

Intravenous Immune Globulin

HMSA Medicare Advantage- Prior Authorization Request

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. **Please respond below and fax this form to CVS Caremark toll-free at 1-866-237-5512.** If you have questions regarding the prior authorization, please contact CVS Caremark at **1-808-254-4414.** For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

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Patient's Name:			Date:	
Patient's ID:			Patient's Date of Birth:	
Patient's Phone Number:				
Physician's Name:				
Specialty:			NPI#:	
Physician Office Telephone:			Physician Office Fax:	
			a accordance with FDA-approved labeling, lence-based practice guidelines.	
Additional Demographic Info	ormation:			
Patient Weight:		kg		
Patient Height:	ft	inches		
Indicate where the drug is bei				
☐ Military Facility ☐ Sk	illed Nursing Factorial Psychiatric Ropharmacy (1997) Ing administered (1997) Home (1997)	facility \(\begin{align*} \Pi \) Nur esidential Treas Other d: oatient Hospital	tment ☐ End Stage Renal Facility	
What is the requested product? Alyglo Bivigam Gammagard Liquid Gammaked Gamunex-C Panzyga Other	☐ Asceniv☐ Fleboga☐ Gamma☐ Gamma☐ Octagai☐ Privige	amma DIF agard S/D aplex m		
What is the ICD-10 code?				

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Criteria Questions:

100. Is IVIG prescribed for a patient with an ICD-10 code(s) listed in Appendix B: ICD-10 Codes That Support Medical Necessity?
☐ Yes, Continue to #101
□ No, No Further Questions
101. What is the diagnosis? Primary immunodeficiency (includes congenital agammaglobulinemia, X-lined agammaglobulinemia, hypogammaglobulinemia, common variable immunodeficiency, X-linked immunodeficiency with hyper IgM syndrome, severe combined immunodeficiency, and Wiskott-Aldrich syndrome), Continue to #150
Bone marrow/hematopoietic stem cell transplant recipient, <i>Continue to #200</i>
☐ Kidney transplant recipient, Continue to #250
Heart transplant recipient, Continue to #250
☐ Pre-kidney transplantation, No Further Questions
Pre-heart transplantation, No Further Questions
☐ Idiopathic thrombocytopenic purpura (ITP), Continue to #300
Chronic lymphocytic leukemia (CLL), Continue to #400
☐ Multiple myeloma, No Further Questions
☐ Kawasaki disease (mucocutaneous lymph node syndrome), <i>No Further Questions</i> ☐ HIV infection (pediatric), <i>Continue to #500</i>
☐ Guillain-Barre syndrome, <i>Continue to #600</i>
☐ Myasthenia gravis, Continue to #600
☐ Chronic inflammatory demyelinating polyneuropathy (CIDP) and variants excluding MMN, Continue to #800
☐ Multifocal motor neuropathy, <i>Continue to #900</i>
☐ Polymyositis, No Further Questions
☐ Dermatomyositis, <i>Continue to #700</i>
☐ Relapsing-remitting multiple sclerosis, <i>Continue to #700</i>
☐ Lambert-Eaton myasthenic syndrome, <i>No Further Questions</i> ☐ Autoimmune mucocutaneous blistering disease (eg, pemphigus vulgaris, pemphigus foliaceus, bullous pemphigoid), <i>Continue to #1100</i> ☐ Autoimmune retinopathy, <i>Continue to #1200</i>
☐ None of the above, <i>No Further Questions</i>
Primary Immunodeficiency
150. Is this request for a new start or continuation of therapy? ☐ New start, <i>Continue to #152</i> ☐ Continuation of therapy, <i>Continue to #151</i>
151. Was IVIG previously authorized by HMSA/CVS for this member? ☐ Yes, No Further Questions ☐ No, Continue to #152 ☐ Unknown, Continue to #152
152. Does the member have laboratory evidence of immunoglobulin deficiency, defined as any of the following?

