

Cutaquig

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 1-855-330-1720**. If you have questions regarding the prior authorization, please contact CVS Caremark at **1-888-877-0518**. 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:	
Specialty:	
Physician Office Telephone:	
<u>Referring</u> Provider Info:	8
Fax:	Phone:
	ing Provider 🛛 Same as Requesting Provider
Name:	NPI#:
Fax:	Phone:

Approvals may be subject to dosing limits in accordance with FDA-approved labeling, accepted compendia, and/or evidence-based practice guidelines.

Required Demographic Information:

Patient Weight: _____kg

Patient Height: _____cm

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Criteria Questions:

- 1. What is the diagnosis? Myasthenia gravis □ Macrophage Activation Syndrome (MAS) Chronic inflammatory demyelinating polyneuropathy (CIDP) □ Hyperimmunoglobulinemia E syndrome □ Immune thrombocytopenic purpura (ITP) □ Multiple myeloma B-cell chronic lymphocytic leukemia (CLL) Opsoclonus-myoclonus □ Stiff-person syndrome □ Post-transfusion purpura Dermatomyositis □ Solid organ transplantation Polymyositis □ Stevens-Johnson syndrome Multifocal motor neuropathy □ Toxic necrotizing fasciitis Human immunodeficiency virus (HIV) infection Toxic epidermal necrolysis Guillain-Barré syndrome □ Toxic shock syndrome Lambert-Eaton myasthenic syndrome □ Kawasaki syndrome (pediatric) □ Parvovirus B19-induced pure red cell aplasia □ Isoimmune hemolytic disease of newborn Neonatal hemochromatosis □ Fetal/neonatal alloimmune thrombocytopenia □ Immune checkpoint inhibitor related toxicity □ Acquired red cell aplasia □ CAR-T therapy related hypogammaglobulinemia □ Acute disseminated encephalomyelitis Rasmussen encephalitis Autoimmune neutropenia □ Enteroviral meningoencephalitis Autoimmune hemolytic anemia Systemic lupus erythematosus Autoimmune neutropenia Hematophagocytic lymphohistiocytosis (HLH) BK virus associated nephropathy □ Major surgery associated secondary immunosuppression □ Churg-Strauss syndrome □ Major burns associated secondary immunosuppression Birdshot retinochoroidopathy □ Hematologic malignancy associated secondary immunosuppression
 - Collagen-vascular disease associated secondary immunosuppression
 - Bone marrow transplant/hematopoietic stem cell transplant recipient

□ Autoimmune mucocutaneous blistering disease (includes pemphigus vulgaris, pemphigus foliaceus, bullous pemphigoid, mucous membrane pemphigoid, and epidermolysis bullosa aquisita)

Drimary immunodeficiency (eg, common variable immunodeficiency, X-linked agammaglobulinemia, severe combined immunodeficiency, Wiskott-Aldrich syndrome)

- Dediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS)
- □ Pediatric acute onset neuropsychiatric syndrome (PANS)
- Other
- 2. What is the ICD-10 code?

Complete the following section based on the patient's diagnosis, if applicable.

Section A: Primary Immunodeficiency

- Is this a request for continuation of immune globulin therapy? \Box Yes \Box No If No, skip to #8 3.
- Has the patient experienced a reduction in the frequency of bacterial infections since starting immune globulin 4.
- 5. Does the prescriber measure trough IgG levels at least once per year? □ Yes □ No □ Not applicable to diagnosis If Not applicable to diagnosis, no further questions.
- 6. Is the most recent trough IgG level at or above the lower range of normal for age? ACTION REQUIRED: If Yes, attach a copy of the laboratory report with a recent IgG trough level. If Yes or Not applicable, no further questions. \Box Yes \Box No \Box Not applicable for diagnosis
- 7. Will the prescriber re-evaluate the dose of immune globulin and consider a dose adjustment (when clinically appropriate)? Yes No Not applicable/not clinically appropriate *No further questions*.

