

## CIDP PRESCRIPTION REFERRAL FORM

Please fax completed form to 1-855-710-7035

20	<b>imunex-c</b>
	immune globulin injection (human), 10%
2-	caprylate/chromatography purified

PATIENT INFORMATION	:						_
Name:				Date of birth:	/ /	Male	Female
Address:				Height:	(ft/in) Weig	ght: lb	kg
City:	S	State: Z	IP:	Diagnosis: ICD-10	☐CIDP G61.81	Other:	
Phone:	Email:			Current therapy:			
Parent/guardian name (if pat	tient is under 18):						
PATIENT INSURANCE INFO	ORMATION: <b>Please pr</b>	ovide a co	py of the <u>fror</u>	nt and back of ins	urance card		
COORDINATION OF CARE:							
Preferred site of care: Out	tpatient infusion center	Physicia	n office H	ome Other:			
Facility name:	cility name: Contact name:					Phone:	
PRESCRIBING INFORMATI with a loading dose of 2 g/l						study dosing p	rotocol
LOADING DOSE:	g/kg IV divided	over:	day(s)				
MAINTENANCE DOSE:	g/kg IV divided	over:	day(s)	every:	week(s)		
ROUTE: Peripheral IV	Central line	REFILLS	: times				
PREMEDICATE: Yes	No Acetaminophen	(oral) 650 m	g Diphenh	nydramine (oral)			
Other medications (specify): Drug allergies:							
Nursing services:							
Anaphylaxis kit:							
Provide any medical/ancil	lary supplies as necessa	ary to safely	administer pres	scribed medication			
Other order(s):							
PRESCRIBER INFORMATION	ON AND SIGNATURE: F	Required to pro	ocess				
Physician name (print):				Office contact:			
City:		State:		ZIP:			
Phone:	Email:			Fax:			
Facility or prescriber tax ID#	:	NPI#:		DEA#:			
I verify that the patient and prescrib based on my professional judgmen other personal information as may and contracted third parties to forw than 18 years I attest that I have of	t and medical necessity. I als be necessary to Gamunex C vard this prescription electron	so attest that I honnexions and/ nically, by facsin	nave obtained the p or their agents. I a nile, or by mail to t	patient's affirmative author authorize Grifols and its a	orization to release the ffiliated companies, a	e above informati gents and repres	ion and such entatives,
PHYSICIAN SIGNATURE:				Date:			
DISPENSE AS WRITTEN: E	Exact terminology may l	oe based on	state regulation	ns. Please provide st	ate-specific presci	ription languaç	ge here:



If you have questions, please call Gamunex Connexions toll free at 1-888-MYGAMUNEX (694-2686), Monday to Friday from 8 AM to 8 PM EST.

Please see Important Safety Information on the next page and refer to accompanying full Prescribing Information for GAMUNEX-C. CIDP=chronic inflammatory demyelinating polyneuropathy.

## **Important Safety Information**

GAMUNEX®-C (immune globulin injection [human], 10% caprylate/chromatography purified) is indicated for the treatment of primary humoral immunodeficiency disease (PIDD) in patients 2 years of age and older, idiopathic thrombocytopenic purpura (ITP) in adults and children, and chronic inflammatory demyelinating polyneuropathy (CIDP) in adults.

Thrombosis may occur with immune globulin products, including GAMUNEX-C. Risk factors may include: advanced age, prolonged immobilization, hypercoagulable conditions, history of venous or arterial thrombosis, use of estrogens, indwelling central vascular catheters, hyperviscosity, and cardiovascular risk factors. Thrombosis may occur in the absence of known risk factors. For patients at risk of thrombosis, administer GAMUNEX-C at the minimum dose and infusion rate practicable. Ensure adequate hydration in patients before administration. Monitor for signs and symptoms of thrombosis and assess blood viscosity in patients at risk for hyperviscosity.