TION REQUIRED: Please attach a copy of the laboratory report

Agammaglobulinemia (total IgG less than 200 mg/dL) or

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 Persistent hypogammaglobulinemia (total IgG less than 400 mg/dL, or at least two standard deviations below normal, on at least two occasions) or Absence of B lymphocytes
☐ Yes, Continue to #153
□ No, Continue to #153
153. Does the member have a documented inability to mount an adequate response to inciting antigens, defined as either of the following? <i>ACTION REQUIRED: Please attach a copy of the laboratory report</i> • Lack of appropriate rise in antibody titer following provocation with a polysaccharide antigen. For example, an adequate response to the pneumococcal vaccine may be defined as at least a 4-fold increase in titers for at least 50% of serotypes tested, or • Lack of appropriate rise in antibody titer following provocation with a protein antigen. For example, an adequate response to tetanus/diphtheria vaccine may be defined as less than a 4-fold rise in titers 3-4 weeks after vaccine administration. ☐ Yes ☐ No
Bone Marrow/Stem Cell Transplant Recipient
200. Will IVIG be used for treatment of stem cell transplantation rejection (antibody-mediated)? ☐ Yes, No Further Questions ☐ No, Continue to #201
201. Has the patient received transplantation for a Medicare-approved indication? ☐ Yes, Continue to #202 ☐ No, Continue to #202
202. Is the patient 20 years of age or older? ☐ Yes, Continue to #203 ☐ No, Continue to #203
203. Was the patient cytomegalovirus (CMV) seropositive before transplantation? ☐ Yes, No Further Questions ☐ No, Continue to #204
204. Did the patient undergo allogeneic transplantation for a hematologic malignancy with bone marrow/stem cells from a CMV-seropositive donor? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
Kidney and Heart Transplant
250. Will IVIG be used for treatment of antibody-mediated kidney or heart transplantation rejection? ☐ Yes, No Further Questions ☐ No, No Further Questions
Idiopathic Thrombocytopenic Purpura (ITP)
300. Is IVIG requested for a pregnant woman with idiopathic thrombocytopenic purpura (ITP)? ☐ Yes, Continue to #301 ☐ No, Continue to #306

301. Has the patient previously delivered an infant with autoimmune thrombocytopenia? ☐ Yes, Continue to #304 ☐ No, Continue to #302
302. Does the patient have a platelet count of less than 75,000/mm3 during the current pregnancy? ☐ Yes, Continue to #304 ☐ No, Continue to #303
303. Does the patient have a past history of splenectomy? ☐ Yes, Continue to #304 ☐ No, Continue to #304
304. Has other therapy failed or does the patient have a contraindication to other therapy? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>Continue to #305</i>
305. Does the patient have a rapidly progressive form of the disease? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
306. Does the patient have acute idiopathic thrombocytopenic purpura (ITP)? ☐ Yes, Continue to #307 ☐ No, Continue to #310
307. Will IVIG be used for management of acute bleeding due to severe thrombocytopenia (platelet count less than 30,000/mm3)? ☐ Yes, No Further Questions ☐ No, Continue to #308
308. Will IVIG be used to increase platelet counts prior to an invasive major surgical procedure (eg, splenectomy)? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>Continue to #309</i>
309. Will IVIG be used for a patient with severe thrombocytopenia (platelet count less than 20,000/mm3) considered to be at risk for intracerebral hemorrhage? ☐ Yes, No Further Questions ☐ No, No Further Questions
310. Does the patient have chronic idiopathic thrombocytopenic purpura (ITP)? ☐ Yes, <i>Continue to #311</i> ☐ No, <i>Continue to #311</i>
311. Will IVIG be used for first-line/initial treatment of ITP? ☐ Yes, Continue to #312 ☐ No, Continue to #314
312. Is IVIG requested for a pediatric patient? ☐ Yes, No Further Questions