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- 8. What is the specific immunodeficiency disorder?
 - □ Severe combined immunodeficiency (SCID), *specify*: ___
 - Congenital agammaglobulinemia (eg, X-linked or autosomal recessive agammaglobulinemia)
 - □ Other non-SCID combined immunodeficiency disorder, *specify*:
 - Common variable immunodeficiency (CVID)
 - □ Hypogammaglobulinemia (unspecified) or other predominant antibody deficiency disorder
 - □ Selective IgA deficiency
 - □ Selective IgM deficiency
 - □ IgG subclass deficiency
 - □ Specific antibody deficiency
 - □ Other immunodeficiency disorder/none of the above, *specify*:
- 9. ACTION REQUIRED: Please indicate and attach a copy of the following pre-treatment laboratory information (where applicable):
 - IgG (total) level: _____ mg/dL • a) Is IgG (total) level within the normal reference range? \Box Yes \Box No b) If No, is the IgG level greater than or equal to (\geq) 2 SD below the mean for age? \Box Yes \Box No
 - IgG subclass levels:
 - a) IgG1 _____ mg/dL; b) IgG2 _____ mg/dL; c) IgG3 _____ mg/dL; d) Other

e) Are the IgG subclass levels within the normal reference range? \Box Yes \Box No f) If No, is the level(s) greater than or equal to (\geq) 2 SD below the mean for age? \Box Yes \Box No

- g) Were IgG subclass levels measured on at least 2 different occasions? \Box Yes \Box No
- IgA level: _____ mg/dL; Is the IgA level within the normal reference range? \Box Yes \Box No
- IgM level: mg/dL; Is the IgM level within the normal reference range? \Box Yes \Box No
- 10. If diagnosis is severe combined immunodeficiency, are maternal T cells present in the circulation?
- 11. If diagnosis is severe combined immunodeficiency, what is the patient's CD3 T cell count? ACTION REQUIRED: Attach a copy of the laboratory report with lymphocyte subset enumeration by flow cytometry. _____ per microliter
- 12. Was the diagnosis confirmed by molecular or genetic testing? ACTION REQUIRED: Please attach a copy of the laboratory report or other medical record that shows the results of molecular/genetic testing. □ Yes □ No □ Not applicable to diagnosis
- 13. If the diagnosis is common variable immunodeficiency, have other causes of immune deficiency been excluded (eg, drugs, infectious disease, malignancy)? 🗆 Yes 📮 No 📮 Not applicable to diagnosis
- 14. Does the patient have a history of recurrent bacterial infections (eg, pneumonia, otitis media, sinusitis, sepsis, gastrointestinal infections)? Yes No
- 15. Was the immune globulin therapy initiated in the hospital setting? \Box Yes \Box No
- 16. Has the patient demonstrated an impaired antibody response to vaccination with a pneumococcal polysaccharide vaccine? ACTION REQUIRED: If Yes, please attach a copy of the laboratory report with post-vaccination *titers.* \Box Yes \Box No \Box Not applicable

Section B: Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)

- 17. Is this a request for continuation of immune globulin therapy? If Yes, skip to $\#21 \square$ Yes \square No
- 18. Is the disease course progressive or relapsing/remitting for 2 months or longer? \Box Yes \Box No
- 19. Does the patient have moderate to severe functional disability? \Box Yes \Box No
- 20. Were electrodiagnostic studies (electromyography [EMG] or nerve conduction studies [NCS]) and the evaluation of cerebrospinal fluid (when available) performed to confirm the diagnosis? ACTION REOUIRED: If 'yes', attach a copy of the EMG or NCS test results and CSF analysis. \Box Yes \Box No No further questions

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- 21. Has the patient demonstrated significant improvement in disability and maintenance of improvement since starting IG therapy? IYes No
- 22. Is IG being used at the lowest effective dose and frequency? \Box Yes \Box No

Section C: Multifocal Motor Neuropathy (MMN)

