

MEDICARE FORM

Immune Globulin (IG) Therapy Medication and/or Infusion Precertification Request

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(All fields must be completed and legible for precertification review.)

Gammaplex, Hyqvia, and Panzyga are non-preferred. The preferred products are Gammaked, Gamunex-☐ Start of treatment: Start date / / Please indicate: C, Hizentra, Octagam, Privigen and Continuation of therapy: Date of last treatment ____/_ / Xembify. Precertification Requested By: ___ Phone: ___ A. PATIENT INFORMATION First Name: Last Name: DOB: ZIP: Address: City: State: Work Phone: Cell Phone: Home Phone: Email: Current Weight: ___ lbs or ____ kgs Height: _ inches or cms Allergies: **B. INSURANCE INFORMATION** Aetna Member ID #: Does patient have other coverage? ☐ Yes ☐ No Group #: _____ If yes, provide ID#: _____ Carrier Name: ____ Insured: ___ Insured: ____ **Medicaid:** ☐ Yes ☐ No If yes, provide ID #: **Medicare:** ☐ Yes ☐ No If yes, provide ID #: C. PRESCRIBER INFORMATION (Check One): M.D. D.O. N.P. P.A. First Name: Last Name: Address: City: State: ZIP: UPIN: Phone: Fax: St Lic #: NPI#: DEA #: Office Contact Name: Provider Email: Phone: D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION **Dispensing Provider/Pharmacy:** Place of Administration: ☐ Self-administered ☐ Physician's Office ☐ Physician's Office ☐ Retail Pharmacy Home ☐ Specialty Pharmacy ☐ Mail Order ☐ Other: Outpatient Infusion Center Phone: ____ Center Name: ____ ☐ Home Infusion Center Phone: _____ Address: Agency Name: __ Phone: _____ Fax: ____ Administration code(s) (CPT): Address: PIN: TIN: E. PRODUCT INFORMATION Request is for: Asceniv ☐ Bivigam ☐ Cutaquig ☐ Cuvitru ☐ Flebogamma ☐ Gamastan S/D ☐ Gammaked ☐ Gammagard ☐ Gammaplex ☐ Gamunex-C ☐ Hizentra ☐ HyQvia ☐ Octagam
☐ Panzyga
☐ Privigen Xembify _____ Frequency: __ HCPCS Code: _____ ____ | IV | IM | SC F. DIAGNOSIS INFORMATION – Please indicate primary ICD Code and specify any other where applicable. Primary ICD Code: Secondary ICD Code: Other ICD Code: **G. CLINICAL INFORMATION** – Required clinical information must be completed in its entirety for all precertification requests. Please provide the current immunoglobulin levels: Immunoglobulin A (IgA) level and date obtained: _____ Date: ____/ Date: ____/ / Immunoglobulin G (IgG) level and date obtained: Immunoglobulin M (IgM) level and date obtained: For All Requests: (Clinical documentation required for all requests) Note: Asceniv, Bivigam, Cutaquig, Cuvitru, Flebogamma, Gammagard, Gammaplex, Hyqvia and Panzyga, are non-preferred. The preferred products are Gammaked, Gamunex-C, Hizentra, Octagam, Privigen and Xembify. ☐ Yes ☐ No Has the patient had prior therapy with the requested immune globulin product within the last 365 days? ☐ Yes ☐ No Has the patient had a trial and failure, intolerance, or contraindication to Gammaked, Gamunex-C, Hizentra, Octagam, Privigen or Xembify? Please explain if there are any other medical reason(s) that the patient cannot use Gammaked, Gamunex-C. Hizentra, Octagam, Privigen or Xembify. ☐ Yes ☐ No Is the patient changing to a different immunoglobulin product? ☐ Yes ☐ No Does the patient have immunoglobulin A (IgA) deficiency with anti-IgA antibodies?

Continued on next page

For New Jersey HMO D-SNP:

For other lines of business:

Please use other form.

1-833-322-0034 PHONE: 1-844-362-0934 (TTY: 711)

Note: Asceniv, Bivigam, Cutaquig, Cuvitru, Flebogamma, Gammagard,

FAX:



