

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.)

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Please indicate:	☐ Start of treatment: Start date/ _/ ☐ Continuation of therapy: Date of last treatment			
	(All fields must be completed and legible for precertification review.)			

For Ohio MMP:

FAX: 1-855-734-9389 **PHONE:** 1-855-364-0974

For other lines of business:

Please use other form.

Note: Asceniv, Bivigam, Cutaquig, Cuvitru, Flebogamma, Gammagard, Gammaked, Gammaplex, Gamunex-C, Hyqvia, Octagam, Panzyga, and Xembify are non-preferred. The preferred products are Privigen

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Precertification Requested By:		Phone:		Fax:	
A. PATIENT INFORMATION					
First Name:	Las	t Name:			
Address:	City	:		State:	ZIP:
Home Phone:	Work Phone:	1	Cell Phone:	Į.	L
DOB: Allergies:			Email:		
	gs Height:	inches or			
B. INSURANCE INFORMATION	go rioight				
Aetna Member ID #:	Does patient have othe	r coverage?	es 🗌 No		
Group #:	If yes, provide ID#:	_			
Insured:	Insured:				
C. PRESCRIBER INFORMATION					
First Name:	Last Name:		(Check One	e): 🔲 M.D. 🗀] D.O. 🗌 N.P. 🗌 P.A.
Address:		City:		State:	ZIP:
Phone: Fax:	St Lic #:	NPI #:	DEA #:	ι	JPIN:
Provider Email:	Office Contact Name:		Phone:	L	
D. DISPENSING PROVIDER/ADMINISTRATION IN	FORMATION				
Self-administered ☐ Physician's Office Outpatient Infusion Center Phone: Center Name: ☐ Home Infusion Center Phone: Agency Name: ☐ Administration code(s) (CPT): ☐ Address: ☐		☐ Physician's Offic ☐ Specialty Pharm Name: Address: Phone: TIN:	nacy 🗌 Mail	Order Otl	
E. PRODUCT INFORMATION					
Request is for: Asceniv Bivigam Gammagard Gammaplex Gamunex-C Dose: Frequency:	☐ Hizentra ☐ HyQvia	☐ Octagam [HCPCS Code:	☐ Panzyga	☐ Privigen	☐ Xembify
F. DIAGNOSIS INFORMATION – Please indicate pri	mary ICD Code and specify any	other where applicable			
Primary ICD Code:	Secondary ICD Code:		Other ICD Co	ode:	_
G. CLINICAL INFORMATION – Required clinical info	ormation must be completed in it	s <u>entirety</u> for all precert	ification reques	ts.	
Please provide the current immunoglobulin levels Immunoglobulin A (IgA) level and date obtained: Immunoglobulin G (IgG) level and date obtained: Immunoglobulin M (IgM) level and date obtained:				Date:	1 1
For All Requests: (Clinical documentation require Note: Asceniv, Bivigam, Cutaquig, Cuvitru, Flebo are non-preferred. The preferred products are Pri Yes No Has the patient had prior therapy wit Yes No Has the patient had a trial and failure Please explain if there are any other medical reason(gamma, Gammagard, Gamma vigen and Hizentra. h the requested immune globulir e, intolerance, or contraindication	n product within the last n to Privigen or Hizentra	365 days?	ia, Octagam,	Panzyga, and Xembify