- 23. Is this a request for continuation of immune globulin therapy? If Yes, skip to #26 🛛 Yes 🖓 No
- 24. Has the patient experienced progressive, multifocal, asymmetrical weakness without objective sensory loss in 2 or more nerves for at least 1 month? \Box Yes \Box No
- 25. Were electrodiagnostic studies (electromyography [EMG] or nerve conduction studies [NCS]) performed to confirm the diagnosis? *ACTION REQUIRED: If 'yes', attach a copy of the EMG or NCS test results.*□ Yes □ No
- 26. Has the patient demonstrated significant improvement in disability and/or maintenance of improvement since starting IG therapy? □ Yes □ No

Section D: Dermatomyositis (DM) or Polymyositis (PM)

- 27. Is this request for continuation of immune globulin therapy? If Yes, skip to #32 🛛 Yes 🖓 No
- 28. Does the patient exhibit any of the following clinical features? Indicate ALL that apply.
 - □ Proximal muscle weakness (upper or lower extremity and trunk)
 - Elevated serum creatine kinase (CK) or aldolase level
 - □ Muscle pain on grasping or spontaneous pain
 - □ Myogenic changes on EMG (short-duration, polyphasic motor unit potentials with spontaneous fibrillation potentials)
 - Desitive anti-Jo-1 (histadyl tRNA synthetase) antibody
 - □ Non-destructive arthritis or arthralgias

□ Systemic inflammatory signs (fever: more than 37°C at axilla, elevated serum CRP level or accelerated ESR of more than 20 mm/h by the Westergren method

□ Pathological findings compatible with inflammatory myositis (inflammatory infiltration of skeletal evidence of active regeneration may be seen)

□ None of the above

- 29. Were electrodiagnostic studies (electromyography [EMG] or nerve conduction studies [NCS]) and the muscle biopsy (when available) performed to confirm the diagnosis? *ACTION REQUIRED: If 'Yes', attach a copy of the EMG or NCS test results.* □ Yes □ No
- 30. Were standard first-line (corticosteroids) and second-line (immunosuppressants) treatments tried but were unsuccessful or not tolerated? *ACTION REQUIRED: If 'Yes', attach supporting chart note(s) describing previous treatments and no further questions.* □ Yes □ No
- 31. Is the patient unable to receive standard first-line and second-line therapy because of a contraindication or other clinical reason? ACTION REQUIRED: If 'Yes', attach supporting chart note(s) describing previous treatments.
 □ Yes □ No No further questions
- 32. Has the patient demonstrated significant improvement in disability and/or maintenance of improvement since starting IG therapy? IYes No

Section E: Parvovirus B19-induced Pure Red Cell Aplasia (PRCA)

- 33. Does the patient have severe, refractory anemia associated with bone marrow suppression? \Box Yes \Box No
- 34. Does the patient have parvovirus B19 viremia? ACTION REQUIRED: If 'yes', attach test result confirming presence of parvovirus B19. □ Yes □ No

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Section F: Myasthenia Gravis

- 35. What is the primary reason IG is being prescribed?
 - □ Refractory myasthenia gravis, *skip to #38*
 - □ Acute exacerbation/crisis
 - □ Worsening weakness, *skip to #37*
 - Dere-operative management (eg, prior to thymectomy), no further questions
 - Other _____
- 36. Does the patient have severe swallowing difficulty and/or respiratory failure? *If Yes, no further questions.* □ Yes □ No
- 37. Does the patient have weakness with an increase in any of the following symptoms: diplopia, ptosis, blurred vision, difficulty speaking (dysarthria), difficulty swallowing (dysphagia), difficulty chewing, impaired respiratory status, fatigue, or limb weakness? □ Yes □ No *No further questions*
- 38. Has the patient tried and failed 2 or more standard therapies (eg, corticosteroids, azathioprine, cyclosporine, mycophenolate mofetil, rituximab)? ACTION REQUIRED: If 'Yes', attach supporting chart note(s) describing previous treatments. □ Yes □ No