□ No, Continue to #313
313. Will IVIG be used in combination with corticosteroids (if a rapid response is required or to avoid splenectomy) or does the patient have a contraindication to corticosteroids? Test, No Further Questions No, No Further Questions
314. Has the patient received previous treatment with corticosteroids and has had a splenectomy? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>Continue to #315</i>
315. Are the platelet counts persistently at or below 20,000/mm3? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
Chronic Lymphocytic Leukemia
400. Is IVIG prescribed for a patient with chronic lymphocytic leukemia and associated hypogammaglobulinemia? ☐ Yes, <i>Continue to #401</i> ☐ No, <i>Continue to #401</i>
401. At the time of initiation of IVIG therapy, does/did the patient have repeated bacterial infections? ☐ Yes, <i>Continue to #402</i> ☐ No, <i>Continue to #402</i>
402. At the time of initiation of IVIG therapy, does/did the patient have an IgG level less than 600 mg/dL or is/was there evidence of specific antibody deficiency? ☐ Yes, No Further Questions ☐ No, No Further Questions
Pediatric HIV Infection
500. Is IVIG prescribed for a child less than 13 years of age? ☐ Yes, Continue to #501 ☐ No, Continue to #501
 501. Does the patient meet both of the following criteria? Entry CD4+ lymphocyte counts greater than or equal to 200/mm3 and Clinically symptomatic or asymptomatic, but immunologically abnormal ☐ Yes, No Further Questions ☐ No, No Further Questions
Guillain-Barre Syndrome or Myasthenia Gravis,
600. Has other therapy failed or does the patient have a contraindication to other therapy? ☐ Yes, Continue to #750 ☐ No, Continue to #601
601. Does the patient have a rapidly progressive form of the disease? ☐ Yes, Continue to #750 Send completed form to: CVS Caremark Specialty Programs. Fax: 1-866-237-5512

□ No, Continue to #750
Dermatomyositis or Relapsing-Remitting MS
700. Has other therapy failed or does the patient have a contraindication to other therapy? ☐ Yes, Continue to #750 ☐ No, Continue to #750
Neurological Disorders – Continuation Criteria
750. Is the patient a new start or continuing with IVIG therapy? ☐ New start, <i>Continue to #751</i> ☐ Continuation of therapy, <i>Continue to #752</i>
751. Will the benefit of IVIG therapy be measured using quantitative monitoring tools or any accepted metric such as MRC scale and activities of daily living (ADLs)? The Yes, No Further Questions No, No Further Questions
752. Has the patient demonstrated significant clinical improvement with IVIG therapy? ☐ Yes, <i>Continue to #753</i> ☐ No, <i>Continue to #753</i>
753. What is the total duration of treatment with IVIG? ☐ Less than 2 years, No Further Questions ☐ Greater than or equal to 2 years, Continue to #754
754. Has IVIG dose reduction or withdrawal of treatment been periodically attempted, and the effects measured to validate continued use? ☐ Yes, No Further Questions ☐ No, No Further Questions ☐ Not applicable/condition is not stable, No Further Questions
Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)
800. Does the patient have chronic inflammatory demyelinating polyneuropathy (CIDP) or other neuropathy resulting from diabetes mellitus, dysproteinemia, renal failure or malnutrition? Tyes, Continue to #801 No, Continue to #801
801. Does the patient demonstrate clinical signs and symptoms consistent with a diagnosis of CIDP? Refer to Appendix D Yes, Continue to #802 No, Continue to #802
802. Does electrodiagnostic and laboratory evidence support the diagnosis of CIDP, including the following? Conduction block at sites not prone to compression

Conduction slowing from moderate to severe axonal loss has been excluded

demyelinating polyneuropathy

Motor nerves show segmental conduction slowing and increased distal latencies consistent with a

Cerebrospinal fluid (CSF) analysis shows cytoalbuminologic dissociation (occurs in >90% of cases)

