

INTRAVENOUS IMMUNOGLOBULIN (IVIg) Authorisation Request Form (Effective from 1 Sept 2015) HAEMATOLOGICAL INDICATIONS FOR NEUROLOGICAL & IMMUNOLOGICAL INDICATIONS PLEASE USE DEDICATED FORM



PLEASE FAX COMPLETED FORM TO RCH BLOOD BANK LAB ON 9345 5817

| Once complete, signed and dated, please FAX: For enquiries and urgent requests please PHONE: AFTER HOURS PHONE: | | | | | | |
|--|---|--|--|--|--|--|
| State/Territory: VICTORIA | | | | | | |
| Requesting Medical Officer Name: | Position: | | | | | |
| Pager/Mobile: Phone: | Fax: Date: | | | | | |
| PATIENT DETAILS (or affix hospital label) | PRODUCT DELIVERY INSTRUCTIONS | | | | | |
| Surname: | Dispenser (hospital blood bank/pathology laboratory/pharmacy/private pathology) Dispenser name: | | | | | |
| Given names: | Street: | | | | | |
| DOB: | Suburb: | | | | | |
| Gender: Female Male | State/Territory: | | | | | |
| UR: | Postcode: | | | | | |
| Hospital: | Phone: Fax: | | | | | |
| Weight: kg Height: cm | Email: | | | | | |
| Previous Immunoglobulin treatment: | | | | | | |
| IVIg 🔄 Subcutaneous Immunoglobulin (SCIg) 📄 Unknown 📄 Normal Human Immunoglobulin (NHIg) 🦳 | dditional delivery instructions: | | | | | |
| Please provide details (including date, product and response, if known): | | | | | | |
| | | | | | | |
| | | | | | | |
| ITP Adult Paediatric In pregnancy Refractory to steroids Steroids contraindicted Feto-maternal/neonatal alloimmune Maternal Neonatal thrombocytopenia Post transfusion purpura For all these conditions, please indicate: Platelet Count Detail Bleeding Detail other treatment including steriod use: Other haematological conditions (please specify) | TACHED TO DEMONSTRATE THAT ALL QUALIFYING CRITERIA HAVE BEEN MET. Acquired hypogammaglobulinaemia secondary to haematological malignancies (please select one of the below) CLL Multiple Myeloma NHL HSCT Other relevant haematological malignancy (please specify) Recurrent or severe bacterial infection(s) Yes No Details of bacterial infection(s) Total serum IgG g/L Date Performed: Clinically active bronchiectasis Yes No Anti B-cell therapy e.g. Rituximab Yes No | | | | | |
| Dose required: g OR Number of doses planne | ed (e.g. 2x24g): DOSE/kg: | | | | | |
| Frequency: Once only Monthly Other (please specify) Date required: | | | | | | |
| IMPORTANT: Your patient will be allocated either Intragam P 6% or an imported IVIg product provided your order meets policy requirements for the supply of IVIg for clinical indications funded under the Criteria. Some hospitals have local policies for imported IVIg product. Please check with your blood and blood products Dispenser (blood bank, pathology laboratory, pharmacy or private pathology). | | | | | | |
| Please indicate your preferred imported IVIg product: Available until 31 December 2015* Kiovig 10% Octagam 5% Octagam 10% | | | | | | |
| * These products are available for existing patients until 3 | | | | | | |
| Available from 1 November 2015 | Privigen 10% Flebogamma 5% Flebogamma 10% | | | | | |
| OFFICE USE ONLY (Blood Service authorisation) Delegate: Designation (MO/TN/Other): | | | | | | |
| Qualifying Criteria met not met Request approved yes | no Referred to JDO/IVIg Group for Review: yes no | | | | | |
| Product: | g Frequency: Australian Red Cross | | | | | |
| Review required by: (continuing supply will be conditional on this review) DLUUD SCRVIDE This fax message and any attached files may contain information that is confidential including health information intended only for use by the individual or entity to whom they are addressed. If you are not This fax message and any attached files may contain information that is confidential including health information intended only for use by the individual or entity to whom they are addressed. If you are not | | | | | | |

the intended recipient or the person responsible for delivering the message to the intended recipient, be advised that you have received this message in error. To protect the privacy of individuals in this form you should notify the sender immediately and shred the fax.