Section G: Stiff-Person Syndrome

- 39. Has the diagnosis been confirmed by anti-glutamic acid decarboxylase (GAD) antibody testing? *ACTION REQUIRED: If 'Yes', attach GAD antibody test results.* □ Yes □ No
- 40. Has the patient received first-line treatment with benzodiazepines and/or baclofen and experienced an inadequate response? *ACTION REQUIRED: If 'Yes', attach supporting chart note(s) describing previous treatments.*□ Yes □ No

Section H: Immune Thrombocytopenic Purpura (ITP)

41. Is the patient a pregnant woman? □ Yes □ No If yes, please provide estimated date of delivery and no further questions:_____

- 42. Is the patient an adult with refractory ITP after splenectomy? \Box Yes \Box No If No, skip to #45
- 43. What is the current pre-treatment platelet count? *ACTION REQUIRED: Attach lab report with platelet count.* ______ per mcL *If less than 30,000/mcL, no further questions.*
- 44. Does the patient have significant bleeding symptoms (eg, mucosal bleeding or other moderate to severe bleeding)? □ Yes □ No *No further questions*
- 45. Is the patient at high risk for bleeding or does the patient require a rapid increase in platelets? If yes, please indicate the risk factors for bleeding or reason for a rapid increase in platelets.
 - Undergoing a medical or dental procedure where blood loss is anticipated
 - Comorbidity (eg, peptic ulcer disease or hypertension)
 - □ Mandated anticoagulation therapy
 - □ Profession or lifestyle predisposes the patient to trauma (eg, construction worker, fireman, professional athlete)
 - Other_
 - □ No, not at high risk or does not require rapid increase in platelets
- 46. What is the current pre-treatment platelet count? *ACTION REQUIRED: Attach lab report with platelet count.* _____mcL
- 47. Does the patient have significant bleeding symptoms (eg, mucosal bleeding or other moderate to severe bleeding)? □ Yes □ No
- 48. What is the classification of ITP?
 - □ Newly-diagnosed ITP (diagnosed within the past 3 months), no further questions if patient is < 18 years old
 - □ Previously untreated ITP (initial therapy), no further questions if patient is less than 18 years old
 - Chronic or persistent ITP (greater than or equal to 3 months from diagnosis), *skip to #51*
 - □ ITP unresponsive to first-line treatment, *skip to #51*
 - Other ____

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- 49. Please indicate the prescribed regimen.
 - □ IG monotherapy
 - □ IG in combination with corticosteroid, *no further questions*
 - Other _____
- 50. Is corticosteroid therapy contraindicated? Yes No *No further questions*
- 51. Does the patient have relapsed ITP after a previous response to IG therapy? *If Yes, no further questions.* □ Yes □ No
- 52. Does the patient have a history of inadequate response, intolerance or a contraindication to corticosteroid or anti-D therapy? *ACTION REQUIRED: If 'Yes', attach supporting chart note(s) describing previous treatments or contraindication*. □ Yes □ No

Section I: B-Cell Chronic Lymphocytic Leukemia (CLL), Bone Marrow Transplant/Hematopoietic Stem Cell Transplant Recipient

- 53. Is this request for continuation of immune globulin therapy? If Yes, skip to #58 🛛 Yes 🖓 No
- 54. Is IG prescribed for prophylaxis of bacterial infections? \Box Yes \Box No
- 55. What is the patient's pre-treatment IgG level? ACTION REQUIRED: If IgG is less than 500 mg/dL, attach a copy of the laboratory report with the pre-treatment IgG level. _____ mg/dL
- 56. *If diagnosis is B-cell chronic lymphocytic leukemia*, does the patient have a history of recurrent sinopulmonary infections requiring intravenous antibiotics or hospitalization? □ Yes □ No *No further questions*
- 57. *If diagnosis is bone marrow transplant/hematopoietic stem cell transplant recipient,* has the patient received a bone marrow/hematopoietic stem cell transplant within the past 100 days? Wes No *No further questions*
- 58. Has the patient experienced a reduction in the frequency of bacterial infections since starting IG therapy? □ Yes □ No