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• Serum tests do NOT support an alternative diagnosis (eg, IgG or IgM monoclonal gammopathy, anti-MAG antibodies, or anti-GM1 antibodies)
☐ Yes, Continue to #803
□ No, Continue to #803
803. Does the patient have any other explanation or any condition below that is associated with CIDP?
• HIV disease
Distally predominant diabetic neuropathy
Diabetic amyotrophy
 Diabetic cachectic neuropathy Distal acquired demyelinating symmetric neuropathy with an IgM paraprotein
☐ Yes, Continue to #804
□ No, Continue to #804
804. Is there evidence of another treatable cause of polyneuropathy or hereditary demyelinating neuropathy?
Yes, Continue to #805
□ No, Continue to #805
805. Does the patient have CIDP that predominantly affects sensory nerves and IVIG is prescribed principally for pain control?
☐ Yes, Continue to #806
□ No, Continue to #808
806. Prior to starting IVIG therapy, did the patient experience a measurable response to a therapeutic trial of prednisone?
☐ Yes, Continue to #807
□ No, Continue to #807
807. Did the patient receive a consultation from a neurologist or rheumatologist who is an expert in the field of CIDP and who validated the need for IVIG to control pain?
☐ Yes, Continue to #809
□ No, Continue to #809
 808. Did the patient receive a consultation from a neurologist or rheumatologist who is an expert in the field of CIDP that resulted in all of the following? Comprehensive history and examination
 Validation of correct CIDP diagnosis Validation of need for IVIG treatment, recommended regimen and appropriate measures of therapeutic benefit, and follow-up
Tyes, Continue to #809
□ No, Continue to #809
809. Has other therapy failed or does the patient have a contraindication to other therapy?
☐ Yes, Continue to #811
□ No, Continue to #810
810. Does the patient have a rapidly progressive form of the disease?
☐ Yes, Continue to #811
□ No, Continue to #811
811. Is the patient a new start or continuing with IVIG therapy for CIDP?

☐ New start, Continue to #812 ☐ Continuation of therapy, Continue to #813
812. Will the benefit of IVIG therapy be measured using quantitative monitoring tools or any accepted metric such as MRC scale and activities of daily living (ADLs)? The Yes, No Further Questions No, No Further Questions
813. Has the patient demonstrated significant clinical improvement with IVIG therapy (and when relevant, a reduction in the level of sensory loss)? ☐ Yes, Continue to #814 ☐ No, Continue to #814
814. What is the total duration of treatment with IVIG for CIDP? ☐ Less than 2 years, <i>No Further Questions</i> ☐ Greater than or equal to 2 years, <i>Continue to #815</i>
815. Has IVIG dose reduction or withdrawal of treatment been periodically attempted, and the effects measured, to validate continued use? Yes, No Further Questions No, No Further Questions Not applicable/condition is not stable, No Further Questions
<u>Multifocal Motor Neuropathy (MMN)</u>
900. Does the patient demonstrate clinical signs and symptoms consistent with a diagnosis of multifocal motor neuropathy (MMN)? Refer to Appendix E Yes, Continue to #901 No, Continue to #901
901. Does electrodiagnostic and laboratory evidence support the diagnosis of MMN? Yes, Continue to #902 No, Continue to #902
 902. Did the patient receive a consultation from a neurologist or rheumatologist who is an expert in the field of MMN that resulted in all of the following? Comprehensive history and examination Validation of correct MMN diagnosis Validation of need for IVIG treatment, recommended regimen and appropriate measures of therapeutic benefit, and follow-up Yes, Continue to #903 No, Continue to #903
903. Is the patient a new start or continuing with IVIG therapy for MMN? ☐ New start, <i>Continue to #904</i> ☐ Continuation of therapy, <i>Continue to #905</i>
904. Will the benefit of IVIG therapy be measured using quantitative monitoring tools or any accepted metric such as MRC scale and activities of daily living (ADLs)? Yes, No Further Questions No, No Further Questions Send completed form to: CVS Caremark Specialty Programs. Fax: 1-866-237-5512