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HAEMATOLOGICAL INDICATIONS

FOR NEUROLOGICAL & IMMUNOLOGICAL INDICATIONS PLEASE USE DEDICATED FORM

| Patient details | : | | | | |
|--|---------------------------|-------------------------------|--------------|--------|--|
| Surname: | | | Given names: | | |
| DOB: | | | Hospital: | | |
| Requesting Me | edical Officer Name: | | Position: | | |
| Pager/Mobile: | | Phone: | Fax: | Date: | |
| Consultant Co | nfirming Diagnosis: | | Specialty: | | |
| Name: | | | Phone: | | |
| Email: | | | Mobile: | | |
| Postal Address: | | | | | |
| IMPORTANT: The addresses above will be used for any relevant future correspondence, including the patient treatment review outcome notification form for patients requiring continuing access to government funded immunoglobulin product. For some conditions, the Criteria require review by a specialist in a specific discipline – please refer to the Criteria. | | | | | |
| Prescriber acknowledgement and confirmation (to be completed by the treating medical specialist or appropriate delegate following discussion with their patient) I acknowledge the governance and management arrangements for the appropriate supply and use of immunoglobulin products, funded under the national blood arrangements, and the provision of information required to support authorisation. To the best of my knowledge, the information provided in this form and attachments is true and correct. I have provided and/or explained to my patient (or parent/carer/guardian) the Privacy Statement and Notice (Notice) and Patient Information Brochure and they have had the opportunity to ask questions. I believe that they are aware of and understand: the risks and benefits of treatment with immunoglobulin products and alternative treatments (where these exist), the national blood arrangements, including that immunoglobulin products may need to change from time to time (for patients requiring ongoing treatment only) the nature of ongoing monitoring and review and that access to product will cease if response to treatment does not demonstrate clinical benefit. I confirm that my patient (or parent/carer/guardian) has provided express consent (explicit verbal or written consent) to: the collection and recording of personal information (including sensitive health information) in secure databases, held by the Australian Red Cross Blood Service) and the National Blood Authority (NBA). the use of limited identifying details (for example, name, date of birth, sex and hospital identifiers) within search functions of the above mentioned databases to ensure that patients are correctly identified. the disclosure and use of this information in a manner which will not readily identify them, (such as through the removal of directly identifying personal information in a manner which will not readily identify them, (such as through the removal of directly identifying per | | | | | |
| My patient understands that any additional use of information held by the Blood Service and NBA will only be undertaken in accordance with the requirements of the Privacy Act 1988 (Cth) and any relevant state/territory laws, and that the information may be made available for medical or public health research only with approval of a properly constituted human research ethics committee (HREC). | | | | | |
| Signature: | τεσεαιτη υπιγ ωτη αρφιονά | οι α ριορειιγ τοποτιτατέα ηθη | Date | | |
| Name: | | | | ition: | |
| The Australian Red Cross Blood Service is contracted by the National Blood Authority to perform the roles of Authoriser and Distributor of immunoglobulin products supplied and funded under the national blood arrangements. | | | | | |
| YOU MUST SELECT YOUR STATE OR TERRITORY BEFORE PRINTING PRINT | | | | | |
| This fax message and any attached files may contain information that is confidential including health information intended only for use by the individual or entity to whom they are addressed. If you are not the intended recipient or the person responsible for delivering the message to the intended recipient, be advised that you have received this message in error. To protect the privacy of individuals in this form you should notify the sender immediately and shred the fax. Form ID - NBA301001 Effective from 1 Sept 2015 Uncontrolled version when printed Page 2 of 2 | | | | | |