Renal dysfunction, acute renal failure, osmotic nephrosis, and death may occur with immune globulin intravenous (IVIG) products in predisposed patients. Patients predisposed to renal dysfunction include those with any degree of preexisting renal insufficiency, diabetes mellitus, age greater than 65, volume depletion, sepsis, paraproteinemia, or patients receiving known nephrotoxic drugs. Renal dysfunction and acute renal failure occur more commonly in patients receiving IVIG products containing sucrose. GAMUNEX-C does not contain sucrose. For patients at risk of renal dysfunction or failure, administer GAMUNEX-C at the minimum concentration available and the minimum infusion rate practicable.

GAMUNEX-C is contraindicated in patients who have had an anaphylactic or severe systemic reaction to the administration of human immune globulin. It is contraindicated in IgA-deficient patients with antibodies against IgA and history of hypersensitivity.

Severe hypersensitivity reactions may occur with IVIG products, including GAMUNEX-C. In case of hypersensitivity, discontinue GAMUNEX-C infusion immediately and institute appropriate treatment.

Monitor renal function, including blood urea nitrogen (BUN), serum creatinine, and urine output in patients at risk of developing acute renal failure.

Hyperproteinemia, increased serum viscosity, and hyponatremia may occur in patients receiving IVIG treatment, including GAMUNEX-C.

There have been reports of aseptic meningitis, hemolytic anemia, and noncardiogenic pulmonary edema (transfusion-related acute lung injury [TRALI]) in patients administered with IVIG, including GAMUNEX-C.

The high-dose regimen ( $1g/kg \times 1-2 \text{ days}$ ) is not recommended for individuals with expanded fluid volumes or where fluid volume may be a concern.

Because GAMUNEX-C is made from human blood, it may carry a risk of transmitting infectious agents, eg, viruses, the variant Creutzfeldt-Jakob disease (vCJD) agent, and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent.

Do not administer GAMUNEX-C subcutaneously in patients with ITP because of the risk of hematoma formation.

Periodic monitoring of renal function and urine output is particularly important in patients judged to be at increased risk of developing acute renal failure. Assess renal function, including measurement of BUN and serum creatinine, before the initial infusion of GAMUNEX-C and at appropriate intervals thereafter.

Consider baseline assessment of blood viscosity in patients at risk for hyperviscosity, including those with cryoglobulins, fasting chylomicronemia/markedly high triacylglycerols (triglycerides), or monoclonal gammopathies, because of the potentially increased risk of thrombosis.

If signs and/or symptoms of hemolysis are present after an infusion of GAMUNEX-C, perform appropriate laboratory testing for confirmation.

If TRALI is suspected, perform appropriate tests for the presence of antineutrophil antibodies and anti-HLA antibodies in both the product and patient's serum.

After infusion of IgG, the transitory rise of the various passively transferred antibodies in the patient's blood may yield positive serological testing results, with the potential for misleading interpretation.

In clinical studies, the most common adverse reactions with GAMUNEX-C were headache, pyrexia, hypertension, chills, rash, nausea, arthralgia, and asthenia (in CIDP); cough, rhinitis, pharyngitis, headache, asthma, nausea, fever, diarrhea, and sinusitis with intravenous use (in PIDD) and local infusion-site reactions, fatigue, headache, upper respiratory tract infection, arthralgia, diarrhea, nausea, sinusitis, bronchitis, depression, allergic dermatitis, migraine, myalgia, viral infection, and pyrexia with subcutaneous use (in PIDD); and headache, ecchymosis, vomiting, fever, nausea, rash, abdominal pain, back pain, and dyspepsia (in ITP).

The most serious adverse reactions in clinical studies were pulmonary embolism (PE) in 1 subject with a history of PE (in CIDP), an exacerbation of autoimmune pure red cell aplasia in 1 subject (in PIDD), and myocarditis in 1 subject that occurred 50 days post-study drug infusion and was not considered drug related (in ITP).

Please see accompanying full Prescribing Information for GAMUNEX-C.

References: 1. GAMUNEX®-C (immune globulin injection [human], 10% caprylate/chromatography purified) Prescribing Information. Grifols. 2. Hughes RAC, Donofrio P, Bril V, et al. Intravenous immune globulin (10% caprylate chromatography purified) for the treatment of chronic inflammatory demyelinating polyradiculoneuropathy (ICE study): a randomised placebo-controlled trial. *Lancet Neurol*. 2008;7(2):136-144.



