V dC	LIIG	Immune Globulin (IG) Therapy Medication and/or Infusion Precertification Request Page 1 of 3 (All fields must be completed and legible for precertification review.)				PHONE: <u>1-855-676-5772 (TTY: 711)</u> For other lines of business: Please use other form. Note: Asceniv, Bivigam, Cutaquig, Cuvitru, Flebogamma, Gammagard, Gammaplex, Hyqvia, and Panzyga				
Please indicate:			atment: Start date / _/ on of therapy: Date of last treatment / _/				are non-preferred. The preferred products are Gammaked, Gamunex- C, Hizentra, Octagam, Privigen and Xembify.			
Precertification R	lequested By:				Phone:			Fax:		
A. PATIENT INFOR	RMATION									
First Name:			Last Name:					DOB:		_
Address:				City	/:			State:	ZIP:	
Home Phone:		Work Phone:		Cel	I Phone:			Email:		
Current Weight:	lbs or	kgs Height:	inches or cms	Alle	rgies:					
B. INSURANCE IN	FORMATION									
Aetna Member ID	#:		Does patient have o		•	Yes 🗌 No				
-			If yes, provide ID#:		Ca	arrier Name:				
Insured:			Insured:							
Medicare: Yes	□ No If yes, pr	ovide ID #:	N	Nedi	caid: 🗌 Yes 🗌 No	If yes, provide	e ID :	#:		
C. PRESCRIBER II	NFORMATION									
First Name:			Last Name:	<u> </u>		(Check Or		□ M.D. □ D	1	□ P.A.
Address:					City:	<u> </u>		I	ZIP:	
Phone:	Fax		St Lic #:		NPI #:	DEA #:		UPI	N:	
Provider Email:			fice Contact Name:			Phone:				
		NISTRATION INFORM	IATION							
	red Phy sion Center ame:	Phone:	_ Home		Dispensing Provi	ffice	tail P il Ore	der 🗌 Other		
	lame:				Address:					
					Phone:					
Address:					TIN:			PIN:		
E. PRODUCT INFO										
Request is for: Gammagard Dose:	Gammaplex		☐ Cutaquig    Cuv ☐ Hizentra    ☐ HyC		☐ Flebogamma ☐ Octagam HCPCS Code: _	🗌 Panzyga		Privigen	ammaked Xembify ] IV          IM	-
F. DIAGNOSIS INF	ORMATION - P	lease indicate primary	ICD Code and specify	any	other where applicab	ole.				
Primary ICD Code:	·	Secor	ndary ICD Code:			Other ICD 0	Code	э:		
G. CLINICAL INFO	RMATION - Red	quired clinical informat	tion must be completed	in its	entirety for all prece	ertification reque	ests.			
Please provide the Immunoglobulin A ( Immunoglobulin G (	(IgA) level and da (IgG) level and da	oglobulin levels: ate obtained: ate obtained:						Date:	     	
For All Requests: ( Note: Asceniv, Biv are Gammaked, G Yes No Hat Yes No Hat Please explain if the Yes No Is t	(Clinical docume vigam, Cutaquig amunex-C, Hize s the patient had s the patient had ere are any other the patient chang	entation required for J, Cuvitru, Flebogamn entra, Octagam, Privig prior therapy with the a trial and failure, intol r medical reason(s) tha ping to a different immu	all requests) ma, Gammagard, Gam gen and Xembify. requested immune glot plerance, or contraindica at the patient cannot use unoglobulin product?	bulin ation e Ga	olex, Hyqvia and Par product within the las to Gammaked, Gam mmaked, Gamunex-0	<b>nzyga, are non</b> ist 365 days? iunex-C, Hizenti	ra, O	<b>ferred. The pr</b> o	<b>eferred pro</b> Jen or Xemb	
	es the patient hav	ve immunoglobulin A (	(IgA) deficiency with ant	ti-IgA	vantibodies?					

For Michigan MMP:

FAX:

1-844-241-2495

PHONE: 1-855-676-5772 (TTY: 711)

**♦aetna** 

**MEDICARE FORM** 



## **MEDICARE FORM**

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

Page 2 of 3

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

For Michigan MMP: FAX: <u>1-844-241-2495</u> PHONE: <u>1-855-676-5772 (TTY: 711)</u>

For other lines of business: Please use other form.

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.

