Cuidalines
	Grade of recommendations: International Guidelines
	WHO Collaborating Center For Training and Research in Newborn Care. Management of neonatal seizures. <i>Newbornwhocc.org</i> . http://www.newbornwhocc.org/2014_pdf/Neonatal seizures 2014.pdf. Published 2014. Accessed May 11, 2016.
All	Neurology/Seizures
	Phenobarbital
Item 69	
	Stop antiepileptic drugs if seizure-free for >72 hours in neonates with normal neurological examination and/or normal electroencephalography.
Stop	In neonates in whom seizure control is achieved with a single antiepileptic drug, the drug can be discontinued abruptly without any tapering of doses. In neonates requiring more than one antiepileptic drugs for seizure control, the drugs may be stopped one by one, with phenobarbital being the last drug to be withdrawn.
	Grade of recommendations: International Guidelines
	WHO Collaborating Center For Training and Research in Newborn Care. Management of neonatal seizures. <i>Newbornwhocc.org.</i> http://www.newbornwhocc.org/2014_pdf/Neonatal seizures 2014.pdf. Published 2014. Accessed May 11, 2016.
All	Neurology/Seizures
	Phenobarbital, phenytoin, lidocaine, benzodiazepines
Item 70	
	Consider pyridoxine only in neonates with recurrent seizures with no obvious cause.
Start	If there are recurrent seizures with no obvious cause consider pyridoxine dependency. A therapeutic trial of pyridoxine IV 50 -100 mg may be helpful (this may be considered during EEG).
	Crade of recommendations: International Cuidelines
	Grade of recommendations: International Guidelines
	WHO Collaborating Center For Training and Research in Newborn Care. Management of neonatal seizures. <i>Newbornwhocc.org.</i> http://www.newbornwhocc.org/2014_pdf/Neonatal seizures 2014.pdf. Published 2014. Accessed May 11, 2016.
All	Neurology/Seizures





Pyridoxine

8. PAIN, SEDATION & NEONATAL ABSTINENCE SYNDROME Pain, Analgesia & Sedation

Item 71

Start pain management in neonates with non-pharmacological techniques (incl. Sucrose) if aproppriate.

If moderate-severe pain is evident (including post-surgery, severe illness, major injury, congenital malformations or palliative care), progress to pharmacological agents.

Stop

Grade of recommendations: Institutional Guidelines

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

ΑII

Pain, sedation & neonatal abstinence syndrome/Pain, Analgesia & Sedation Analgesics

Item 72

Start paracetamol in neonates who are still in pain despite adequate nonpharmacological interventions.

Recommended doses are 10-15 mg/kg orally or 20-25 mg/kg rectally administered every 6-8 hours. Recommendations for intravenous paracetamol, are a loading dose of 20 mg/kg, followed by 10 mg/kg every 6-8 hours. Maximum doses should not exceed 40 mg/kg/day for infants at 26–32 weeks PMA and 60 mg/kg/day for infants at 32–42 weeks PMA. Hepatotoxicity occurs very rarely, if ever, following routine administration of Start / stop paracetamol in neonates. Priority to the oral route.

Grade of recommendations: International Guidelines

Barrington K, Batton D, Finley G, Wallman C, Canadian Paediatric Society. Prevention and Management of Pain in the Neonate: An Update. Paediatr Child Heal. 2007;12(2):137-138. doi:10.1542/peds.2006-2277.

Ballantyne JC, Cousins MJ, Giamberardino MA, et al. Acute Pain Management in Newborn Infants. Pain Clin Updat. 2011;19(6):1-6.

ΑII

Pain, sedation & neonatal abstinence syndrome/Pain, Analgesia & Sedation Paracetamol, opioids





Item 73

Do not use nonsteroidal antiinflamatory agents (NSAID) as analgesics.

NSAIDs are not recommended for neonatal analgesia, as safer and more effective agents are available. Treatment should be switched to other pharmacologic classes.

Stop

Grade of recommendations: International Guidelines

Ballantyne JC, Cousins MJ, Giamberardino MA, et al. Acute Pain Management in Newborn Infants. *Pain Clin Updat.* 2011;19(6):1-6.

Batton DG, Barrington KJ, Wallman C, American Academy of Pediatrics, Canadian Paediatric Society. Prevention and Management of Procedural Pain in the Neonate: An Update. *Pediatrics*. 2006;118(5):2231-2241. doi:10.1542/peds.2006-2277.

ΑII

Pain, sedation & neonatal abstinence syndrome/Pain, Analgesia & Sedation

NSAID

Item 74

Start morphine as first line treatment for pain relief in neonates who are still in pain despite adequate non-pharmacological techniques and paracetamol treatment.

Morphine is recommended as the first-line strong opioid for the treatment of persistant moderate to severe pain in children with medical illnesses. There is insufficient evidence to recommend any alternative opioid in preference to morphine as the opioid of first choice. Intermittent dose 50-100 μ g/kg IV every 4-8 hours, Infusion dose 10-30 μ g/kg/h (for opioid naive patients). Start at the lower dose and titrate carefully to effect using small incremental doses. When opioids are administered for greater than 4 days, physical dependence and tolerance may develop. This means that higher opioid doses are required in order to maintain patient confort and that treatment should be weaned over a period of days at the rate of 10% of the prescribed dose per day, based on the clinical assessment of the neonate. Caution should be taken when treating newborns with opioids, especially preterm neonates, as they are more sensitive to opioids and are at risk for respiratory depression, apnea, hypotension and urinary retention.

Start / stop

Grade of recommendations: International Guidelines

World Health Organization. WHO Guidelines on the Pharmacological Treatment of Persisting Pain in Children with Medical Illnesses. Geneva, Switzerland: *World Health Organization*; 2012.

Anand KJ, International Evidence-Based Group for Neonatal. Consensus statement for the prevention and management of pain in the newborn. *Arch Pediatr Adolesc Med.* 2001;155(2):173-180.

 $\textbf{Lexicomp.} \ \ \textbf{Morphine: Pediatric drug information.} \ \textit{UpToDate.com.}$

http://www.uptodate.com/contents/morphine-pediatric-drug-

information?source=search_result&search=morphine&selectedTitle=2~150#F11443893. Published 2016. Accessed June 22, 2016.

Australian and New Zealand Neonatal Network. Assessment and Management of Neonatal Pain. *Best Pract Clin Guidel Assess.* 2007;(September):1-14. http://www.acnn.org.au/acnn-resources/clinical-guidelines/newborn-pain-practice-guideline-2007.pdf.

ΑII

Pain, sedation & neonatal abstinence syndrome/Pain, Analgesia & Sedation

Paracetamol, opioids





Item 75

Start opioids as first line treatment for postoperative analgesia, and use them as long as pain assessment scales deem necessary.

Morphine dosage: intermittent dose 50-100 μ g/kg IV every 4-8 hours, Infusion dose 10-30 μ g/kg/h (for opioid naive patients). Clinical titration using small incremental doses (5–20 μ g/kg) may be required When opioid are administered for greater than 4 days, physical dependence and tolerance may develop. This means that higher opioid doses are required in order to maintain patient confort and that treatment should be weaned over a period of days at the rate of 10% of the prescribed dose per day, based on the clinical assessment of the neonate. Caution should be taken when treating newborns with opioids, especially preterm neonates, as they are more sensitive to opioids and are at risk for respiratory depression, apnea, hypotension and urinary retention.

Start

Grade of recommendations: International Guidelines

Barrington K, Batton D, Finley G, Wallman C, Canadian Paediatric Society. Prevention and Management of Pain in the Neonate: An Update. *Paediatr Child Heal.* 2007;12(2):137-138. doi:10.1542/peds.2006-2277.

Anand KJ, International Evidence-Based Group for Neonatal. Consensus statement for the prevention and management of pain in the newborn. *Arch Pediatr Adolesc Med.* 2001;155(2):173-180.

Lexicomp. Morphine: Pediatric drug information. *UpToDate.com*. http://www.uptodate.com/contents/morphine-pediatric-drug-information?source=search_result&search=morphine&selectedTitle=2~150#F11443893. Published 2016. Accessed June 22, 2016.

Australian and New Zealand Neonatal Network. Assessment and Management of Neonatal Pain. *Best Pract Clin Guidel Assess.* 2007;(September):1-14. http://www.acnn.org.au/acnn-resources/clinical-guidelines/newborn-pain-practice-guideline-2007.pdf.

ΑII

Pain, analgesia & neonatal abstinence syndrome/Pain & analgesia





Item 76

Reassess the indication of morphine or fentanyl in chronically ventilated preterm neonates without pain.

In the absence of pain, discomfort or difficulties for improving gas exchange, use of continuous infusions of morphine or fentanyl in chronically ventilated preterm neonates is not recommended. There is insufficient evidence to recommend routine use of opioids in mechanically ventilated newborns. Opioids should be used selectively, when indicated by clinical judgment and evaluation of pain indicators. If sedation is required, morphine is safer than midazolam. Newborns, especially preterm neonates, are more sensitive to opioids and are at risk for respiratory depression, apnea, hypotension, and urinary retention

Stop

Grade of recommendations: National Guidelines

Barrington K, Batton D, Finley G, Wallman C, Canadian Paediatric Society. Prevention and Management of Pain in the Neonate: An Update. *Paediatr Child Heal.* 2007;12(2):137-138. doi:10.1542/peds.2006-2277.

Ballantyne JC, Cousins MJ, Giamberardino MA, et al. Acute Pain Management in Newborn Infants. *Pain Clin Updat.* 2011;19(6):1-6.

Bellù R, de Waal K, Zanini R. Opioids for neonates receiving mechanical ventilation: a systematic review and meta-analysis. Arch Dis Child Fetal Neonatal Ed. 2010;95:F241-51. doi:10.1136/adc.2008.150318.

<37w GA

Pain, sedation & neonatal abstinence syndrome/Pain, Analgesia & Sedation Morphine, fentanyl, opioids

Item 77

Do not use ketamine treatment for routine management of pain.

In spite of its theoretical advantages, ketamine is a potent anesthetic that has received minimal study in neonates. As such, it should only be used for surgery or highly invasive procedures.

Stop

Grade of recommendations: FRN

Ballantyne JC, Cousins MJ, Giamberardino MA, et al. Acute Pain Management in Newborn Infants. *Pain Clin Updat.* 2011;19(6):1-6.

Witt N, Coynor S, Edwards C, Bradshaw H. A Guide to Pain Assessment and Management in the Neonate. *Curr Emerg Hosp Med Rep.* 2016;4:1-10. doi:10.1007/s40138-016-0089-y.

All

Pain, sedation & neonatal abstinence syndrome/Pain, Analgesia & Sedation Ketamine





Neonatal Abstinence Syndrome (NAS)

Item 78

Consider non-pharmacological interventions for the initial management of all infants suspected of having or at risk of developing NAS. This may mitigate the need for medication.

Stop

Infants at risk for NAS should be monitored diligently during the initial days after birth. At present, the modified Finnegan scores remains the most common tool that is used. Start modified Finnegan scoring within 24h of birth and monitor score every 3-4h. Treatment is indicated when the average of three scores is ≥8 or when the average of two scores is ≥12.