Section J: HIV Infection: Prophylaxis or Thrombocytopenia

- 59. Is the requested drug being prescribed for prophylaxis of bacterial infections in a pediatric patient? *If Yes, skip to #70* □ Yes □ No
- 60. Is the requested drug being prescribed for treatment of thrombocytopenia associated with HIV? 🗖 Yes 🗖 No
- 61. Is the patient an adult? Yes No If No, skip to #66
- 62. Does the patient have significant bleeding? \Box Yes \Box No
- 63. What is the patient's platelet count? _____ / mcL
- 64. Is the patient Rh-positive? \Box Yes \Box No If No, no further questions.
- 65. Has the patient failed treatment with RhIG? Yes No *No further questions*
- 66. What is the patient's pre-treatment IgG level? ACTION REQUIRED: If IgG is less than 400 mg/dL, attach a copy of the laboratory report with the pre-treatment IgG level. _____ mg/dL
- 67. Has the patient had 2 or more bacterial infections in a 1-year period despite antibiotic chemoprophylaxis with TMP-SMZ or another active agent? *If Yes, no further questions.* □ Yes □ No
- 68. Does the patient have HIV-associated thrombocytopenia despite anti-retroviral therapy? *If Yes, no further questions.* □ Yes □ No
- 69. What is the patient's T4 cell count? _____ / mm3 □ Unknown If greater than or equal to 200/mm3, no further questions. If less than 200/mm3 or unknown, skip to #76
- 70. Is this request for continuation of immune globulin therapy? If Yes, skip to $\#79 \square$ Yes \square No

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- 71. Please indicate whether IG will be used for primary or secondary prophylaxis. □ Primary prophylaxis □ Secondary prophylaxis, *skip to #73* □ Other _____, *skip to #74*
- 72. What is the patient's pre-treatment IgG level? ACTION REQUIRED: If IgG is less than 400 mg/dL, attach a copy of the laboratory report with the pre-treatment IgG level. _____ mg/dL
 If less than 400 mg/dL, no further questions.
 If greater than or equal to 400 mg/dL, skip to #74
- 73. Does the patient have a history of recurrent bacterial infections (greater than 2 serious bacterial infections in a 1-year period)? *If Yes, no further questions.* □ Yes □ No
- 74. Has the patient failed to form antibodies to common antigens, such as measles, pneumococcal, and/or Haemophilus influenzae type b vaccine? *If Yes, no further questions.* □ Yes □ No
- 75. Is this request for a single dose of immune globulin for a patient who has been exposed to measles? *If Yes, no further questions.* □ Yes □ No
- 76. Does the patient live in an area where measles is highly prevalent? \Box Yes \Box No If No, skip to #78
- 77. Has the patient failed to develop an antibody response after two doses of measles, mumps, and rubella live virus vaccine? *If Yes, no further questions.* □ Yes □ No
- 78. Does the patient have chronic bronchiectasis that is suboptimally responsive to antimicrobial and pulmonary therapy? □ Yes □ No *No further questions*
- 79. Has the patient experienced a reduction in the frequency of bacterial infections since starting IG therapy? □ Yes □ No

Section K: Lambert-Eaton Myasthenic Syndrome

- 80. Is this request for continuation of immune globulin therapy? If Yes, skip to #86 🛛 Yes 🗋 No
- 81. Has the diagnosis been confirmed by neurophysiology studies (e.g., electromyography) or a positive anti- P/Q type voltage-gated calcium channel antibody test? ACTION REQUIRED: If 'yes', attach a copy of the laboratory report, neurophysiology study report or other supporting medical record(s).
 2 Van Neurophysiology study report of other supporting medical record(s).