905. Has the patient demonstrated significant clinical improvement with IVIG therapy (and when relevant, a reduction in the level of sensory loss)?
☐ Yes, Continue to #906
□ No, Continue to #906
906. What is the total duration of treatment with IVIG for MMN?
☐ Less than 2 years, No Further Questions
☐ Greater than or equal to 2 years, <i>Continue to #907</i>
907. Has IVIG dose reduction or withdrawal of treatment been periodically attempted, and the effects measured, to validate continued use?
☐ Yes, No Further Questions
□ No, No Further Questions
☐ Not applicable/condition is not stable, <i>No Further Questions</i>
Autoimmune Mucocutaneous Blistering Diseases
1100. Has treatment with conventional therapy failed or is conventional therapy contraindicated?
☐ Yes, Continue to #1102
□ No, Continue to #1101
1101. Does the patient have rapidly progressive disease in which a clinical response could not be affected quickly enough using conventional agents?
☐ Yes, Continue to #1102
□ No, Continue to #1102
1102. Will IVIG be used for short-term therapy (NOT maintenance therapy)?
☐ Yes, No Further Questions
□ No, No Further Questions
Autoimmune Retinopathy
1200. How long has the patient received treatment with IVIG for autoimmune retinopathy?
□ 0 months to up to 3 months, <i>No Further Questions</i>
□ 3 months or more, <i>Continue to #1201</i>
1201. Has the condition improved with IVIG therapy?
☐ Yes, No Further Questions
□ No, No Further Questions

APPENDICES

Appendix A: IVIG CPT/HCPCS Codes

Appendix A.	1710 01 1/1101 00 00dc3
J1459	INJECTION, IMMUNE GLOBULIN (PRIVIGEN), INTRAVENOUS, NONLYOPHILIZED (E.G.
	LIQUID), 500 MG
J1556	INJECTION, IMMUNE GLOBULIN (BIVIGAM), 500 MG
J1557	INJECTION, IMMUNE GLOBULIN, (GAMMAPLEX), INTRAVENOUS, NONLYOPHILIZED
	(E.G. LIQUID), 500 MG
J1561	INJECTION, IMMUNE GLOBULIN, (GAMUNEX-C/ GAMMAKED), NONLYOPHILIZED (E.G.
	LIQUID), 500 MG
J1566	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, LYOPHILIZED (E.G. POWDER), NOT
	OTHERWISE SPECIFIED, 500 MG
J1568	INJECTION, IMMUNE GLOBULIN, (OCTAGAM), INTRAVENOUS, NONLYOPHILIZED (E.G.
	LIQUID), 500 MG
J1569	INJECTION, IMMUNE GLOBULIN, (GAMMAGARD LIQUID), NONLYOPHILIZED, (E.G.
	LIQUID), 500 MG
J1572	INJECTION, IMMUNE GLOBULIN, (FLEBOGAMMA/FLEBOGAMMA DIF), INTRAVENOUS,
	NONLYOPHILIZED (E.G. LIQUID), 500 MG
J1599	INJECTION, IMMUNE GLOBULIN, INTRAVENOUS, NONLYOPHILIZED (E.G. LIQUID),
	NOT OTHERWISE SPECIFIED, 500 MG

Appendix B: ICD-10 Codes That Support Medical Necessity

ICD-10 Code	Description
B20*	Human immunodeficiency virus [HIV] disease
B25.0	Cytomegaloviral pneumonitis
B25.1	Cytomegaloviral hepatitis
B25.2	Cytomegaloviral pancreatitis
B25.8	Other cytomegaloviral diseases
C90.00	Multiple myeloma not having achieved remission
C90.02	Multiple myeloma in relapse
C91.10	Chronic lymphocytic leukemia of B-cell type not having achieved remission
C91.11	Chronic lymphocytic leukemia of B-cell type in remission
C91.12	Chronic lymphocytic leukemia of B-cell type in relapse
D59.0	Drug-induced autoimmune hemolytic anemia
D59.1	Other autoimmune hemolytic anemias
D61.01*	Constitutional (pure) red blood cell aplasia
D69.3	Immune thrombocytopenic purpura
D69.42	Congenital and hereditary thrombocytopenia purpura
D69.49	Other primary thrombocytopenia
D80.0	Hereditary hypogammaglobulinemia
D80.1	Nonfamilial hypogammaglobulinemia
D80.2	Selective deficiency of immunoglobulin A [IgA]
D80.3	Selective deficiency of immunoglobulin G [IgG] subclasses
D80.4	Selective deficiency of immunoglobulin M [IgM]