Grade of recommendations: National Guidelines

Kocherlakota P, American Academy of Pediatrics. Neonatal Abstinence Syndrome. *Pediatrics*. 2014;134(2):e547-61. doi:10.1542/peds.2013-3524.

Wiles JR, Isemann B, Ward LP, et al. Current management of neonatal abstinence syndrome secondary to intrauterine opioid exposure. *J Pediatr.* 2014;165(3):440-446. doi:10.1016/j.jpeds.2014.05.010.

ΑII

Pain, analgesia & neonatal abstinence syndrome/NAS

Opioids

Item 79

Start morphine as the first line pharmacological treatment for NAS when opioids are used by the mother and supportive measures failed.

Start

Morphine is indicated when the average of three Modified Finnegan Scores is ≥ 8 on the scoring tool or when the average of two scores is ≥ 12 . Recommended doses for oral morphine are variable. Try 50 μ g/kg every 3–4 h, then 10% or 50 μ g increments to a maximum dose of 1300 μ g/kg/day. Phenobarbital should be considered at this point.

Grade of recommendations: National Guidelines

Kocherlakota P, American Academy of Pediatrics. Neonatal Abstinence Syndrome. *Pediatrics*. 2014;134(2):e547-61. doi:10.1542/peds.2013-3524.

Canadian Society of Pharmacology and Therapeutics, Dow K, Ordean A, et al. Neonatal Abstinence Syndrome: Clinical Practice Guidelines For Ontario. *J Popul Ther Clin Pharmacol*. 2012;19(3):e488-e506.

All

Pain, analgesia & neonatal abstinence syndrome/NAS





Item 80

Item 66	
	Start weaning of morphine as soon as Modified Finnegan scores are <8 for 24 to 48 hours in neonates with NAS.
Stop	Initiate weaning of morphine when Modified Finnegan scores are <8 for 24 to 48 hours by a 10% decrease of the total daily dose with each wean occurring no more frequently than every 48 to 72 hours. Morphine can be discontinued when scores are stable for 48 to 72 hours on a dose of 0.05 to 0.1 mg/kg/day.
· ·	
	Grade of recommendations: Institutional Guidelines
	Provincial Council for Maternal and Child Health. Neonatal Abstinence Syndrome (NAS). Perinatal Quality Collaborative of North Carolina. http://www.pqcnc.org/documents/nas/nasresources/NASGuidelines.pdf. Published 2012. Accessed June 21, 2016.
All	Pain, analgesia & neonatal abstinence syndrome/NAS
	Morphine
Item 81	·
TICHT OT	Do not use morphine in neonates with NAS when the drugs used by the mother are non-opioids.
Ston	If needed, use phenobarbital (see phenobarbital recommendations).
Stop	
	Grade of recommendations: D
All	Pain, analgesia & neonatal abstinence syndrome/NAS
	Morphine, opioids





09. INFECTIOLOGY Meningitis Item 82 Start empirical antibiotic treatment with high dose amoxicilline and gentamicin in neonates with diagnosed or strongly suspected meningitis. Follow your institution guidelines for dosage. Start **Grade of recommendations: Textbook** Département de l'enfant et de l'adolescent - HUG. Cahier de l'interne. Hôpitaux Univ Genève. 2015;(Décembre). Edwards MS, Baker CJ. Bacterial meningitis in the neonate: Treatment and outcome. UpToDate.com. http://www.uptodate.com/contents/bacterial-meningitis-in-the-neonate-treatment-andoutcome#H26. Published 2016. Accessed May 30, 2016. Infectiology/Meningitis ΑII Item 83 Check results of cerebro-spinal fluid (CSF) culture as soon as they are available in order to reassess the need for treatment or the choice of antibiotics in neonates with suspected meningitis treated with empirical antibiotics. If low clinical suspicion of meningitis, stop antibiotics after 48 hr if: - CSF glucose >2/3 simultaneous blood glucose and - CSF protein <1 g/L, culture results are negative and baby remains well. Other **Grade of recommendations: Institutional Guidelines** Polin RA, Committee on Fetus and Newborn, American Academy of Pediatrics. Management of Neonates With Suspected or Proven Early-Onset Bacterial Sepsis. Pediatrics. 2012;129(5):1006-1015. doi:10.1542/peds.2012-0541. The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1,

All Infectiology/Meningitis
Antibiotics



2016.



Item 84

Do not use corticosteroids for the treatment of neonates with suspected or confirmed bacterial meningitis. Reassess the corticosteroid indication.

At present, there is insufficient data to make a recommendation on the use of adjunctive corticosteroids in neonates with bacterial meningitis. Very low-quality data from two randomised controlled trials suggest that some reduction in death and hearing loss may result from use of adjunctive steroids alongside standard antibiotic therapy for treatment of patients with neonatal meningitis. Benefits are not yet seen with regards to a reduction in neurological consequences.

Stop

Grade of recommendations: FRN

Ogunlesi TA, Odigwe CC, Oladapo OT. Adjuvant corticosteroids for reducing death in neonatal bacterial meningitis. *Cochrane database Syst Rev.* 2015;11(11):CD010435. doi:10.1002/14651858.CD010435.pub2.

National Collaborating Centre for Women's and Children's Health. Bacterial meningitis and meningococcal septicaemia. *NICE Clin Guidel* 102. 2010;(September):271. doi:10.1136/bmj.c3209.

Tunkel AR, Hartman BJ, Kaplan SL, et al. Practice guidelines for the management of bacterial meningitis. *ClinInfectDis.* 2004;39(9):1267-1284. doi:10.1086/425368.

All Infectiology/Meningitis

Corticosteroids





Sepsis

Sepsis: According to the onset of age, neonatal sepsis is divided into early-onset sepsis (EOS) and late-onset sepsis (LOS). EOS reflects transplacental or, more frequently, ascending infections from the maternal genital tract, whereas LOS is associated with the postnatal nosocomial or community environment, with the peak incidence reported to be between the 10th and 22nd day of life. The onset of LOS is most frequently defined at 72 h after birth, a cut-off time point considered to adequately differentiate LOS from EOS in terms of the spectrum of causative pathogens

Early-Onset Sepsis (EOS): Onset of sepsis symptoms during the first 72 hours of life. Risk factors of EOS:

- Maternal group B Streptococcus colonisation;
- Signs of chorioamnionitis (maternal fever >38 °C plus at least two of the following symptoms: maternal leucocytosis (>15 G/I), foetal tachycardia (>160/min), uterine tenderness, foul-smelling amniotic fluid);
- Prolonged rupture of membranes (>18 hours before delivery),
- Preterm birth <37 weeks,
- Previous neonate with an invasive group B streptococcus infection;

Definition

- Suspected infection in a sibling in the case of a multiple pregnancy.

Late-Onset Sepsis: Onset of sepsis symptoms at 72 hours of life or later. Risk factors of LOS:

- Risk of infection is inversely related to gestational age and birth weight and directly related to severity of illness at birth, reflecting need for invasive interventions e.g. prolonged ventilation, central venous access and parenteral nutrition.
- Delayed introduction of enteral feeds is associated with higher infection rates
- Increased risk of sepsis after gut surgery especially if enteral feeds slow to establish e.g. post-gastroschisis or necrotising enterocolitis (NEC) with stoma

Symptoms of sepsis:

- -Tachypnoea, respiratory distress, apnoea;
- -Tachycardia/bradycardia, poor peripheral perfusion (i.e. capillary refill time >3 seconds), mottling;
- -Temperature instability (hyperthermia >38.0 °C or hypothermia <36.0 °C);
- -Lethargy, irritability, altered muscular tone or floppiness;
- -Vomiting, poor feeding.

Dong Y, Speer CP. Late-onset neonatal sepsis: recent developments. *Arch Dis Child Fetal Neonatal Ed.* 2015;100(3):F257-63. doi:10.1136/archdischild-2014-306213.

Stocker M, Berger C, McDougall J, Giannoni E, Swiss Society of Neonatology, Paediatric Infectious Disease Group of Switzerland. Recommendations for term and late preterm infants at risk for perinatal bacterial infection. Swiss Med Wkly. 2013;143(September):1-5. doi:10.4414/smw.2013.13873.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.





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Item 85	
	Do not use empirical antibiotic therapy for asymptomatic neonates with a single risk factor of infection (incl. mother with suspected chorioamnionitis or unexplained premature delivery).
	Only close observation for the first 48 hours is advised.
Stop	Grade of recommendations: National Guidelines
	Stocker M, Berger C, McDougall J, Giannoni E, Swiss Society of Neonatology, Paediatric Infectious Disease Group of Switzerland. Recommendations for term and late preterm infants at risk for perinatal bacterial infection. Swiss Med Wkly. 2013;143(September):1-5. doi:10.4414/smw.2013.13873. National Institute For Health and Care Excellence. Neonatal infection: antibiotics for prevention and treatment. NICE Clin Guidel 149. nice.org.uk/guidance/cg149. Published 2012. Accessed May 26, 2016.
All	Infectiology/Sepsis
	Antibiotics
Item 86	
	Start empirical antibiotic treatment after blood cultures have been drawn in all newborn infants with suggestive signs of neonatal infection.
	Use amoxicillin and gentamicin at doses recommended by your institution. There is a need
	for therapeutic drug monitoring for aminoglycoside therapy (aiming for a residual gentamicine blood concentration of 0.5-2 mg/L before 3rd dose).
Start	for therapeutic drug monitoring for aminoglycoside therapy (aiming for a residual
Start	for therapeutic drug monitoring for aminoglycoside therapy (aiming for a residual
Start	for therapeutic drug monitoring for aminoglycoside therapy (aiming for a residual gentamicine blood concentration of 0.5-2 mg/L before 3rd dose). Grade of recommendations: National Guidelines Stocker M, Berger C, McDougall J, Giannoni E, Swiss Society of Neonatology, Paediatric Infectious Disease Group of Switzerland. Recommendations for term and late preterm infants at risk for perinatal bacterial infection. Swiss Med Wkly. 2013;143(September):1-5. doi:10.4414/smw.2013.13873.
Start	for therapeutic drug monitoring for aminoglycoside therapy (aiming for a residual gentamicine blood concentration of 0.5-2 mg/L before 3rd dose). Grade of recommendations: National Guidelines Stocker M, Berger C, McDougall J, Giannoni E, Swiss Society of Neonatology, Paediatric Infectious Disease Group of Switzerland. Recommendations for term and late preterm infants at risk for perinatal bacterial infection. Swiss Med Wkly. 2013;143(September):1-5.





