Ontario MOHLTC IG Request Form For Neurology Use Only

Patient Name:

Patient Hospital/Medical Record#:

Patient DOB (YYYY/MM/DD):

Gender M/F:

Location:

Ontario Health Insurance#:

ALL FIELDS BELOW ARE MANDATORY

SECTION A: Physician & Hospital Information

Date of Request (YYYY/MM/DD)	Date Required (YYYY/MM/DD)	Hospital Transfusion Service (HTS) Fax Number
Name of Ordering Physician	Physician's Contact Phone Number	Physician's Email
Is the patient being seen by a Neurologist/ Neuromuscular Specialist? Yes No	Is the request for a hospital inpatient? □ Yes □ No	Hospital where patient will receive IG

SECTION B: Request Type

Initial Request: Maximum 6 month approval	Renewal Request: A reassessment should be done to confirm IG treatment continues to be effective and
	that minimum effective dose is being applied. Maximum 12 month approval.

SECTION C: Clinical Indication Refer to Ontario IG Management Utilization Guidelines for additional indications where IG may be appropriate

ApprovedCondition	Guidelines for INITIAL Request	Guidelines for RENEWAL Request	
Guillain–Barré Syndrome (GBS) including Miller Fisher Syndrome and other variants	 IG recommended for Grade 3 severity (able to walk with aid greater; or less than Grade 3 severity that are progressing. IG should be given within 2 weeks of symptom onset. Adult: Total Dose of 2 g/kg divided over 2 to 5 days. Pediatric: Total Dose of 2 g/kg divided over 2 days. 	 IG treatment for GBS is typically one-time/in the acute setting. Re-treatment for patients who do not respond may be considered. Repeat treatment with IVIG at 2g/kg divided over 2-5 days. 	
Myasthenia Gravis (MG)	 IG is recommended as first-line treatment in moderate-sever MG or in myasthenic crisis. Induction Dose: 2g/kg divided over 2-5 days. Initial requests may be made for induction plus two maintenance doses; fill out Section D accordingly. 	 IG in combinations with immunosuppressive therapy can be considered in refractory cases. If additional IG is required, dose should be adjusted depending upon response and titrated to the minimum effective dose. Maintenance Dose: 1g/kg 	
Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)	 IG is recommended as first-line therapy in CIDP. Induction Dose: 2 g/kg divided over 2 to 5 days. All patients receiving IG for chronic treatment of CIDP should followed by a neuromuscular specialist. 	 Immunosuppressive therapy in combination with IG can be considered in refractory cases. Continued use should be based on objective measures of sustained effectiveness. Aim for minimum effective dose. Maintenance Dose: 1g/kg every3 weeks. 	
Multifocal Motor Neuropathy (MMN)	 IG is recommended as first-line treatment for MMN. Induction Dose: 2g/kg divided over 2-5 days. 	 Maintenance Dose: Tailor to the lowest dose that maintains clinical efficacy, usually 1g/kg or less per treatment course. Some patients may require higher doses for efficacy, up to 2g/kg every 4 weeks. 	
Other (please specify the These requests will require unapproved indication.	· · ·	ding treatment to date and documentation to support IG treatment for an	
Has the patient used ot	her therapies to treat this condition? \Box Yes, specify	ther treatments below 🛛 No	
Treatment	Dose (if applicable)	Duration of treatment What was the outcome?	
		□ No response □ Contraindications □ Intolerance	

Other Comments: (include notes regarding response to IG therapy)

SECTION D: Dosage Information (Verification of dose using Dose Calculator tool is recommended. Refer to http://ivig.transfusionontario.org/dose/

Patient Weight:	kg	Patient Height:	cm	BMI:	Dose must	be adjusted for BMI greater	r than or equal to 30
Induction/One-time dose		g/kg = Total dose of		g; divided over	days		
Maintenance dose		g/kg = Total dose of		g; divided over	days; every	weeks; Duration:	months

Dose Calculator Used?
Yes No If No, why was it not used?

SECTION E: For Transfusion Medicine Use Only

Section E. For Transitision medicine ose only					
Dose verified	Dose adjusted to:	By (signature req'd):			
Confirmed with ordering physician		Date:			
Approved	Denied				
Signature of Approving Physici	ian or designate:		Date:		

□ Intolerance

 \Box No response \Box Contraindications