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ICD-10 Code	Description
D80.5	Immunodeficiency with increased immunoglobulin M [IgM]
D80.6	Antibody deficiency with near-normal immunoglobulins or with hyperimmunoglobulinemia
D80.7	Transient hypogammaglobulinemia of infancy
D81.0	Severe combined immunodeficiency [SCID] with reticular dysgenesis
D81.1	Severe combined immunodeficiency [SCID] with low T- and B-cell numbers
D81.2	Severe combined immunodeficiency [SCID] with low or normal B-cell numbers
D81.5	Purine nucleoside phosphorylase [PNP] deficiency
D81.6	Major histocompatibility complex class I deficiency
D81.7	Major histocompatibility complex class II deficiency
D81.89	Other combined immunodeficiencies
D81.9	Combined immunodeficiency, unspecified
D82.0	Wiskott-Aldrich syndrome
D82.1	Di George's syndrome
D82.4	Hyperimmunoglobulin E [IgE] syndrome
D83.0	Common variable immunodeficiency with predominant abnormalities of B-cell numbers and function
D83.1	Common variable immunodeficiency with predominant immunoregulatory T-cell disorders
D83.2	Common variable immunodeficiency with autoantibodies to B- or T-cells
D83.8	Other common variable immunodeficiencies
D83.9	Common variable immunodeficiency, unspecified
G11.3	Cerebellar ataxia with defective DNA repair
G25.82	Stiff-man syndrome
G35	Multiple sclerosis
G60.3	Idiopathic progressive neuropathy
G61.0	Guillain-Barre syndrome
G61.81*	Chronic inflammatory demyelinating polyneuritis
G61.82	Multifocal motor neuropathy
G65.0	Sequelae of Guillain-Barre syndrome
G70.00	Myasthenia gravis without (acute) exacerbation
G70.01	Myasthenia gravis with (acute) exacerbation
G70.81	Lambert-Eaton syndrome in disease classified elsewhere
G73.1	Lambert-Eaton syndrome in neoplastic disease
G73.3	Myasthenic syndromes in other diseases classified elsewhere
L10.0**	Pemphigus vulgaris
L10.1**	Pemphigus vegetans
L10.2**	Pemphigus foliaceous
L10.3**	Brazilian pemphigus [fogo selvagem]

ICD-10 Code	Description			
L10.4**	Pemphigus erythematosus			
L10.5**	Drug-induced pemphigus			
L10.81**	Paraneoplastic pemphigus			
L10.89**	Other pemphigus			
L10.9**	Pemphigus, unspecified			
L12.0**	Bullous pemphigoid			
L12.1**	Cicatricial pemphigoid			
L12.8**	Other pemphigoid			
L12.9**	Pemphigoid, unspecified			
L13.8**	Other specified bullous disorders			
M30.3	Mucocutaneous lymph node syndrome [Kawasaki]			
M31.1	Thrombotic microangiopathy			
M33.00	Juvenile dermatopolymyositis, organ involvement unspecified			
M33.01	Juvenile dermatopolymyositis with respiratory involvement			
M33.02	Juvenile dermatopolymyositis with myopathy			
M33.09	Juvenile dermatopolymyositis with other organ involvement			
M33.10	Other dermatopolymyositis, organ involvement unspecified			
M33.11	Other dermatopolymyositis with respiratory involvement			
M33.12	Other dermatopolymyositis with myopathy			
M33.19	Other dermatopolymyositis with other organ involvement			
M33.20	Polymyositis, organ involvement unspecified			
M33.21	Polymyositis with respiratory involvement			
M33.22	Polymyositis with myopathy			
M33.29	Polymyositis with other organ involvement			
M33.90	Dermatopolymyositis, unspecified, organ involvement unspecified			
M33.91	Dermatopolymyositis, unspecified with respiratory involvement			
M33.92	Dermatopolymyositis, unspecified with myopathy			
M33.99	Dermatopolymyositis, unspecified with other organ involvement			
M34.83	Systemic sclerosis with polyneuropathy			
M36.0	Dermato(poly)myositis in neoplastic disease			
T86.01	Bone marrow transplant rejection			
T86.02	Bone marrow transplant failure			
T86.09	Other complications of bone marrow transplant			
T86.11	Kidney transplant rejection			
T86.12	Kidney transplant failure			
T86.19	Other complication of kidney transplant			
T86.21	Heart transplantation rejection			
T86.22	Heart transplant failure			