Item 87

Reassess the need for antibiotics after 48 hours in neonates treated empirically with antibiotics for suspected sepsis. This is important since clinical and laboratory signs of neonatal infection are nonspecific and symptomatic neonates are treated empirically. In most cases, based on the clinical course, negative culture results and laboratory parameters, a decision can be made to stop safely antibiotic therapy after this time (C-reactive protein (CRP) and procalcitonin (PCT) have a high negative predictive value and can be used to stop empirically started antibiotic therapy early). Frequently, prolonged antibiotic therapy (>5 days) causes an increased mortality and a higher incidence of necrotising enterocolitis in preterm infants. This emphasises the need to stop empirical treatment in the absence of proven infection as early as possible and at the latest after 48-72 hours. Stop **Grade of recommendations: National Guidelines** Stocker M, Berger C, McDougall J, Giannoni E, Swiss Society of Neonatology, Paediatric Infectious Disease Group of Switzerland. Recommendations for term and late preterm infants at risk for perinatal bacterial infection. Swiss Med Wkly. 2013;143(September):1-5. doi:10.4414/smw.2013.13873. National Institute For Health and Care Excellence. Antibiotics for neonatal infection. NICE quality standard 75. guidance.nice.org.uk/qs75. Published 2014. Accessed May 26, 2016. Brady MT, Polin RA. Prevention and Management of Infants With Suspected or Proven Neonatal Sepsis. Pediatrics. 2013;132(1):166-168. doi:10.1542/peds.2013-1310. ΑII Infectiology/Sepsis Item 88 Do not use cephalosporins as first-line treatment in infant with suspected neonatal infection, because of the high risk of developing resistance. Use is restricted to special cases

Stop

Grade of recommendations: National Guidelines

Stocker M, Berger C, McDougall J, Giannoni E, Swiss Society of Neonatology, Paediatric Infectious Disease Group of Switzerland. Recommendations for term and late preterm infants at risk for perinatal bacterial infection. Swiss Med Wkly. 2013;143(September):1-5. doi:10.4414/smw.2013.13873.

All Infectiology/Sepsis
Cephalosporins





Item 89

	Do not use intravenous immunoglobulin in the treatment of suspected or proven neonatal sepsis.
	Therapy with intravenous immunoglobulin had no effect on the outcomes of suspected or proven neonatal sepsis and should be discontinued.
Stop	proven neonatal sepsis and should be discontinued.
	Grade of recommendations: Randomized Controlled Trial
	The INIS Collaborative Group. Treatment of Neonatal Sepsis with Intravenous Immune Globulin. <i>N Engl J Med.</i> 2011;365(13):1201-1211. doi:10.1056/NEJMoa1100441.
All	Infectiology/Sepsis
	Intravenous immunoglobulin (IVIG)
Item 90	
	Do not use Vancomycin as prophylaxis against sepsis in preterm neonates.
Stop	Vancomycin prophylaxis should not be undertaken in preterm infants with additional risk factors for infection such as a birth weight less than 1500 grams, use of central venous catheters, and administration of intravenous hyperalimentation.
	Grade of recommendations: FRN
	Ap C, Finer N, Kj B, Craft AP, Finer N, Barrington KJ. Vancomycin for prophylaxis against sepsis in preterm neonates. <i>Cochrane Database Syst Rev.</i> 2000;(1). doi:10.1002/14651858.CD001971.
<37w GA	Infectiology/Sepsis OR Prevention/Sepsis
	Vancomycin





Hepatitis Item 91 Administer an initial dose of hepatitis-B vaccine within 12 hours of birth in infants born to HBsAg-positive mothers, including infants weighing <2000g. Administer Hepatitis-B immune globulins (HBIG) 200 IU concurrently but at a different anatomic site. The later HBIG is administered after exposure, the less it is effective. The interval of effectiveness is unlikely to exceed 7 days. For infants who weigh less than 2000g at birth, the initial vaccine dose should not be counted in the required 3-dose schedule. Give lowbirth-weight and premature babies full neonatal dose of hepatitis B vaccine. Monitor infants born <28 weeks of gestational age for 72h after HBIG. **Grade of recommendations: National Guidelines** Start Groupe de travail "Prévention de la transmission mère-enfant de l'hépatite B"; Commission fédérale pour les vaccinations; Office de la santé publique., Anderau R, Bachmann G, et al. Recommandations pour la prévention de la transmission mère-enfant de l'hépatite B. Paediatrica. 2007;18(2):20-26. Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012. Brook G, Soriano V, Bergin C. European guideline for the management of hepatitis B and C virus infections, 2010. Int J STD AIDS. 2010;21:669-678. doi:10.1258/ijsa.2010.010234. Frieden TR, Jaffe HW, Cono J, Richards CL, lademarco MF. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006. ΑII Infectiology/Hepatitis Item 92 Do not use early hepatitis-B vaccine in infants born to mothers whose HBsAg and HBeAg status is negative but with positive anti-HBs status (prior infection or at risk of infection). They should receive three hepatitis-B vaccine doses at 2, 3 and 4 months of life along with Stop other recommended vaccines (DTPa-IPV-Hib). **Grade of recommendations: Institutional Guidelines** Département de l'enfant et de l'adolescent - HUG. Cahier de l'interne. Hôpitaux Univ Genève.



2015;(Décembre).

Infectiology/Hepatitis





Human Immunodeficiency Virus (HIV) Item 93 Start HIV prophylaxis with zidovudine as close to birth as possible for at least 4 weeks or consider tritherapy in neonates born to HIV-infected mothers who did not follow proper antenatal treatment or whose viremia are detectable. Follow your institution guidelines for doses. Grade of recommendations: National Guidelines Start Commission fédérale pour la santé sexuelle (CFSS) en matière de prévention contre la transmission du VIH de la mère à l'enfant. Maladies transmissibles : Recommandations de la Commission fédérale pour la santé sexuelle (CFSS) en matière de prévention contre la transmission du VIH de la mère à l'enfant. Office Féréral de la Santé Publique (OFSP), Suisse. 2016;4:80-81. AIDSinfo. Recommendations for Use of Antiretroviral Drugs in Pregnant HIV-1-Infected Women for Maternal Health and Interventions to Reduce Perinatal HIV Transmission in the United States. U.S. Department of Health and Human Services (HHS). https://aidsinfo.nih.gov/contentfiles/lvguidelines/peri_recommendations.pdf. Published 2015. Accessed June 6, 2016. Département de l'enfant et de l'adolescent - HUG. Cahier de l'interne. Hôpitaux Univ Genève. 2015;(Décembre). ΑII Infectiology/HIV Item 94 Start tritherapy immediately in the neonate aged <72 hr if the mother is diagnosed postpartum with HIV infection. Follow your institution guidelines for the choice of molecule and doses. **Grade of recommendations: Institutional Guidelines** Start The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshireshropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.



ΑII



Infectiology/HIV

Respiratory Syncytial Virus (RSV) Item 95 Start respiratory syncytial virus prophylaxis with palivizumab in neonates with severe bronchopulmonary dysplasia (BPD). Use palivizumab 15 mg/kg/dose IM once a month during RSV season (~october till february). In those with moderate BPD, consider palivizumab prophylaxis. Palivizumab is not indicated for neonates with mild BPD. Start / Stop Grade of recommendations: National Guidelines Aebi C, Barazzone C, Hammer J, Kind C, Nadal D, Pfister RE. Consensus concernant la prévention des infections par le virus respiratoire syncitial (VRS) avec l'anticorps humanisé nonoclonal palivizumab (Synagis). Paediatrica. 2004;15(6):17-19. Infectiology/RSV AND Pneumology/BPD All Palivizumab Item 96 Start respiratory syncytial virus prophylaxis with palivizumab in neonates with haemodynamically significant congenital heart disease AND other associated risk factors. Use palivizumab 15 mg/kg/dose once every month during RSV season (~october till february). It is not advised to administer such prophylaxis in infants solely with haemodynamically significant congenital heart disease. Associated risk factors: cyanotic Start / stop heart disease, severe pulmonary hypertension and overt heart failure. **Grade of recommendations: National Guidelines** Aebi C, Barazzone C, Hammer J, Kind C, Nadal D, Pfister RE. Consensus concernant la prévention des infections par le virus respiratoire syncitial (VRS) avec l'anticorps humanisé

nonoclonal palivizumab (Synagis). Paediatrica. 2004;15(6):17-19.

Infectiology/RSV AND Cardiology/congenital heart failure

Palivizumab



ΑII



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Item 97	
	Dot not use respiratory syncytial virus prophylaxis with palivizumab routinely in preterm neonates.
Stop	This statement was made considering: - The modest efficacy of palivizumab; - That in Switzerland the outcomes of hospitalization due to RSV infection in former preterm neonates without additionnal risk factors did not differ substantially from those of term neonates; - The high cost of palivizumab.
	Grade of recommendations: National Guidelines
	Grade of recommendations. National Guidelines
	Aebi C, Barazzone C, Hammer J, Kind C, Nadal D, Pfister RE. Consensus concernant la prévention des infections par le virus respiratoire syncitial (VRS) avec l'anticorps humanisé nonoclonal palivizumab (Synagis). <i>Paediatrica</i> . 2004;15(6):17-19.
All	Infectiology/RSV AND Prevention/RSV
	Palivizumab
Item 98	
	Do not use palivizumab for the treatment of respiratory syncytial virus (RSV) infection. Stop the treatment, even if it was given before the infection.
Stop	Grade of recommendations: National Guidelines
	Aebi C, Barazzone C, Hammer J, Kind C, Nadal D, Pfister RE. Consensus concernant la prévention des infections par le virus respiratoire syncitial (VRS) avec l'anticorps humanisé nonoclonal palivizumab (Synagis). <i>Paediatrica</i> . 2004;15(6):17-19.
	American Academy of Pediatrics. Updated guidance for palivizumab prophylaxis among infants and young children at increased risk of hospitalization for respiratory syncytial virus infection. <i>Pediatrics</i> . 2014;134(2):415-420. doi:10.1542/peds.2014-1665.
	Robinson JL, Le Saux N, Canadian Paediatric Society. Preventing hospitalizations for respiratory syncytial virus infection. <i>Paediatr Child Health.</i> 2015;20(6):321-333.
All	Infectiology/RSV
	Dali: in was ab





Palivizumab

Toxoplasmosis

Confirmed Case:

- Clinical illness in a child with laboratory evidence of Toxoplasma gondii infection born to a mother with documented seroconversion during pregnancy (post-conceptually)
 OR
- > Laboratory confirmation of infection in the neonate with or without clinical illness:
- Detection of IgA and/or IgM antibodies to T. gondii from a single peripheral blood specimen from the neonate OR
- Demonstration of rising T. gondii IgG titres in sequential sera from the neonate OR
- Detection of T. gondii nucleic acid (e.g., PCR) in amniotic fluid, placental tissue, fetal or neonatal tissue, blood or CSF OR

Definition

- Isolation of T. gondii from blood or body fluid of the neonate by mouse inoculation OR
- Microscopic demonstration of T. gondii in an appropriate neonatal clinical specimen.

Clinical illness: Fetal infection early in pregnancy may manifest as fetal death, chorioretinitis, brain damage with intracerebral calcifications, hydrocephaly, microcephaly, fever, jaundice, rash, hepatosplenomegaly or convulsions. Fetal infection later in pregnancy results in mild or subclinical disease with delayed manifestations (recurrent or chronic chorioretinitis, developmental delay, hearing loss or blindness).

Probable Case:

- > Clinical illness in a child with laboratory evidence of T. gondii infection born to a seropositive mother OR
- > Clinical illness in a neonate born to a female with reactivated toxoplasma infection (rare).