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.						
For All requests continued: Please indicat	e which of the following applies to th	e patient and answer subsequent que	estions			
Acquired red cell aplasia						
☐ Acute disseminated encephalomyelitis☐ Autoimmune mucocutaneous blistering of	lianana					
Please select which applies to the		☐ Epidermolysis bullosa acquisita	a ☐ Gestational Pemphigoid			
Flease select which applies to the	Linear IgA disease	☐ Mucous membrane pemphigoid				
	☐ Pemphigus vulgaris	☐ Pemphigus foliaceus	□ None of the above			
☐ Yes ☐ No Has patient failed			Thomas of the above			
Yes No Does the patient have contraindications to conventional therapy?						
	Yes No Does the patient have rapidly progressive disease in which a clinical response could not be					
		uickly enough using conventional age	ents?			
Autoimmune hemolytic anemia (refractor	ry)					
Autoimmune neutropenia (refractory)						
B-cell chronic lymphocytic leukemia (CLL)						
Yes No Does the patient have hypogammaglobulinemia associated with CLL?						
Yes No Does the patient have recurrent infections or specific antibody deficiency? Birdshot (vitiligenous) retinochoroidopathy						
☐ BK virus associated nephropathy						
☐ Chronic inflammatory demyelinating polyneuropathy (CIDP)						
Yes No Has the patient responded to previous intravenous immune globulin (IVIG) therapy?						
☐ Churg-Strauss Syndrome (CSS) (allergic granulomatosis)						
☐ Yes ☐ No Will IVIG be used as adjunctive therapy for persons with severe active illness?						
	rentions been unsuccessful, become					
☐ Dermatomyositis	ich applies: Unsuccessful Int	olerable				
1 — _ ' _	as adjunctive therapy for persons w	no have had an inadequate response	to first and second line theranies?			
Yes No Will this be used as adjunctive therapy for persons who have had an inadequate response to first and second line therapies? Enteroviral meningoencephalitis						
☐ Guillain-Barre Syndrome (GBS) and GBS variants						
Yes No Has the patient been diagnosed during the first 2 weeks of illness?						
☐ Yes ☐ No Does the patient require aid to walk? (must be severely affected)						
☐ Yes ☐ No Does the patient have any contraindications to IVIG?						
☐ Hematophagocytic lymphohistiocytosis (HLH) or macrophage activation syndrome (MAS)						
☐ Yes ☐ No Does the patient have hypogammaglobulinemia? Please indicate the IgG level: ☐ Less than 400mg/dL ☐ 400mg/dl or greater						
☐ Hemolytic disease of newborn	s the IgG level two standard deviatio	is below the mean for age?				
_ ,	decrease the need for exchange tra	nsfusion?				
☐ HIV infected children						
☐ Yes ☐ No Is this request for bacterial control or prevention of infection?						
☐ HIV- associated thrombocytopenia (pediatric or adult)						
☐ Hyperimmunoglobulinemia E Syndrome						
Yes No Is this request for						
Immune or Idiopathic thrombocytopenic	. ,	racry to control evenesive blooding of	or to defer or avoid enlangetemy)?			
		rgery, to control excessive bleeding, or	Date:/			
☐ Kawasaki Disease	arrent platelet count and date collect	cu	Batc			
☐ Lambert-Eaton myasthenic syndrome						
☐ Moersch-Woltmann (Stiff-man) syndrome (unresponsive to other therapies)						
Multifocal motor neuropathy						
Yes No Does the patient have progressive, symptomatic multifocal motor neuropathy?						
☐ Yes ☐ No Was the diagnosis based on electrophysiologic findings that rule out other possible conditions that may not respond to this treatment?						
☐ Multiple Myeloma ☐ Myasthenia Gravis ☐ Neonatal Alloimmune Thrombocytopenia (NAIT) (also known as Fetal Alloimmune Thrombocytopenia or FAIT)						
 □ Neonatal Hemochromatosis (prophylaxis) □ Opsoclonus-myoclonus □ Paraneoplastic opsoclonus-myoclonus-ataxia associated with neuroblastoma □ Parvovirus B19 infection (chronic with severe anemia) □ Polymyositis in persons who are resistant to first and second line therapies 						
☐ Post-transfusion purpura ☐ Preparatio	n tor tnymoma surgery (to prevent m	yastnenia exacerbation) Primary	numoral immunodeficiency diseases:			



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G. CLINICAL INFORMATION (continued) -		be completed in its <u>entirety</u> for all prece	rtification requests.	
Please indicate which of the following app	•			
	hyperimmunoglobulin M a (Good Syndrome) yndrome)) proaches (i.e., interferons) failed, bee	□ Common variable immunodeficier □ Hypogammaglobulinemia □ Severe combined immunodeficier come intolerable, or contraindicated? aches have become intolerable □ States.	☐ Wiscott- Aldrich Syndrome	
Renal transplantation from live donor with Yes No Is a suitable non-reconstruction of the content of the cont	eactive live or cadaveric donor unaverage donor unaverage donor unaverage donor unaverage donor surgery (such as cardial eases) evere infection for persons meeting of the for allosensitized members undergoing persons with severe active SLE) intions been unsuccessful, become inconsuccessful Intolerable Comparison of the formula of the	ailable (preparative regimen)? ic transplants) and certain diseases selection criteria ing solid organ transplant? intolerable, or are contraindicated? contraindicated		
☐ Toxic shock syndrome or toxic necrotizing fasciitis due to group A streptococcus				
Yes No Has the patient received IVIC Yes No Does the patient received IVIC	d an adequate response to therapy? or life-threatening infections and date 6 within the past 6 months? oratient have a documented severe an one previous infusion?	•	er's current dosage). e event that occurred during or	
H. ACKNOWLEDGEMENT				
Request Completed By (Signature Requ	uired):		Date: //	
Any person who knowingly files a request finsurance company by providing material				

insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.