ICD-10 Code	Description			
T86.298	Other complications of heart transplant			
T86.5	Complications of stem cell transplant			
Z48.21	Encounter for aftercare following heart transplant			
Z48.22	Encounter for aftercare following kidney transplant			
Z76.82	Awaiting organ transplant status			
Z86.19	Personal history of other infectious and parasitic diseases			
Z87.01	Personal history of pneumonia (recurrent)			
Z94.0	Kidney transplant status			
Z94.1	Heart transplant status			
Z94.81	Bone marrow transplant status			
Z94.84	Stem cells transplant status			

^{*} Group 1 Medical Necessity ICD-10 Codes Asterisk Explanation:

B20 is only payable for children under 13 years of age.

D61.01 is only to be used when patient has failed all first line therapies.

G61.81 is not payable when associated with diabetes mellitus, dysproteinemias, renal failure, or malnutrition.

Appendix C: Conditions for Which IVIG is NOT Reasonable and Necessary

- Epilepsy
- Amyotrophic lateral sclerosis (ALS)
- Paraneoplastic neurological syndromes
- Undiagnosed neuropathy or weakness
- Malignancies with no causal link to coexisting neurological dysfunctions

Appendix D: Clinical Signs and Symptoms of Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)

- Typical CIDP
 - Symmetrical muscle weakness affects proximal and distal muscles of all four limbs.
 - Sensory loss may affect the distal limbs and usually involves large fiber modalities.
 - The clinical evolution tends to be gradually progressive, evolving over periods of more than 8
 weeks although patients typically present to clinicians within 6 months of onset.
 - Decreased or absent reflexes in affected nerve distributions occur in nearly all CIDP presentations, and develop during the acute phase typically within 8 weeks of symptom onset.
 - The patient should have a neurologic function assessment score of at least 3 or greater on the Rankin Scale at the time of initial therapy. However, IVIg can be used in patients with rapidly worsening weakness regardless of the Rankin score.
- A multifocal variant of CIDP (multifocal acquired demyelinating sensory and motor neuropathy or MADSAM)
 - Leads to sensory and motor dysfunction in multiple individual nerve distributions (for example, ulnar or median).
 - Weakness may affect the upper or lower limbs, but it most commonly affects distal musculature and is more common in the hands.
 - Progression tends to be step-wise with episodes of weakness compiling over time to cause gradually increasing debility.
- Predominant sensory CIDP
 - Occasionally, a patient with CIDP may have only sensory symptoms. The sensory loss may affect the upper or lower limbs and tends to be relatively symmetrical.
 - Like more common sensory-motor CIDP presentations, patients typically seek medical attention within 6-9 months from onset.

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^{**} Refer to Local Coverage Article for ICD-10 codes (A54641)

- The sensory loss may begin relatively acutely and progresses in a stepwise or gradual fashion.
 The sensory distribution is usually not simply limited to the feet or in a stocking distribution, but takes on unusual patterns involving the trunk, arms, or proximal legs.
- The condition is rare compared with the relatively common purely sensory neuropathies such as distal diabetic, toxic, alcoholic, and idiopathic neuropathies.
- Pure sensory CIDP also must be distinguished from distal demyelinating neuropathies associated with an IgM paraprotein, which is not responsive to IVIg or prednisone.

Appendix E: Clinical Signs and Symptoms of Multifocal Motor Neuropathy (MMN)

- MMN is a purely motor syndrome that tends to affect the hands.
- Weakness affects the distribution of individual nerves and tends to progress in a step-wise fashion over time.
- Patients may have subjective sensory complaints but objective sensory findings are not present.
- The diagnosis is generally made using motor and sensory nerve conduction studies. MMN responds to IVIg but not to prednisone. Therefore, prednisone is never indicated in this condition.

I attest that this informa	tion is accura	te and true,	and that docum	entation suppo	rting this
information is available	for review if i	requested by	CVS Caremark	or the benefit	plan sponsor.

X	
Prescriber or Authorized Signature	Date (mm/dd/yy)