Alberta Health and Wellness. Public Health Notifiable Disease Management Guidelines: Congenital Toxoplasmosis. *Government of Alberta.* http://www.ncbi.nlm.nih.gov/pubmed/23761154. Published 2011.

Item 99

Administer a combination of pyrimethamine-sulfadiazine-folinic acid during the first year of life to neonates in whom a diagnosis of congenital toxoplasmosis is confirmed or probable.

Follow your institution guideline for dosage.

Start

Grade of recommendations: Textbook

Guerina NG, Lee J, Lynfield R. Congenital toxoplasmosis: Treatment, outcome, and prevention. *UpToDate.com.* http://www.uptodate.com/contents/congenital-toxoplasmosis-treatment-outcome-and-prevention. Published 2015. Accessed June 9, 2016.

Rudin C, Boubaker K, Raeber PA, et al. Toxoplasmosis during pregnancy and infancy a new approach for Switzerland Swiss Working Group on congenital Toxoplasmosis. *Swiss Med Wkly.* 2008;138(49-50 SUPPL. 168):1-8.

Département de l'enfant et de l'adolescent - HUG. Cahier de l'interne. Hôpitaux Univ Genève. 2015;(Décembre).

All Infectiology/Toxoplasmosis

Item 100

Do not use spiramycin in neonates. Stop treatment and screen for potential QT interval prolongation.

Stop

Spiramycin is no longer used in suspected toxoplasmosis. It was indicated in the past while waiting for a conclusive diagnosis, but no benefit has ever been demonstrated and it can cause cardiac toxicity (QT prolongation).

Grade of recommendations: Review

Tomasoni LR, Meroni V, Bonfanti C, et al. Multidisciplinary approach to congenital Toxoplasma infection: An Italian nationwide survey. *New Microbiol.* 2014;37(3):347-354.

All Infectiology/Toxoplasmosis
Spiramycin





Cytomegalovirus (CMV)

Life-threatening disease:

- Viral sepsis-like syndrome
- Pneumonitis
- Myocarditis
- Severe hepatitis
- Enterocolitis

- Definition Severe and refractory thrombocytopenia
 - Sight-threatening retinitis
 - Severe neurologic disease
 - Underlying primary immune disorder (eg, severe combined immunodeficiency [SCID]) regardless of degree of symptoms

Severe focal disease is defined as severe hepatitis, severe bone marrow suppression, colitis or pneumonia.

Demmler-Harrison GJ. Congenital cytomegalovirus infection: Management and outcome. UpToDate.com. http://www.uptodate.com/contents/congenital-cytomegalovirus-infection-management and-outcome. Published 2015. Accessed June 13, 2016.

D'Oronzio U, Arlettaz MR, Hagmann C, Swiss Society of Neonatology. Congenital cytomegalovirus infection. Swiss Soc Neonatol. 2015;(May):1-22.

Item 101

Start antiviral treatment as soon as virologic testing is confirmed and within the first 30 days of life in symptomatic cytomegalovirus (CMV) infected newborns with central nervous system involvement or if life-threatening.

Treatment can be considered in symptomatic newborns with severe focal disease. IV ganciclovir and oral valganciclovir can be used, depending on the severity of the disease. Follow your institution guideline for dosage. Monitor full blood count, liver function tests, creatinine, urea and electrolytes. Suspend treatment if absolute neutrophil count < 500/µL or platelet count < 25 000/µL. Asymptomatically infected or mild/moderate symptomatic neonates should not be treated with antiviral agents.

Start / Stop

Grade of recommendations: National Guidelines

D'Oronzio U, Arlettaz MR, Hagmann C, Swiss Society of Neonatology. Congenital cytomegalovirus infection. Swiss Soc Neonatol. 2015;(May):1-22.

Swanson EC, Schleiss MR. Congenital Cytomegalovirus Infection : New Prospects for Prevention and Therapy. Pediatr Clin North Am. 2013;60(2):1-17. doi:10.1016/j.pcl.2012.12.008.

Buonsenso D, Serranti D, Gargiullo L, et al. Congenital cytomegalovirus infection: Current strategies and future perspectives. Eur Rev Med Pharmacol Sci. 2012;16(7):919-935.

Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.) . Elk Grove Village, IL: American Academy of Pediatrics; 2012.

All Infectiology/CMV





Item 102

	Stop antiviral treatment in neonates with asymptomatic cytomegalovirus infection
Stop	Antiviral therapy is not recommended routinely in neonates and young infants because of possible toxicities, including neutropenia in a significant proportion of recipients.
	Grade of recommendations: International Guidelines
	Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.
	Buonsenso D, Serranti D, Gargiullo L, et al. Congenital cytomegalovirus infection: Current strategies and future perspectives. <i>Eur Rev Med Pharmacol Sci.</i> 2012;16(7):919-935.
	Coll O, Benoist G, Ville Y, et al. Guidelines on CMV congenital infection. <i>J Perinat Med.</i> 2009;37(5):433-445. doi:10.1515/JPM.2009.127.
All	Infectiology/CMV
	Antiviral agents





Herpes Simplex Virus (HSV)

Clinical suspicion of neonatal herpes infection:

Since most neonatal heroes infections occur where the mother has no history of genital herpes, an HSV infection must be suspected immediately if the neonate exhibits suspicious symptoms (B).

Definition The possibility of neonatal herpes infection must be especially considered in case of:

- characteristic skin or mucosal lesions
- conjunctivitis, particularly if there is injection of the conjunctiva, bulbi, or keratitis
- seizures and/or lethargy without any other explanation
- fever or other systemic symptoms without any other explanation.

Swiss Herpes Management Forum. Swiss recommendations for the management of genital herpes and herpes simplex virus infection in neonate. Swiss Med Wkly. 2004;134:205-2014.

Item 103

Start aciclovir IV in neonates with Herpes Simplex Virus (HSV) disease, regardless of maternal history or pending laboratory confirmation or exclusion of HSV.

Localised HSV infections (skin, eyes and mouth) are treated for 14 days with parenteral aciclovir and CNS forms or disseminated infection for 21 day (C). Follow your institution guideline for dosage. For infants with CNS disease, CSF should be sampled near the end of a 21-day course of therapy. If the PCR remains positive, treatment should be extended with weekly CSF sampling and aciclovir stopped when a negative result is obtained (D). After acute HSV treatment, suppressive therapy with oral aciclovir should be given for six months to infants with CNS disease (D). Do not treat acute HSV infection with oral aciclovir because this leads to non-therapeutic drug levels (D).

Start

Grade of recommendations: National Guidelines

Swiss Herpes Management Forum. Swiss recommendations for the management of genital herpes and herpes simplex virus infection in neonate. Swiss Med Wkly. 2004;134:205-2014.

Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006.

Allen UD, Robinson JL, Canadian Paediatric Society. Prevention and management of neonatal herpes simplex virus infections. Paediatr Child Heal. 2014;19(4):201-206.

Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.) . Elk Grove Village, IL: American Academy of Pediatrics; 2012.

Infectiology/HSV ΑII





Item 104

	Start a topical antiviral treatment in combination with aciclovir IV in neonates with Herpes Simplex Virus (HSV) disease with ocular involvment.
	An ambiduard asiant consultation about the advised
	An ophtalmological consultation should be advised.
Start	Grade of recommendations: National Guidelines
	Swiss Herpes Management Forum. Swiss recommendations for the management of genital herpes and herpes simplex virus infection in neonate. <i>Swiss Med Wkly.</i> 2004;134:205-2014.
	Allen UD, Robinson JL, Canadian Paediatric Society. Prevention and management of neonatal herpes simplex virus infections. <i>Paediatr Child Heal.</i> 2014;19(4):201-206.
	Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.
All	Infectiology/HSV





Varicella-Zoster Virus (VZV) Item 105 Start Varicella-Zoster Immune Globulin (VZIG) 125IU IM as soon as exposure is known and within a 72 hour period, independent of maternal history of varicella, in neonates born at <28 weeks of gestational age or who weighed <1000g at birth who have been significantly exposed to Varicella-Zoster Virus (VZV). Start **Grade of recommendations: National Guidelines** Swiss Herpes Management Forum, Meylan P, Kempf W, et al. Recommandations suisses pour la prise en charge des infections dues au virus de la varicelle-zoster. Forum Med Suisse. 2007;7:895-Centers for Disease Control and Prevention. Updated Recommendations for Use of VariZIG — United States, 2013. Morbidity and Mortality Weeekly Report (MMWR). https://www.cdc.gov/mmwr/preview/mmwrhtml/mm6228a4.htm. Published 2013. Accessed June 29, Cobelli-Kett J. Perinatal varicella. Pediatr Rev. 2013;34(1). doi:10.1542/pir.34-1-49. <28w GA or Infectiology/VZV <1000g BW Item 106 Start Varicella-Zoster Immune Globulin (VZIG) 125IU IM in neonates ≥28w GA or ≥1000g BW who have been significantly exposed postnatally to Varicella-Zoster Virus (VZV), only if born to mother who has no or unknown history of varicella. VZIG must be administered as soon as exposure is known and within a 72 hour period. **Grade of recommendations: National Guidelines** Start / stop Swiss Herpes Management Forum, Meylan P, Kempf W, et al. Recommandations suisses pour la prise en charge des infections dues au virus de la varicelle-zoster. Forum Med Suisse. 2007;7:895-Centers for Disease Control and Prevention. Updated Recommendations for Use of VariZIG — United States, 2013. Morbidity and Mortality Weeekly Report (MMWR). https://www.cdc.gov/mmwr/preview/mmwrhtml/mm6228a4.htm. Published 2013. Accessed June 29,

Cobelli-Kett J. Perinatal varicella. Pediatr Rev. 2013;34(1). doi:10.1542/pir.34-1-49.

≥28w GA or ≥1000g BW 2016.

Infectiology/VZV





Item 107

Start Varicella-Zoster Immune Globulin (VZIG) 125IU IM as soon as possible, after birth or with onset of maternal illness, in term or late preterm neonates whose mother had varicella disease <u>5 days prior</u> to or 2 days after delivery.

Grade of recommendations: Institutional Guidelines

Start

Wilson L, Bowers L. Chicken Pox (Varicella Zoster Virus) VZ Immunoglobulin (VZIG) Information. *Newborn Services Clinical Guideline - Auckland District Health Board.* http://www.adhb.govt.nz/newborn/Guidelines/Infection/Varicella/VZIGInformation.htm. Published 2009. Accessed June 29, 2016.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

Cobelli-Kett J. Perinatal varicella. Pediatr Rev. 2013;34(1). doi:10.1542/pir.34-1-49.

All Infectiology/VZV

Item 108

Start aciclovir IV in neonates who develop systemic symptoms or severe cutaneous Varicella-Zoster disease, or who are at high risk of infection.

Infants at high risk of infection are those who did not receive Varicella-Zoster immunoglobulin (VZIG) as indicated, and/or are immunocompromised, and/or are <28 weeks' GA at birth. Do not give oral aciclovir as absorption is unpredictable in neonates.