Yes – Neurophysiology studies

□ Yes – Positive anti- P/Q type voltage-gated calcium channel antibody test

🗆 No

- 82. Has the patient tried an anticholinesterase (e.g., pyridostigmine) but it was unsuccessful or not tolerated? □ Yes □ No
- 83. Has the patient tried amifampridine (eg 3,4-diaminopyridine phosphate, Firdapse) but it was unsuccessful or not tolerated? □ Yes □ No
- 84. Does the patient have severe weakness? If Yes, no further questions. \Box Yes \Box No
- 85. Is there difficulty with venous access for plasmapheresis? \Box Yes \Box No No further questions
- 86. Has the patient experienced stability or improvement in symptoms relative to the natural course of LEMS? □ Yes □ No

Section L: Immune Checkpoint Inhibitor-Related Adverse Events

- 87. Has the patient experienced a moderate or severe adverse event to a PD-1 inhibitor (e.g., pembrolizumab, nivolumab) or PD-L1 inhibitor (e.g., atezolizumab, avelumab, durvalumab)?
- 88. Is the offending drug being temporarily held or has it been discontinued permanently? \Box Yes \Box No
- 89. Which of the following adverse events did the patient experience?
 - □ Pneumonitis □ Peripheral neuropathy

□ Myasthenia gravis □ Encephalitis

- Transverse myelitis
- Severe inflammatory arthritis

Other _____

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Section M: Hypogammaglobulinemia from CAR-T Therapy

- 90. Has the patient received treatment with CAR-T therapy (e.g., tisagenlecleucel [Kymriah] or axicabtagene ciloleucel [Yescarta]? Yes No
- 91. What is the patient's IgG level? ACTION REQUIRED: If IgG is less than 400 mg/dL', attach a copy of the laboratory report with the pre-treatment IgG level. _____mg/dL □ Unknown

Section N: Guillain-Barre Syndrome (GBS)

- 92. Does the patient have severe disease with significant weakness (e.g., inability to stand or walk without aid, respiratory weakness)? □ Yes □ No
- 93. Did the onset of neurologic symptoms occur less than 4 weeks from the anticipated start of immunoglobulin therapy? Yes No

Section O: Acute Disseminated Encephalomyelitis

94. Has the patient had an insufficient response to intravenous corticosteroid treatment? \Box Yes \Box No

Section P: Autoimmune Mucocutaneous Blistering Disease (includes Pemphigus Vulgaris, Pemphigus Foliaceus, Bullous Pemphigoid, Mucous Membrane Pemphigoid, and Epidermolysis Bullosa Aquisita)

- 95. Has the diagnosis been proven by biopsy and confirmed by pathology report? Yes No
- 96. Is the condition rapidly progressing, extensive, or debilitating? \Box Yes \Box No
- 97. Has the patient failed or experienced significant complications (eg diabetes, steroid-induced osteoporosis) from standard treatment (corticosteroids, immunosuppressive agents)? UYes No

Section Q: Autoimmune Hemolytic Anemia

- 98. Which type of autoimmune hemolytic anemia does the patient have? □ Warm type □ Cold type □ Other _____
- 99. Has the patient tried corticosteroids with inadequate response? If Yes, no further questions. \Box Yes \Box No
- 100. Has the patient has a splenectomy with inadequate response? If Yes, no further questions. \Box Yes \Box No
- 101.Does the patient have a contraindication to corticosteroids or splenectomy? \Box Yes \Box No

Section R: Autoimmune Neutropenia

102.Is treatment with G-CSF (granulocyte colony stimulating factor) an appropriate option? Examples of G-CSF include Fulphila, Granix, Leukine, Neulasta, Neuopogen, Udenyca, Zarxio. Yes No

Section S: Birdshot Retinochoroidopathy

103.Has the patient tried immunosuppressant therapy (e.g., corticosteroids, cyclosporine) with inadequate response? □ Yes □ No

Section T: Churg-Strauss Syndrome

104. Does the patient have severe, active disease? \Box Yes \Box No

105. Will immune globulin be used as adjunctive therapy? \Box Yes \Box No

106. Has the patient experienced failure, intolerance, or is contraindicated to other interventions? \Box Yes \Box No

Section V: Hematophagocytic Lymphohistiocytosis (HLH) and Macrophage Activation Syndrome (MAS) 108. What is the patient's total IgG level? ACTION REQUIRED: Attach a copy of the laboratory report with the pretreatment IgG level._____ mg/dL If less than 400 mg/dL, no further questions.