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 indicate	which of the following applies to the	e patient and answer subsequent quest	lions					
Acquired red cell aplasia								
Acute disseminated encephalomyelitis								
Autoimmune mucocutaneous blistering di	seases							
Please select which applies to the p		🗌 Epidermolysis bullosa acquisita	Gestational Pemphigoid					
	🗌 Linear IgA disease	Mucous membrane pemphigoid (	cicatrical pemphigoid)					
	🗌 Pemphigus vulgaris	Pemphigus foliaceus	None of the above					
Yes No Has patient failed								
$\rightarrow$ Yes $\square$ No Does the patient have contraindications to conventional therapy?								
		have rapidly progressive disease in wh						
Autoimmune hemolytic anemia (refractory		uickly enough using conventional agen	15 ?					
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) retinochoroidopath		, ,						
BK virus associated nephropathy								
Chronic inflammatory demyelinating polyr	neuropathy (CIDP)							
Yes No Has the patient re	sponded to previous intravenous imi	mune globulin (IVIG) therapy?						
Churg-Strauss Syndrome (CSS) (allergic								
	as adjunctive therapy for persons w							
	entions been unsuccessful, become							
	ch applies: 🗌 Unsuccessful 📋 Inte	blerable  Contraindicated						
Dermatomyositis	a adjunctive therapy for persons wh	a have had an inadequate reasonable to	first and second line therenics?					
Enteroviral meningoencephalitis	as adjunctive therapy for persons wi	o have had an inadequate response to	first and second line therapies?					
Guillain-Barre Syndrome (GBS) and GBS	variants							
	een diagnosed during the first 2 wee	ks of illness?						
☐ Yes ☐ No Does the patient r								
$\Box$ Yes $\Box$ No Does the patient h	-	<i>, , , , , , , , , ,</i>						
☐ Hematophagocytic lymphohistiocytosis (H	•	rome (MAS)						
Yes No Does the patient h								
	e IgG level: Less than 400mg/dL	☐ 400mg/dl or greater						
	☐ Yes ☐ No Is the IgG level two standard deviations below the mean for age?							
Hemolytic disease of newborn								
	decrease the need for exchange trar	isfusion?						
Yes No Is this request for bacterial control or prevention of infection?								
HIV- associated thrombocytopenia (pedia	tric or adult)							
☐ Hyperimmunoglobulinemia E Syndrome ☐ Yes ☐ No Is this request for treatment of severe eczema?								
Immune or Idiopathic thrombocytopenic p								
		gery, to control excessive bleeding, or	to defer or avoid splenectomy)?					
	irrent platelet count and date collecte	ed:	Date: / /					
🗌 Kawasaki Disease			2001 ,					
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)								
	Parvovirus B19 infection (chronic with severe anemia) Polymyositis in persons who are resistant to first and second line therapies							
Post-transfusion purpura Preparation	for thymoma surgery (to prevent my	asthenia exacerbation) 🔲 Primary h	umoral immunodeficiency diseases:					



## **MEDICARE FORM**

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

Page 3 of 3

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

For Michigan MMP: FAX: <u>1-844-241-2495</u> PHONE: <u>1-855-676-5772 (TTY: 711)</u>

For other lines of business: Please use other form.

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.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB						
G. CLINICAL INFORMATION (continued) –		be completed in its <u>entirety</u> for all	precertification requests.						
• • • •	Please indicate which of the following applies to the patient:								
Congenital agammaglobulinemi X-linked immunodeficiency with Immunodeficiency with thymom Rasmussen encephalitis (Rasmussen's Sy Relapsing-remitting multiple sclerosis (MS		Wiscott- Aldrich Syndrome eficiency None of the Above ted?							
Please select: 🗌 Standard approaches have failed 🔲 Standard approaches have become intolerable 🔲 Standard approaches are contraindicated									
<ul> <li>Renal transplantation from live donor with ABO incompatibility or positive cross-match</li> <li>Yes No Is a suitable non-reactive live or cadaveric donor unavailable (preparative regimen)?</li> <li>Secondary immunosuppression associated with major surgery (such as cardiac transplants) and certain diseases (extensive burns, or collagen-vascular diseases)</li> <li>Selective IgG subclass deficiencies with severe infection for persons meeting selection criteria</li> <li>Solid organ transplantation</li> <li>Yes No Will IVIG be used for allosensitized members undergoing solid organ transplant?</li> <li>Staphylococcal Toxic Shock Syndrome</li> <li>Stem cell or bone marrow transplantation</li> <li>Systemic lupus erythematosus (SLE) (for persons with severe active SLE)</li> <li>Yes No Have other interventions been unsuccessful, become intolerable, or are contraindicated?</li> <li>Please select: Unsuccessful Intolerable Contraindicated</li> <li>Toxic epidermal necrolysis (Lyell's syndrome) and Steven-Johnson Syndrome</li> <li>Toxic shock syndrome or toxic necrotizing fasciitis due to group A streptococcus</li> </ul>									
For Continuation Requests:(Clinical docum			tion of the potiont's progress						
Yes No Has the patient demonstrated an adequate response to therapy? If Yes, please send documentation of the patient's progress (include specific significant or life-threatening infections and dates of occurrences as well as the member's current dosage).									
<ul> <li>☐ Yes ☐ No Has the patient received IVIG within the past 6 months?</li> <li>☐ Yes ☐ No Does the patient have a documented severe and/or potentially life-threatening adverse event that occurred during or following the previous infusion?</li> <li>☐ Yes ☐ No Could the adverse reaction be managed through pre-medication in the home or office setting?</li> </ul>									
H. ACKNOWLEDGEMENT									
Request Completed By (Signature Requ	ired):		Date: //						
Any person who knowingly files a request for insurance company by providing material insurance act, which is a crime and subject	y false information or conceals	material information for the pu							

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