Start

Grade of recommendations: National Guidelines

Swiss Herpes Management Forum, Meylan P, Kempf W, et al. Recommandations suisses pour la prise en charge des infections dues au virus de la varicelle-zoster. *Forum Med Suisse*. 2007;7:895-005

Royal Berkshire, Boden J. Varicella Infection in the Neonate GL366. *NHS Found Trust.* 2009;(October).

Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (*American Academy of Pediatrics, ed.*). Elk Grove Village, IL: American Academy of Pediatrics; 2012.

All Infectiology/VZV





Item 109

Stop	Stop Varicella-Zoster immunoglobulin (VZIG) if neonatal chickenpox has developed.
	Varicella-Zoster immunoglobulin (VZIG) is of no benefit once neonatal chickenpox has developed and treatment should be discontinued.
	Grade of recommendations: Institutional Guidelines
	The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.
	Infectiology/VZV
	Varicella-Zoster immunoglobulin (VZIG)





Chlamydia	
Item 110	
Stop	Do not use prophylactic antibiotic treatment in neonates at high risk of chlamydial infection (born to mothers who have untreated chlamydia).
	Neonates born to mothers who have untreated chlamydia are at high risk of infection; however, prophylactic antibiotic treatment is not indicated, as the efficacy of such treatment is unknown. Infants should be monitored to ensure appropriate treatment if symptoms develop.
	Grade of recommendations: National Guidelines
	Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006.
All	Infectiology/Chlamydia OR Prevention/Chlamydia Antibiotics
Item 111	
Start	Start erythromycin orally for 14 days in neonates with chlamydial conjunctivitis or with suspected or confirmed chlamydial pneumonia.
	Follow your institution guideline for dosage. Because the efficacy of erythromycin therapy is approximately 80%, a second course may be required, and a follow-up of infants is recommended. Neonates treated with erythromycin or azithromycin should be observed for signs and symptoms of hypertrophic pyloric stenosis.
	Grade of recommendations: National Guidelines
	Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006. Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.) . Elk Grove Village, IL: American Academy of Pediatrics; 2012.
All	Infectiology/Chlamydia





Item 112

Start azithromycin as second line treatment when erythromycin is not avaliable in neonates with chlamydial conjunctivitis or with suspected or confirmed chlamydial pneumonia. Azithromycin suspension should be administered for 3 days. Neonates treated with erythromycin or azithromycin should be observed for signs and symptoms of hypertrophic pyloric stenosis. Start Grade of recommendations: National Guidelines Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006. Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.) . Elk Grove Village, IL: American Academy of Pediatrics; 2012. Infectiology/Chlamydia

ΑII

Item 113	
	Stop topical antibiotics for the treatment of chlamydial conjunctivitis in neonates.
	Topical antibiotic therapy alone is inadequate for the treatment of ophthalmia neonatorum caused by chlamydia and is unnecessary when systemic treatment is administered.
04	
Stop	Grade of recommendations: National Guidelines
	Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006. Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove
	Village, IL: American Academy of Pediatrics; 2012.
All	Infectiology/Chlamydia
	Topical antibiotics





Gonorrhea

Definition

Suspicion of Gonorrhea infection:

Gonococcal ophthalmia is strongly suspected when intracellular gram-negative diplococci are identified on Gram stain of conjunctival exudate.

Frieden, T. R., Jaffe, H. W., Cono, J., Richards, C. L., & lademarco, M. F. (2015). Sexually transmitted diseases treatment guidelines, 2015. MMWR Recommendations and Reports, 64(RR-3), 1–140. http://doi.org/10.1097/00008480-200308000-00006

Item 114

Administer 1 dose of ceftriaxone IV or IM in all neonates born to mothers who have untreated gonorrhea.

Follow your institution guideline for dosage. Treatment must be preceded by testing the infant for gonorrhea at exposed sites. Avoid giving ceftriaxone to premature infants till 41 weeks postmenstrual age, hyperbilirubinemic infants and those receiving calciumcontaining intravenous fluids. When ceftriaxone must not be used, cefotaxime can be considered.

Start

Grade of recommendations: National Guidelines

Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. *MMWR Recomm Reports.* 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006.

Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.

ΑII

Infectiology/Gonorrhea OR Prevention/Gonorrhea

Item 115

Administer 1 dose of ceftriaxone IV or IM in neonates with suspected or confirmed gonococcal ophtalmia neonatorum or other localized gonococcal infection.

Start

Follow your institution guideline for dosage. Infant should receive eye irrigations with saline solution immediately and at frequent intervals until discharge is eliminated. Avoid giving ceftriaxone to premature infants till 41 weeks total age, hyperbilirubinemic infants and those receiving calcium-containing intravenous fluids. When ceftriaxone must not be used, cefotaxime can be considered.

Grade of recommendations: Textbook

Speer ME. Gonococcal infection in the newborn. *UpToDate.com.* http://www.uptodate.com/contents/gonococcal-infection-in-the-newborn#H19. Published 2015. Accessed May 31, 2016.

All Infectiology/Gonorrhea





Item 116

item 116	
	Stop topical antibiotics in neonates with suspected or confirmed gonococcal ophtalmia neonatorum.
Stop	They should receive IV antimicrobial therapy and eye irrigations with saline solution immediately and at frequent intervals until discharge is eliminated. Topical antimicrobial treatment alone is inadequate and unnecessary when recommended systemic antimicrobial treatment is given. This statement does not apply to gonococcal ophtalmia neonatorum prophylaxis, which is practiced in certain medical centers by giving topical treatment at birth to all newborn infants.
	Grade of recommendations: National Guidelines
	Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the
	Committee on Infectious Diseases, American Academy of Pediatrics, Redbook, Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.
All	Infectiology/Gonorrhea
Item 117	
	Start ceftriaxone IV or IM in neonates with disseminated gonococcal infection.
	Follow your institution guideline for dosage. Avoid giving ceftriaxone to premature infants till 41 weeks total age, hyperbilirubinemic infants and those receiving calcium-containing intravenous fluids. When ceftriaxone must not be used, cefotaxime can be considered.
Start	
	Grade of recommendations: National Guidelines

Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.) . Elk Grove Village, IL: American Academy of Pediatrics; 2012.

All

Infectiology/Gonorrhea

Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. *MMWR Recomm Reports.* 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006. **Committee on Infectious Diseases, American Academy of Pediatrics.** Redbook: Report of the





Item 118 Start vancomycin IV until bacteremia is excluded for localized Methicillinresistant Staphylococcus aureus (MRSA) disease in preterm or very lowbirthweight neonates or in more-extensive forms of the disease involving multiple sites in full-term neonates. Start Grade of recommendations: National Guidelines Liu C, Bayer A, Cosgrove SE, et al. Clinical practice guidelines by the Infectious Diseases Society of America for the treatment of methicillin-resistant Staphylococcus aureus infections in adults and children. Clin Infect Dis. 2011;52(3). doi:10.1093/cid/ciq146. Infectiology/MRSA infections





Syphilis

Confirmed congenital infection:

Treponema pallidum demonstrated by darkfield examination (DFE) or polymerase chain reaction (PCR) in placenta or autopsy material, exudate from suspicious lesions or body fluids, e.g. nasal discharge.

Presumed congenital infection:

- Children with a positive treponemal test for syphilis in combination with one or several of the following:
- persistent rhinitis, condylomata lata, osteitis, periostitis, osteochondritis, ascites, cutaneous and mucous membrane lesions, hepatitis, hepatosplenomegaly, glomerulonephritis, haemolytic anaemia;
- radiological abnormalities of the long bones suggestive of congenital syphilis;

Definition

- a positive Rapid Plasma Reagin (RPR) test / Venereal Diseases Research Laboratory (VDRL) test in the cerebrospinal fluid:
- a fourfold increase or more of the T. pallidum Passive Particle Agglutination (TPPA) / T. pallidum Haemagglutination (TPHA) titre in the child's as opposed to the mother's serum (both obtained simultaneously at birth);
- a fourfold increase or more of the titre of a non-treponemal test in the child's as opposed to the mother's serum (both obtained simultaneously at birth);
- a fourfold increase or more of the titre of a non-treponemal test within 3 months after birth;
- a positive anti-treponemal IgM EIA,19S-IgM-FTA-abs test and/or IgM- immunoblot for T. pallidum in the child's serum;
- a mother, in whom syphilis was confirmed during pregnancy, but who was not adequately treated either before or during pregnancy.

Carlin E, Ziza J, Keat A, Janier M. 2014 European Guideline on the management of sexually acquired reactive arthritis. *Int J STD AIDS*. 2014;25(13):901-912. doi:10.1177/0956462414540617.

Item 119

Administer benzylpenicillin G IV, OR procaine penicillin to neonates with confirmed or presumed congenital syphilis, or born to syphilis infected mothers who have not been treated with penicillin at least four weeks prior delivery.

Follow your institution guideline for dosage.

Start

Grade of recommendations: National Guidelines

Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. *MMWR Recomm Reports.* 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006.

Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.

All Infectiology/Syphilis





Item 120

Start	Administer 1 dose of benzathine penicillin G IM in neonates with normal examination, born to syphilis infected mothers who have been adequately treated during pregnancy more than 4 weeks prior to delivery.
	Follow your institution guideline for dosage. If mother's nontreponemal serologic titer remained low and stable before and during pregnancy and at delivery (Venereal Disease Research Laboratory (VDRL) slide test <1:2; rapid plasma reagin (RPR) test <1:4), no treatment is required. If follow-up is uncertain, a single dose of benzathin penicillin G IM can be considered.
	Grade of recommendations: National Guidelines
	Frieden TR, Jaffe HW, Cono J, et al. Sexually transmitted diseases treatment guidelines, 2015. MMWR Recomm Reports. 2015;64(RR-3):1-140. doi:10.1097/00008480-200308000-00006.
	Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.
All	Infectiology/Syphilis





Ureaplasma Urealyticum Infection

Item 121

Reassess the use of macrolides or other antibiotics for the treatment of Ureaplasma urealyticum in neonates.

Stop

Despite in vitro susceptibility of Ureaplasma spp. to erythromycin and favorable pharmacokinetic activity, trials of erythromycin therapy in ureaplasma-colonized preterm infants have failed to demonstrated efficacy to prevent BPD or to eradicate respiratory tract colonization (C). The efficacy of azithromycin and related macrolide, clarithromycin, to prevent BPD has been assessed in single center studies of at-risk preterm infants, but the safety and optimal dosing regimens for these antibiotics have not been determined in appropriate pharmacokinetic and pharmacodynamic studies. It is currently unknown whether eradicating Ureaplasma spp. from the preterm respiratory tract with appropriate antibiotic therapy will prevent ureaplasma infection-mediated lung injury (FRN).

Grade of recommendations: Review

Viscardi RM. Ureaplasma species: Role in Neonatal Morbidities and Outcomes. *Arch Dis Child Fetal Neonatal Ed.* 2014;99(1):1-14. doi:10.1136/archdischild-2012-303351.