109.Is the total IgG level at least two standard deviations below the mean for age? \Box Yes \Box No

Section W: Hyperimmunoglobulinemia E syndrome

110. Does the patient have severe eczema? \Box Yes \Box No

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Section X: Multiple Myeloma

111.Does the patient have recurrent, serious infections despite the use of prophylactic antibiotics? 🛛 Yes 🖓 No

Section Y: Neonatal Hemochromatosis 112. Is the patient currently pregnant? Yes No

113.Does the patient have a history of pregnancy ending in documented neonatal hemochromatosis? \Box Yes \Box No

Section Z: Opsoclonus-Myoclonus

- 114.Does the patient have paraneoplastic opsoclonus-myoclonus-ataxia associated with neuroblastoma? *If Yes, no further questions.* \Box Yes \Box No
- 115.Does the patient have refractory opsoclonus-myoclonus? Yes No
- 116.Is immune globulin being used as last-resort treatment? \Box Yes \Box No

Section AA: Rasmussen Encephalitis

117.Did the patient try anti-epileptic drugs with no improvement in symptoms? \Box Yes \Box No

118.Did the patient try corticosteroids with no improvement in symptoms? \Box Yes \Box No

Section BB: Solid Organ Transplantation

- 119. Is immune globulin being prescribed for solid organ transplantation in an allosensitized patient? *If Yes, no further questions.* □ Yes □ No
- 120.Is the patient undergoing renal transplantation from a live donor with ABO incompatibility or positive cross match? Yes No

Section CC: Secondary Immunosuppression Due to Surgery, Malignancy, Burns, Collagen-Vascular Diseases 121.Is immune globulin being requested to prevent or modify recurrent bacterial or viral infections? Yes No

122. What is the patient's pre-treatment IgG level? ACTION REQUIRED: If IgG is less than 400 mg/dL, attach a copy of the laboratory report with the pre-treatment IgG level. _____ mg/dL uket Unknown

Section DD: Toxic Epidermal Necrolysis, Stevens-Johnson Syndrome 123. Is the patient's case severe? Yes No

Section EE: Systemic Lupus Erythematosus

124. Does the patient have severe, active disease? \Box Yes \Box No

- 125.Has the patient experienced inadequate response, intolerance, or have a contraindication to first line therapy? □ Yes □ No
- 126.Has the patient experienced inadequate response, intolerance, or have a contraindication to second line therapy? Yes \Box No

Section FF: Toxic Necrotizing Fasciitis

127. Does the patient have toxic necrotizing fasciitis due to invasive group A streptococcal infection?
 ACTION REQUIRED: If 'yes', attach documentation confirming presence of fasciitis and culture or Gram stain. □ Yes □ No

Section GG: Toxic Shock Syndrome

128.Does the patient have toxic shock syndrome due to a staphylococcal or streptococcal infection? *ACTION REQUIRED: If 'yes', attach culture or Gram stain.* □ Yes □ No

129. Is the infection refractory to several hours of aggressive therapy? If Yes, no further questions. \Box Yes \Box No

130. Does the patient have an undrainable focus of infection? If Yes, no further questions. \Box Yes \Box No

131.Does the patient have persistent oliguria with pulmonary edema? \Box Yes \Box No

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Section HH: PANDAS

- 132. Is the requested drug being used to treat either of the following conditions? \Box Yes \Box No
 - A) pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections, or
 - B) pediatric acute onset neuropsychiatric syndrome

I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.

Х

Prescriber or Authorized Signature

Date (mm/dd/yy)

Send completed form to: Case Review Unit CVS Caremark Specialty Programs Fax: 1-855-330-1720 Note: This fax may contain medical information that is privileged and confidential and is solely for the use of individuals named above. If you are not the intended recipient you hereby are advised that any dissemination, distribution, or copying of this communication is prohibited. If you have received the fax in error, please immediately notify the sender by telephone and destroy the original fax message. Cutaquig PANDAS CFT - 10/2020.

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