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
C CLINICAL INFORMATION (continued)	Paguired clinical information must	he completed in its entirely for all pr	coortification requests
G. CLINICAL INFORMATION (continued) –			
For All requests continued: Please indicate	which of the following applies to the	e patient and answer subsequent que	estions
Acquired red cell aplasia			
Autoimmuno muos sutang que blistoring die			
Autoimmune mucocutaneous blistering dis		□ Enidermelysis bulless esquisit	c Contational Remphissis
Please select which applies to the pa		☐ Epidermolysis bullosa acquisita	
	☐ Linear IgA disease ☐ Pemphigus vulgaris	☐ Mucous membrane pemphigoi☐ Pemphigus foliaceus	□ None of the above
☐ Yes ☐ No Has patient failed o		☐ Fempiligus ioliaceus	☐ Notice of the above
Tes TWO Tras patient failed to	Does the patient have contraindication	ations to conventional therapy?	
			which a clinical response could not be
		enough using conventional agents?	Willow a difficult respense sould not be
☐ Autoimmune hemolytic anemia (refractory		3 3 3	
☐ Autoimmune neutropenia (refractory)	,		
B-cell chronic lymphocytic leukemia (CLL)			
☐ Yes ☐ No Does the patient h		ciated with CLL?	
☐ Yes ☐ No Does the patient h			
☐ Birdshot (vitiligenous) retinochoroidopathy			
☐ BK virus associated nephropathy			
☐ Chronic inflammatory demyelinating polyn	europathy (CIDP)		
☐ Yes ☐ No Has the patient res	ponded to previous intravenous im	mune globulin (IVIG) therapy?	
☐ Churg-Strauss Syndrome (CSS) (allergic of			
☐ Yes ☐ No Will IVIG be used a			
☐ Yes ☐ No Have other interve			
	h applies: 🗌 Unsuccessful 🔲 Int	olerable	
Dermatomyositis	di ti th		to first and accord line the manice?
Yes No Will this be used a	s adjunctive therapy for persons wi	no nave nad an inadequate response	to first and second line therapies?
☐ Enteroviral meningoencephalitis ☐ Guillain-Barre Syndrome (GBS) and GBS	variants		
Yes No Has the patient be		ks of illnoss?	
Yes No Does the patient re			
Yes No Does the patient h		anected)	
☐ Hematophagocytic lymphohistiocytosis (HI	•	rome (MAS)	
Yes No Does the patient h		Torric (MAG)	
	e IgG level: ☐ Less than 400mg/dL	☐ 400mg/dl or greater	
	the IgG level two standard deviation		
☐ Hemolytic disease of newborn	3 -	3	
Yes No Is this request to d	ecrease the need for exchange tra	nsfusion?	
☐ HIV infected children	_		
☐ Yes ☐ No Is this request for b		ection?	
☐ HIV- associated thrombocytopenia (pediat	ric or adult)		
☐ Hyperimmunoglobulinemia E Syndrome			
Yes No Is this request for t			
Immune or Idiopathic thrombocytopenic pu			
Yes No Is a rapid rise in pl			
Please provide cur	rent platelet count and date collect	ed:	Date: //
Kawasaki Disease			
Lambert-Eaton myasthenic syndrome			
Moersch-Woltmann (Stiff-man) syndrome	(unresponsive to other therapies)		
Multifocal motor neuropathy			
Yes No Does the patient h			
			ions that may not respond to this treatment?
☐ Multiple Myeloma ☐ Myasthenia Gravis			
☐ Neonatal Hemochromatosis (prophylaxis)			
Parvovirus B19 infection (chronic with seve			
☐ Post-transfusion purpura ☐ Preparation	τοr τnymoma surgery (to prevent m	yastnenia exacerbation) 🔲 Primary	/ numoral immunodeficiency diseases:



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued) – F	Required clinical information must be com	oleted in its entirety for all precertif	ication requests	
		octed in its <u>charety</u> for an precent	ication requests.	
Please select: ☐ Standard approached ☐ Renal transplantation from live donor with A ☐ Yes ☐ No Is a suitable non-real ☐ Secondary immunosuppression associated (extensive burns, or collagen-vascular diseation ☐ Selective IgG subclass deficiencies with seven ☐ Solid organ transplantation ☐ Yes ☐ No Will IVIG be used for ☐ Staphylococcal Toxic Shock Syndrome ☐ Stem cell or bone marrow transplantation ☐ Systemic lupus erythematosus (SLE) (for per ☐ Yes ☐ No Have other intervented.	(X-linked agammaglobulinemia)	ave become intolerable Stand preparative regimen)? clants) and certain diseases n criteria d organ transplant? ple, or are contraindicated?	☐ Wiscott- Aldrich Syndrome ☐ None of the Above	
Yes No Has the patient received IVIG v Yes No Does the patient following the	an adequate response to therapy? If Yes life-threatening infections and dates of occ	currences as well as the member's stentially life-threatening adverse e	current dosage).	
H. ACKNOWLEDGEMENT				
Request Completed By (Signature Require	red):		Date: / /	
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent 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.