All

Infectiology/Ureaplasma Urealyticum Infection

Macrolides





Urinary Tract Infection (UTI)

Urinary tract infection is defined here as:

1. Pyuria as determined with an "enhanced urinalysis" (hemocytometer counting chamber) of ≥10 WBC/µL;

2. A urine culture colony count of ≥10 000 CFU/mL for a single organism.

Vesico-ureteric reflux (VUR):

Definition

Grade I: urine refluxes into the ureter only;

Grade II: urine refluxes into the ureter and up to the kidney without dilation;

Grade III: urine refluxes into the ureter and kidney and causes mild dilation;

Grade IV: urine refluxes into ureter and kidney and causes dilation without twisting of the ureter:

Grade V: urine refluxes into ureter and kidney and causes significant dilation with twisting of the ureter.

Santoro JD, Carroll VG, Steele RW. Diagnosis and Management of Urinary Tract Infections in Neonates and Young Infants. *Clin Pediatr (Phila)*. 2012;52(2):111-114. doi:10.1177/0009922812471713.

UCSF Department of Urology. Vesicoureteral Reflux (VUR). *Urology care foundation.* https://urology.ucsf.edu/patient-care/children/additional/vesicoureteral-reflux. Published 2016. Accessed June 30, 2016.

Item 122

Start empiric antibiotics after urine samples and cultures are collected in neonates with fever when urinary tract infection is suspected.

Follow your institution guidelines for dosage and choice of antibiotic agents.

Start

Grade of recommendations: International Guidelines

Bradley JS, Nelson JD. Nelson's Pediatric Antimicrobial Therapy. 19th Editi. (*American Academy of Pediatrics, ed.*). San Diego: American Academy of Pediatrics; 2012.

O'Donovan DJ. Urinary tract infections in neonates. UpToDate.com.

http://www.uptodate.com/contents/urinary-tract-infections-in-neonates#H776291741. Published 2015. Accessed June 30, 2016.

European Association of Urology, Stein R, Dogan HS, et al. Urinary tract infections in children: EAU/ESPU guidelines. *Eur Urol.* 2015;67:546-558. doi:10.1016/j.eururo.2014.11.007.

All Infectiology/UTI





Item 123

Consider antibiotic prophylaxis after an urinary tract infection (UTI) only in neonates with grade IV-V vesico-ureteric reflux.

Antibiotic prophylaxis in order to avoid recurrent UTI is no longer routinely recommended after a UTI but may still be considered when a child is known to have a grade IV or V VUR, or a significant uro- logical anomaly. Grade IV-V are defined as urine refluxes into ureter and kidney and causes dilatation with or without twisting of the ureter.

Start / stop

Grade of recommendations: International Guidelines

Canadian Paediatric Society, Robinson JL, Finlay JC, Lang ME, Bortolussi R. Prophylactic antibiotics for children with recurrent urinary tract infections. Paediatr Child Heal. 2015;20(1):45-47. doi:10.1093/jac/25.4.505.

European Association of Urology, Stein R, Dogan HS, et al. Urinary tract infections in children: EAU/ESPU guidelines. Eur Urol. 2015;67:546-558. doi:10.1016/j.eururo.2014.11.007.

Bradley JS, Nelson JD. Nelson's Pediatric Antimicrobial Therapy. 19th Editi. (American Academy of Pediatrics, ed.). San Diego: American Academy of Pediatrics; 2012.

All Infectiology/UTI
Antibiotics





Pertussis

Suspicion of Pertussis infection:

Pertussis should be suspected (regardless of vaccination status or wheezing) in the following patients (see 'Clinical suspicion' above):

- •Infants <4 months with a cough illness, usually without significant fever, who have:
- -Cough that is not improving (of any duration); the cough may or may not be paroxysmal (movie 1)
- -Rhinorrhea in which the nasal discharge remains watery
- -Apnea, seizures, cyanosis, vomiting, or poor weight gain
- -Leukocytosis with lymphocytosis (WBC count ≥20,000 cells/microL with ≥50 percent lymphocytes)
- -Pneumonia

Yeh S, Mink CM. Bordetella pertussis infection in infants and children: Clinical features and diagnosis. *UpToDate.com.* http://www.uptodate.com/contents/bordetella-pertussis-infection-in-infants-and-children-clinical-features-and-diagnosis?source=see_link. Published 2016. Accessed August 30, 2016.

Item 124

Definition

Start azithromycin oral daily for 5 days in neonates with suspected or confirmed pertussis infection, or in those in close contact with confirmed and contagious cases of pertussis.

Follow your institution guideline for dosage. A person is contagious when < 21 days of cough and < 5 days effective antibiotics. Neonates on macrolide should be monitored for infantile hypertrophic pyloric stenosis (IHPS).

Start

Grade of recommendations: National Guidelines

The Royal Children's Hospital Melbourne. Whooping Cough (Pertussis). The *Royal Children's Hospital Melbourne.*

http://www.rch.org.au/clinicalguide/guideline_index/Whooping_Cough_Pertussis/. Published 2016. Accessed June 30, 2016.

Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (*American Academy of Pediatrics, ed.*). Elk Grove Village, IL: American Academy of Pediatrics; 2012.

Amirthalingam G, Pertussis Guidelines Group. HPA guidelines for the public health management of pertussis. *Heal Prot Agency*. 2012;(October):1-45.

http://www.hpa.org.uk/webc/HPAwebFile/HPAweb_C/1287142671506

All Infectiology/Pertussis





Tuberculosis Item 125 Start isoniazid prophylaxis orally in neonates born to mothers with tuberculosis (TB), or those in close contact with people with smear-positive pulmonary or laryngeal TB who have not had at least 2 weeks of antituberculosis treatment. Prophylaxis for babies born to mothers with active TB is 9 months, and 2-3 months in those born to mothers with inactive TB, if the infant remains asymptomatic. Infants exposed postnatally should be treated 2-3 months after last exposure, if the infant remains asymptomatic. Follow your institution guideline for dosage. Start **Grade of recommendations: National Guidelines** Centre for Communicable Diseases and Infection Control. Pediatric Tuberculosis. In: Public Health Agency of Canada, ed. Canadian Tuberculosis Standards. 7th Editio. Public Health Agency of Canada; 2007. http://www.lung.ca/cts-sct/pdf/tbstand07_e.pdf. National Institute For Health and Care Excellence. Tuberculosis. NICE Clin Guidel. 2016;33. nice.org.uk/guidance/ng33. Bradley JS, Nelson JD. Nelson's Pediatric Antimicrobial Therapy. 19th Editi. (American Academy of Pediatrics, ed.). San Diego: American Academy of Pediatrics; 2012. ΑII Infectiology/Tuberculosis OR Prevention/Tuberculosis Item 126 Start anti-tuberculosis treatment in neonates with congenital tuberculosis or postnatal tuberculosis primary pulmonary disease. Follow your institution guideline for dosage. **Grade of recommendations: Textbook** Start Bradley JS, Nelson JD. Nelson's Pediatric Antimicrobial Therapy. 19th Editi. (American Academy of Pediatrics, ed.). San Diego: American Academy of Pediatrics; 2012. Mittal H, Das S, Faridi MMA. Management of newborn infant born to mother suffering from tuberculosis: Current recommendations & gaps in knowledge. Indian J Med Res. 2014;140:32-39.



ΑII



Infectiology/Tuberculosis

10. ENDOCRINOLOGY <u>Metabolic Bone Disorder (MBD)</u>

Definition

Metabolic Bone Disorder

Decreased bone mineral content relative to the expected level of mineralization for a fetus or infant of comparable size or gestational age seen in conjunction with biochemical and/or radiographic changes.

Rustico SE, Calabria AC, Garber SJ. Metabolic bone disease of prematurity. *J Clin Transl Endocrinol*. 2014;1:85-91. doi:10.1016/j.jcte.2014.06.004.

Risk factors for Metabolic Bone Disorder

- <32 weeks' gestation or <1500 g birth-weight
- Male gender
- Inadequate nutrition
 - Suboptimal intake
- Enteral feeds with low mineral content/bioavailability (unfortified expressed breast milk, term formula given to a preterm infant)

- Phosphorus deficiency (primary nutritional reason)

- Definition
- Vitamin D deficiencyProlonged total parenteral nutrition
- Chronic use of drugs that increase mineral excretion (diuretics, dexamethasone, sodium bicarbonate)
- Lack of mechanical stimulation e.g. sedation/paralysis
- Bronchopulmonary dysplasia
- Cholestatic jaundice
- Short gut syndrome (malabsorption of vitamin D and Ca)

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016

Item 127

Administer calcium, phosphate and vitamin D in preterm infants <32 weeks of gestational age or <1500g or infants at risk of metabolic bone disorders.

When receiving enteral feed, neonates should be given a fortifier (in breastfed infants) or preterm formula with calcium (3.5-5.5 mmol/kg/day) and phosphate (2.5-4.5 mmol/kg/day). If parenteral nutrition is needed, use parenteral nutrition with optimised calcium and phosphate content (Ca 1.8 mmol/kg/day and PO4 1.4 mmol/kg/day) and vitamin D 160-400 IU/kg/day. Maximal mineral accretion rates have been reported with Ca/PO4 (mol/mol) ratio 1.3.

Start

In neonates with biochemical features found in metabolic bone disease, aim for the upper end of the recommended range of calcium and phosphate intake to prevent fractures.

Grade of recommendations: Review

Royal Prince Alfred Hospital Care Newborn. Metabolic bone disease. *Sydney Local Health District - New South Wales government.*

http://www.slhd.nsw.gov.au/rpa/neonatal\content/pdf/guidelines/metabolicBD.pdf. Published 2016. Accessed June 24, 2016.

Rustico SE, Calabria AC, Garber SJ. Metabolic bone disease of prematurity. *J Clin Transl Endocrinol*. 2014;1:85-91. doi:10.1016/j.jcte.2014.06.004.

Pereira-da-Silva L, Costa A, Pereira L, et al. Early high calcium and phosphorus intake by parenteral nutrition prevents short-term bone strength decline in preterm infants. *J Pediatr Gastroenterol Nutr.* 2011;52(2):203-209. doi:10.1097/MPG.0b013e3181f8b295.

ΑII

Prevention/Prevention of MBD OR Endocrinology/MBD





Item 128

Start

Administer the maximal recommended doses of calcium, phosphate and vitamin D to prevent fractures in neonates with biochemical features of metabolic bone disease.

Recommended range are:

When receiving enteral feed, neonate should be given fortifier (for breastfed infant) or preterm formula with calcium (3.5-5.5 mmol/kg/day) and phosphate (2.5-4.5 mmol/kg/day). Do not give Ca and PO4 at the same time because they may precipitate; so give at alternate feeds.

If parenteral nutrition is needed, use parenteral nutrition with optimised calcium and phosphate content (Ca 1.8 mmol/kg/day and PO4 1.4 mmol/kg/day) and vitamin D 160-400 IU/kg/day. Maximal mineral accretion rates have been reported with Ca/PO4 (mol/mol) ratio 1.3.

Grade of recommendations: Institutional Guidelines

Royal Prince Alfred Hospital Care Newborn. Metabolic bone disease. *Sydney Local Health District New South Wales government.*

http://www.slhd.nsw.gov.au/rpa/neonatal\content/pdf/guidelines/metabolicBD.pdf. Published 2016. Accessed June 24, 2016.

