Intravenous Immunoglobulin

IVIG

2018

For newborn use only

Alert	If the patient has ANY adverse reaction, stop infusion and call a medical officer			
	IMMEDIATELY.			
	I his formulary is for intragam 10.			
	Intragam 10 is the domestically produced intravenous immunoglobulin (IVig) and is the most likely product that you will receive from the Australian Blood Service			
	Intergram P (6%) is no longer produced as of lune 2018			
	Flehogamma 5% and 10% should not be given to reonates due to undiagnosed bereditary			
	frictose intolerance			
	Other preparations such as Privigen 10 are available for paediatric use, but beyond the scope			
	of this formulary.			
Indication	1 Neonatal alloimmune thrombocytonenia (NAIT)			
	2 Haemolytic disease of the newborn (HDN) (isoimmunisation)			
	3. Immune thrombocytopenic purpura			
	4. Primary immunodeficiency diseases			
	5. Secondary hypogammaglobulinaemia			
	6. Neonatal haemochromatosis – gestational alloimmune liver disease (GALD)			
	7. Neonatal myasthenia gravis			
	8. Severe neonatal enterovirus infection including myocarditis or hepatitis			
	9. Sepsis/infection – prevention and treatment – NOT RECOMMENDED.			
	1, 3-5, and 7 are approved indications by the National Blood Authority of Australia, 6 is a			
	proposed addition as of June 2018.			
	See <u>https://www.criteria.blood.gov.au/</u> for a comprehensive list.			
	Indications not funded under the Criteria for the clinical use of intravenous immunoglobulin			
	in Australia (Criteria), may be provided for locally under Jurisdictional Direct Orders			
	(https://www.blood.gov.au/system/files/documents/jdo-factsheet.pdf).			
Action	Immunoglobulin G (IgG) provides humoral immunity and is an immune modulator. [19]			
Drug Type	Immunoglobulin			
Trade Name	Intragam 10. Contains 1 g of immunoglobulin G in 10 mL.			
Presentation	Intragam 10 is a 10% w/v solution of IgG produced by CSL Behring from voluntary donors to			
	the Australian Red Cross. Intragam [®] 10 comes in 2.5 g in 25 mL, 10 g in 100 mL and 20 g in			
	200 mL. All these strengths provide 1 g of Ig in 10 mL.			
	Donors are screened for antibodies to HIV and Hepatitis B and C.			
Dosage / Interval	Medical officer should prescribe (1) brand of IVIg and the % concentration (e.g. Intragam 10),			
	(2) dose in grams and the volume in mL (e.g. 2 g/20 mL) and (3) Rate of infusion (see			
	Administration section)			
	Isoimmunisation:			
	1 g/kg (range 0.5–1.5 g/kg) IV. Dose may be repeated in 12–24 hours if required.			
	Neonatal alloimmune thrombocytopenia (NAIT):			
	1 g/kg IV. Repeat if required.			
	Immune thrombocytopenic purpura (ITP):			
	1 g/kg IV. Repeat if required. Immunodeficiency:			
	U.4 g/kg IV (dose should be based on number of infections and trough serum IgG			
	Concentration [optimally above 6 g/L, nigner if there is bronchiectasis]).			
	I alka IV daily for 2 days (total docs) 2 alka). If additional therapy required titrate			
	grapher dinical response [0]			
	against clinical response.[9]			
	severe enterovirus infection/myocarditis or nepatitis:			