Rustico SE, Calabria AC, Garber SJ. Metabolic bone disease of prematurity. *J Clin Transl Endocrinol*. 2014;1:85-91. doi:10.1016/j.jcte.2014.06.004.

Pereira-da-Silva L, Costa A, Pereira L, et al. Early high calcium and phosphorus intake by parenteral nutrition prevents short-term bone strength decline in preterm infants. *J Pediatr Gastroenterol Nutr.* 2011;52(2):203-209. doi:10.1097/MPG.0b013e3181f8b295.

All Endocrinology/MBD

Item 129

Stop steroids and furosemide as soon as possible in neonates at risk of metabolic bone disorder.

In the primary prevention and treatment strategy for MBD, limiting the prolonged exposure to commonly prescribed medications that further reduce mineral stores (e.g. loop diuretics and methylxanthines) or increase bone resorption (e.g. glucocorticoids) is equally important to optimizing nutrition.

Stop

Grade of recommendations: National Guidelines

Rustico SE, Calabria AC, Garber SJ. Metabolic bone disease of prematurity. *J Clin Transl Endocrinol.* 2014;1:85-91. doi:10.1016/j.jcte.2014.06.004.

Abrams SA, Committee on Nutrition, American Academy of Pediatrics. Calcium and Vitamin D Requirements of Enterally Fed Preterm Infants. *Pediatrics*. 2013;131(5):e1676-83. doi:10.1542/peds.2013-0420.

All Prevention/Prevention of MBD OR Endocrinology/MBD





Thyroid Disorders (OR Hypothyroidism)

Item 130

Start levothyroxine (L-T4) immediately in neonates with thyroid function test (TFT) that results in either a free T4 (FT4) concentration below norms for age or a venous TSH concentration > 20 mIU/L.

TFT is normaly performed when capillary TSH concentration from blood obtained on neonatal screening was elevated; therefore this recommendation assumes a high capillary TSH value. Imaging should never be allowed to delay the initiation of treatment. Recommended treatment is levothyroxine (L-T4), given at an initial oral dose of 10-15 µg/kg/day. Infants with very low total T4 or free T4 should be treated with the highest initial dose. Any change in source of the L-T4 (brand) or in formulation (liquid vs tablets) requires retitration of the dose. If intravenous treatment is necessary the dose should be no more than 80% of the oral dose. The dose should then be adjusted according to TSH and FT4 levels.

Start

Grade of recommendations: International Guidelines

Léger J, Olivieri A, Donaldson M, et al. European Society for Paediatric Endocrinology Consensus Guidelines on Screening, Diagnosis, and Management of Congenital Hypothyroidism. *J Clin Endocrinol Metab.* 2014;99(2):363–384. doi:10.1159/000358198.

Rose SR, Brown RS, American Academy of Pediatrics, American Thyroid Association, Lawson Wilkins Pediatric Endocrine Society. Update of Newborn Screening and Therapy for Congenital Hypothyroidism. *Pediatrics*. 2006;117(6):2290-2303. doi:10.1542/peds.2006-0915.

ΑII

Endocrinology/Congenital hypothyroidism OR Endocrinology/Thyroid disorders





Hyperglycemia

Hyperglycemia

Definition

There is no established definition of hyperglycemia. However, start management if:

- two blood sugars are ≥14 on 2 occasions measured at least 2 hr apart
- blood sugars are ≥12 on 2 occasions measured at least 2 hr apart with evidence of significant glycosuria (positive on the urine dipstick).

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, & Southern West Midlands Maternity and Newborn Network. (2015). Neonatal Guidelines 2015-17. Retrieved January 1, 2016, from https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines

Item 131

Decrease glucose intake if necessary and decrease or stop drugs that worsen hyperglycemia, in neonates with hyperglycemia.

If glucose delivery rate >10 mg/kg/min, decrease glucose in increments to 6–10 mg/kg/min. If on TPN, 8–10 mg/kg/min is acceptable. Medications that can worsen hyperglycemia include corticosteroids and phenytoin.

Stop

Grade of recommendations: Textbook

Stark AR, Simmons R. Neonatal hyperglycemia. *UpToDate.com.* http://www.uptodate.com/contents/neonatal-hyperglycemia? Published 2015. Accessed July 5, 2016.

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

All Endocrinology/Hyperglycemia







Item 132

Start insulin only in patients with persistent hyperglycemia when other methods of glucose control have failed.

Other methods of glucose control include decrease of glucose infusion rates, stop of medications predisposing patients to hyperglycemia, and correction of underlying causes of hyperglycemia (i.e., sepsis). Starting dose of insulin is usually 0.05 units/kg/hr, then adjusted according to requirements. Do not include insulin in the total daily fluid intake - it should be titrated on top of the prescribed fluid intake. Monitor the blood glucose concentration, initially once every 2 hours, and once stable at least once every 8 hours. Aim for a blood glucose concentration between 6 and 10 mmol/L.

To prevent hypoglycaemia in neonate on insulin:

- 6-8 mmol/L and stable -> maintain insulin infusion
- 6-8 mmol/L with a moderate decrease: reduce insulin infusion rate to 50% of present rate
- <6 mmol/L or 6-8 mmol/L with a rapid decrease: stop infusion

Start / stop

Grade of recommendations: National Guidelines

The Bedside Clinical Guidelines Partnership, Staffordshire Shropshire & Black Country Newborn and Maternity Network, Southern West Midlands Maternity and Newborn Network. Neonatal Guidelines 2015-17. NHS network. https://www.networks.nhs.uk/nhs-networks/staffordshire-shropshire-and-black-country-newborn/neonatal-guidelines. Published 2015. Accessed January 1, 2016.

Stark AR, Simmons R. Neonatal hyperglycemia. *UpToDate.com.*

http://www.uptodate.com/contents/neonatal-hyperglycemia? Published 2015. Accessed July 5, 2016.

Fallon EM, Nehra D, Potemkin AK, et al. A.S.P.E.N. Clinical Guidelines: Hyperglycemia and Hypoglycemia in the Neonate Receiving Parenteral Nutrition. *J Parenter Enter Nutr.* 2012;36(1):81-95. doi:10.1177/0148607111418980.

Alsweiler J. Hyperglycaemia Causes of Hyperglycaemia Complications of Hyperglycaemia Management of Hyperglycaemia Management of Insulin Infusion. *Newborn Services Clinical Guideline - Auckland District Health Board.*

http://www.adhb.govt.nz/newborn/Guidelines/Nutrition/Hyperglycaemia.htm. Published 2013. Accessed July 4, 2016.

ΑII

Endocrinology/Hyperglycemia

Item 133

Do not provide high glucose infusion rates to prevent hypoglycemia in neonates receiving parenteral nutrition.

Stop

Excess glucose delivery should be avoided to maintain optimal blood glucose concentrations in neonates receiving parenteral nutrition as this may lead to hyperglycemia.

Grade of recommendations: National Guidelines

Fallon EM, Nehra D, Potemkin AK, et al. A.S.P.E.N. Clinical Guidelines: Hyperglycemia and Hypoglycemia in the Neonate Receiving Parenteral Nutrition. *J Parenter Enter Nutr.* 2012;36(1):81-95. doi:10.1177/0148607111418980.

All

Prevention/Prevention of hyperglycemia OR Endocrinology/Hyperglycemia Total Parenteral Nutrition (TPN)





Item 134

ILCIII 134	
	Do not use early insulin therapy in neonates at risk of hyperglycemia.
Stop	The use of early insulin therapy to prevent hyperglycemia is not recommended. There has been substantial research regarding the use of early, continuous insulin infusion to prevent hyperglycemia in the neonate. While a number of small studies suggest a benefit, other larger studies have raised significant concerns regarding this practice. Specifically, a large RCT was terminated early due to increased incidence of hypoglycemia and mortality in the early continuous insulin infusion group. A recent Cochrane review also determined that there is insufficient evidence to recommend early, continuous insulin infusion.
	Grade of recommendations: A
	Fallon EM, Nehra D, Potemkin AK, et al. A.S.P.E.N. Clinical Guidelines: Hyperglycemia and Hypoglycemia in the Neonate Receiving Parenteral Nutrition. <i>J Parenter Enter Nutr.</i> 2012;36(1):81-95. doi:10.1177/0148607111418980.
All	Prevention/Prevention of hyperglycemia OR Endocrinology/Hyperglycemia OR
All	Prevention/Prevention of hypoglycemia OR Endocrinology/Hypoglycemia Insulin
	III SUIII





Hypoglycemia

Symptoms of hypoglycemia in newborns include: Irritability, tremors, jitteriness, exaggerated Moro reflex, high-pitched cry, seizures, lethargy, floppiness, cyanosis, apnea, poor feeding.

Neonates at increased risk of hypoglycemia:

- 1. Symptoms of hypoglycemia
- 2. Large for gestational age (even without maternal diabetes)
- 3. Perinatal stress
 - a. Birth asphyxia/ischemia; cesarean delivery for fetal distress
 - b. Maternal preeclampsia/eclampsia or hypertension
 - c. Intrauterine growth restriction (small for gestational age)
 - d. Meconium aspiration syndrome, erythroblastosis fetalis, polycythemia, hypothermia
- 4. Premature or postmature delivery
- 5. Infant of diabetic mother
- 6. Family history of a genetic form of hypoglycemia
- 7. Congenital syndromes (eg, Beckwith-Wiedemann), abnormal physical features (eg, midline facial malformations, microphallus)

Persistent hypoglycemia

Hypoglycemia that persists or occurs for the first time beyond the first 3 days of life.

Adamkin DH. Postnatal Glucose Homeostasis in Late-Preterm and Term Infants Clinical Report—Postnatal Glucose Homeostasis in Late-Preterm and Term Infants. *Pediatrics*. 2013;127(3):575-579. doi:10.1542/peds.2010-3851.

Thornton PS, Stanley CA, De Leon DD, et al. Recommendations from the Pediatric Endocrine Society for Evaluation and Management of Persistent Hypoglycemia in Neonates, Infants, and Children. *J Pediatr.* 2015;167(2):238-245. doi:10.1016/j.jpeds.2015.03.057.

Item 135

Definition

Start IV glucose infusion in asymptomatic neonates with serum glucose level of <2.6 mmol/L if increased enteral caloric intake is not effective.

Start with an initial glucose infusion regime of 5.5 mg/kg/min. Infants with very low glucose levels, particularly those with levels less than 1.8 mmol/L, should be managed with some expedience, confirming response to intervention in a timely fashion (a response to intravenous interventions should occur within 30 min). Breastfeeding may be continued without risk of overhydration because the volume of colostrum is small. Blood glucose levels should be checked frequently until interventions result in stable glucose levels of 2.6 mmol/L or higher. In neonates with persistent hypoglycemia, consider investigation, consultation and pharmacological intervention if target blood glucose levels are not achieved by intravenous glucose.

Start

Grade of recommendations: Institutional Guidelines

Canadian Paediatric Society Fetus and Newborn Committee, Aziz K, Dancey P. Screening guidelines for newborns at risk for low blood glucose. *Paediatr Child Heal.* 2004;(March 2004, reaffirmed in 2016):1-7.

Queensland Clinical Guidelines Queensland Health. Maternity and Neonatal Clinical Guideline Induction of labour. *Queensl Gov Dep Heal.* 2015;(April). www.health.qld.gov.au/qcg.

Weston P, Harris D, Battin M, Brown J, Hegarty J, Harding J. Oral dextrose gel for the prevention of hypoglycaemia in newborn infants. *Cochrane Database Syst Rev.* 2016;(5). doi:10.1002/14651858.CD011027.pub2.

Thornton PS, Stanley CA, De Leon DD, et al. Recommendations from the Pediatric Endocrine Society for Evaluation and Management of Persistent Hypoglycemia in Neonates, Infants, and Children. *J Pediatr.* 2015;167(2):238-245. doi:10.1016/j.jpeds.2015.03.057.

ΑII

Endocrinology/Hypoglycemia





Item 136

Start IV glucose infusion immediately in symptomatic neonates with glucose levels <2.6 mmol/L.

Because severe, prolonged, symptomatic hypoglycemia may result in neuronal injury, prompt intervention is necessary for infants who manifest clinical signs and symptoms. Atrisk infants with glucose levels less than 1.8 mmol/L on one occasion (assuming one effective feed), or repeatedly less than 2.6 mmol/L, require intervention (C). There should be concurrent investigation and management of the underlying cause. Breastfeeding may be continued without risk of overhydration because the volume of colostrum is small. Blood glucose levels should be checked frequently until interventions result in stable glucose levels of 2.6 mmol/L or higher. In neonates with persistent hypoglycemia, consider investigation, consultation and pharmacological intervention if target blood glucose levels are not achieved by intravenous glucose.

Start

Grade of recommendations: National Guidelines

Canadian Paediatric Society Fetus and Newborn Committee, Aziz K, Dancey P. Screening guidelines for newborns at risk for low blood glucose. *Paediatr Child Heal.* 2004;(March 2004, reaffirmed in 2016):1-7.

Adamkin DH, American Academy of Pediatrics Committee on Fetus and Newborn. Postnatal Glucose Homeostasis in Late-Preterm and Term Infants Clinical Report—Postnatal Glucose Homeostasis in Late-Preterm and Term Infants. *Pediatrics.* 2013;127(3):575-579. doi:10.1542/peds.2010-3851.

Thornton PS, Stanley CA, De Leon DD, et al. Recommendations from the Pediatric Endocrine Society for Evaluation and Management of Persistent Hypoglycemia in Neonates, Infants, and Children. *J Pediatr.* 2015;167(2):238-245. doi:10.1016/j.jpeds.2015.03.057.

All Endocrinology/Hypoglycemia





11. PHARMACOLOGY **Drug & Breast-Feeding**

Item 137 Check the possible milk transfer of drugs taken by mothers to breastfed neonates, and monitor for potential adverse drug effects. Different references can be consulted or refere to a specialist (clinical pharmacist or pharmacologist). For exemple, several reported cases of codeine toxicity, in breastfed neonates (including death) have been published. Keep in mind that drugs given to the breastfeeding mother can also interact with the treatment of the neonate. Here are exemples of common sources of information about drugs and lactation: Other - Briggs GG, Freeman RK, Yaffe SJ. Drugs in Pregnancy and Lactation: A Reference Guide to Fetal and Neonatal Risk. 9th Editio. (Wolters Kluwer Health/Lippincott Williams & Wilkins, ed.). California; 2012. - Swiss teratogen information service: http://www.swisstis.ch/ - Centre de référence sur les agents tératogènes (in french): www.lecrat.org Grade of recommendations: D

Pharmacology/Drug & Breast-Feeding ΑII





Drug-drug Interactions

Item 138

Check changes in drug effect when initiating strong inhibitors or inducers of the cytochrome P450 and/or P-glycoprotein.

Although difficult to predict, modification of plasma concentration may occur when initiating strong inhibitors or inducers of the cytochrome P450 and/or P-glycoprotein. Exemples of strong inhibitors and inducers used in neonates are:

Inhibitors: erythromycine, fluconazole.

Inducers: phenobarbital, phenytoin, rifampicin.

Keep in mind that drugs given to the breastfeeding mother can also interact with the treatment of the neonate, if transfer to milk occurs.

Here are exemples of common sources of information about drug interactions:

- Lexicomp's Lexi-Interact: http://www.uptodate.com/crlsql/interact/frameset.jsp
- HUG pocket's card, Drug interaction, cytochrome and P-glycoprotein (validated for adult patient): http://www.hug-

Other

ge.ch/sites/interhug/files/structures/pharmacologie_et_toxicologie_cliniques/documents/interactions medicamenteuses_et_cyp450.pdf

- Theriaque: http://www.theriaque.org/

Grade of recommendations: D

de Wildt SN, Tibboel D, Leeder JS. Drug metabolism for the paediatrician. *Arch Dis Child.* 2014;99(12):1137-1142. doi:10.1136/archdischild-2013-305212.

Tayman C, Rayyan M, Allegaert K. Neonatal pharmacology: extensive interindividual variability despite limited size. *J Pediatr Pharmacol Ther.* 2011;16(3):170-184. doi:10.5863/1551-6776-16.3.170.

Centre d'information thérapeutique et de pharmacovigilance. Interactions Medicamenteuses, Cytochromes P450 et p-Glycoproteine (pgp). *Hôpitaux Universitaires de Genève.* http://www.hug-ge.ch/sites/interhug/files/structures/pharmacologie_et_toxicologie_cliniques/documents/interactions_medicamenteuses_et_cyp450.pdf. Published 2016. Accessed September 16, 2016.

All

Pharmacology/Drug-drug Interactions

Erythromycine, fluconazole, phenobarbital, phenytoin, rifampicin.





Item 139	Various
item 139	
	Do not use ceftriaxone in neonates who are being, or who have recently been given any IV fluids that contain calcium (such as TPN or Ringer Lactate)
	Precipitation may occur even when the two products are administered in different tubes and could be potentially lethal. Ceftriaxone can also induce kernicterus in the neonate by displacing bilirubin from plasmatic proteins, and its use should be avoided when possible.
Stop	Grade of recommendations: Manufacturer
σιορ	Grade of recommendations: Manufacturer
	Bradley JS, Wassel RT, Lee L, Nambiar S. Intravenous ceftriaxone and calcium in the neonate: assessing the risk for cardiopulmonary adverse events. <i>Pediatrics</i> . 2009;123(4):e609-13. doi:10.1542/peds.2008-3080.
	Ainsworth SBA. Neonatal Formulary. 7th ed. (John Wiley & Sons, ed.). Chichester, UK; 2015.
	Choonara I, Sammons H. Paediatric clinical pharmacology in the UK. <i>Arch Dis Child</i> . 2014;99(12):1143-1146. doi:10.1136/archdischild-2014-306853. Roche Laboratories Inc. Product Information: Rocephin IV, IM injection, powder for solution. <i>Swissmedicinfo.ch</i> . Swissmedic.ch. Published 2010. Accessed July 25, 2016.
All	Pharmacology/Various
	Calcium, ceftriaxone
Item 140	
	Do not use trimethoprime - sulfamethoxazole in neonates.
	Trimethannime culformathanarale is control indicated in necessary Culphanamides con
	Trimethoprime-sulfamethoxazole is contra-indicated in neonates. Sulphonamides can induce kernicterus in the neonate by displacing bilirubin from plasmatic proteins.
	made nonnecessor and necessor and appropriate from processor.
Stop	Grade of recommendations: National Guidelines
	Committee on Infectious Diseases, American Academy of Pediatrics. Redbook: Report of the Committee on Infectious Diseases. 29th Editi. (American Academy of Pediatrics, ed.). Elk Grove Village, IL: American Academy of Pediatrics; 2012.
	Choonara I, Sammons H. Paediatric clinical pharmacology in the UK. <i>Arch Dis Child.</i> 2014;99(12):1143-1146. doi:10.1136/archdischild-2014-306853.



All



Pharmacology/Various

 $Trime tho prime - sulfame tho xazole, \ Co-trimo xazole$

Item 141

Check excipients contained in prescribed drug formulations administered orally or parenterally since they can be harmful and responsible for adverse events in neonates, due to immature metabolism.

Benzyl alcohol must not be given to premature babies or neonates due to risks of serious adverse effects (gasping syndrome and deaths have been reported).

Propylenglycol must not be administrated to neonates at a dose higher than 1 mg/kg/day because it could lead to metabolic acidosis.

Excipients identified as known to be harmful to neonates should be avoided as much as possible. The sum of methyl-, ethyl- and propylparaben should not be >10mg/kg/day in oral drugs and propylparaben oral daily intake should not be >2mg/kg/day.

Other excipients known to be harmful in neonates are: polysorbate 80, sodium benzoate, benzalkonium chloride, saccharin sodium, sorbitol and ethanol. High dose and long term use should be avoided when possible.

Grade of recommendations: D

European Commission. Guidelines Medicinal products for human use Safety, environment and information. Excipients in the label and package leaflet of medicinal products for human use. *Pharm Regul Framew Mark Auth.* 2003.

Stop http://www.ema.europa.eu/docs/en_GB/document_library/Scientific_guideline/2009/09/WC50000341

Lass J, Naelapää K, Shah U, et al. Hospitalised neonates in Estonia commonly receive potentially harmful excipients. *BMC Pediatr.* 2012;12:136. doi:10.1186/1471-2431-12-136.

Shehab N, Lewis CL, Streetman DD, Donn SM. Exposure to the pharmaceutical excipients benzyl alcohol and propylene glycol among critically ill neonates. *Pediatr Crit Care Med.* 2009;10(2):256-259. doi:10.1097/PCC.0b013e31819a383c.

Souza A, Santos D, Fonseca S, et al. Toxic excipients in medications for neonates in Brazil. *Eur J Pediatr.* 2014;173(7):935-945. doi:10.1007/s00431-014-2272-z.

Nellis G, Metsvaht T, Varendi H, et al. Potentially harmful excipients in neonatal medicines: a pan-European observational study. *Arch Dis Child.* 2015;100(7):694-699. doi:10.1136/archdischild-2014-307793

Committee for Human Medicinal Products (CHMP). Background review for the excipient propylene glycol. *Eur Med Agency.* 2014;44(EMA/CHMP/334655/2013):1-96.

Committee for Medicinal Products for Human Use (CHMP). Reflection paper on the use of methyland propylparaben as excipients in human medicinal products for oral use. *Eur Med Agency*. 2015;44(EMA/CHMP/SWP/272921/2012):1-3.

http://www.ema.europa.eu/docs/en_GB/document_library/Scientific_guideline/2013/05/WC50014313

Kozarewicz P, European Medicines Agency (EMA). Preservatives: Are they safe? *Eur Med Agency.* 2010;(May).

European Food Safety Authority (EFSA). Opinion of the Scientific Panel on Food Additives, Flavourings, Processing Aids and Materials in Contact with Food on a Request from the Commission related to para hydroxybenzoates (E 214-219). *EFSA J.* 2006;965(October 2000):1-7. doi:10.2903/j.efsa.2007.428.

All Pharmacology/Various