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	2 g/kg IV (up to 2.5 g/kg) as a single dose within 3 days of onset.			
	Sepsis/intection (prevention or treatment) – not recommended:			
	0.5 to 1 g/kg IV repeated at intervals when required has been used.			
	Neonatal haemochromatosis:			
	1–2 g/kg/day IV following exchange transfusion in the first 7 days and then 1 g/kg			
	weekly, as required.			
Maximum daily dose	2 g/kg/day.			
	Enterovirus infection: 2.5 g/kg/day			
Route	Intravenous.			
Preparation/Dilution	Obtain written consent from parent or guardian.			
	All opened bottles must be used immediately.			
	Do not shake bottles to avoid foaming.			
	A 'peel-off' identification label with Batch Number and Expiry Date is to be placed on th			
	patient's Blood Component order form.			
	Allow preparation to reach room temperature and inspect for turbidity or sediments. If seen			
	return to Blood Bank			
Administration	Infusion rate: 0.5 ml /kg/hour for 60 minutes: then 1 ml /kg/hour for next 60 minutes: 2			
Administration	mission rate. 0.5 mis/kg/nour for ob minutes; then 1 mis/kg/hour (at a maximum rate of 25 mis/hour)			
	To be checked by two Peristered Nurses			
	Bequires a surgically clean procedure			
	Civen via intravenous control line long line or part			
	Given via intravenous cannula, central line, long line or port.			
	Administered by musion pump. A blood filter is not required, but may be used			
	A blood filter is not required, but may be used. Sadium shlasida 0.0% assumed as a flush at the and of the infusion			
	• Sodium chloride 0.9% may be used as a flush at the end of the infusion.			
Monitoring	Vital sign monitoring of temperature, heart rate, respiratory rate and blood pressure to be			
	recorded before commencement of infusion.			
	If the patient is unwell or there are any concerns particularly regarding the baseline			
	observations, the medical officer should be contacted before the infusion commences.			
	Vital signs (temperature, heart rate, respiratory rate) should then be checked and recorded:			
	Within 15 minutes after the start of the infusion;			
	Hourly during the infusion;			
	At the end of the infusion.			
Contraindications	Patients who have had an anaphylactic reaction to a human immunoglobulin preparation.			
Precautions				
Drug Interactions	Concurrent use of immunoglobulin and live virus vaccines may result in interference with the			
	immune response to the live vaccine. The Australian Technical Advisory Group on			
	Immunisation (ATAGI) recommendations are below:			
	Hepatitis B vaccine is an inactivated vaccine and can be administered at any time			
	before, after or concurrently with IVIg.			
	Rotavirus vaccine may be administered at any time before, after or concurrently			
	with any blood product, including antibody-containing products.			
	BCG vaccine can be given at any time before or after administration of			
	immunoglobulin or any antibody-containing blood product.			
	Following the receipt of IVIg for ITP treatment, an interval of 8–10 months should			
	elanse before vaccination with an MMR_MMRV or varicella vaccine			
	May result in false-positive Coombs test due to passive transmission of antibodies to			
	ervithrocyte antigens			
	May result in a falsely elevated blood glucose measurement due to assay interforence with			
	the glucoce debudrogeness (purrelequineline guinene) method			
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Adverse Reactions	If adverse reactions occur, the first response should be to stop the infusion, then notify		
	Medical Officer.		
	• Severe reactions are uncommon especially in neonates. In older patients are most like		
	to occur during the first infusion, but may occur subsequently.		
	Anaphylactic reactions are rare: urticaria, angioedema, bronchospasm and hypotension		
	Anaphylactic reactions may require oxygen, adrenaline (epinephrine) and steroids		
	depending on severity of the reaction.		
	• More common reactions are: flushing, fever, headache, pallor, shivering and tachycardia.		
	Other reported reactions: dyspnoea, chest tightness, tachycardia or hypotension without		
	anaphylaxis, transient haemolytic anaemia, abdominal pain and renal failure.		
	• Milder reactions often resolve after the infusion has been stopped. If so, after discussion		
	with medical staff, the infusion may be recommenced at a slower rate after at least 15		
	minutes.		
	Subsequent infusions should be commenced and escalated at a slower rate.		
Compatibility	Sodium chloride 0.9% for priming and flushing. Others not tested.		
	Administer through a separate line.		
Incompatibility	Compatibility with other drugs not established.		
Stability	Do not mix immunoglobulin products of different formulations or from different		
	manufacturers.		
Storage	Store at 2 to 8 °C (Refrigerate. Do not freeze). Protect from light.		
	Once removed from refrigeration, unopened bottles of Intragam 10 must be used within		
	three months.		
	Intragam 10 can only be ordered from the Australian Red Cross Blood Service (ARCBS).		
Special Comments	Newborn infants with isoimmunisation who are considered at risk of exchange transfusion		
	must have intensive prophylactic phototherapy as this is the intervention most likely to		
	prevent the need for exchange transfusion.		
	If not yet done – newborn screening (NBS) should be performed prior to infusion and		
	repeated as per blood transfusion/NBS policy.		
Evidence summary	Refer to full version.		
References	Refer to full version.		

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