

Subject: Therapeutic Apheresis

Guideline #: CG-MED-68

Status: Revised

Publish Date: 10/01/2025

Last Review Date: 02/20/2025

Description

This document addresses therapeutic apheresis, a procedure by which blood is removed from the body, separated into components, manipulated and returned to the individual. There are multiple pheresis procedures that are performed. The therapeutic apheresis procedures addressed in this document utilize devices approved by the United States (U.S.) Food & Drug Administration (FDA) and include the following subcategories:

- Cytapheresis (specifically, erythrocytapheresis, leukocytapheresis, platelet apheresis, red blood cell (RBC) exchange and thrombocytapheresis),
- Immunoadsorption (IA),
- Low-density lipid (LDL) apheresis,
- Plasmapheresis/plasma exchange,
- Selective high-density lipid (HDL) delipidation and therapeutic apheresis.

Notes:

- For criteria related to autologous cellular immunotherapy or ex vivo gene therapy, refer to applicable documents used by the plan.
- For criteria related to apheresis following mobilization for the purpose of acquiring cells to be used when conducted as part of the development of an ex vivo gene therapy for the treatment of a genetic disease, see
 - TRANS.00029 Hematopoietic Stem Cell Transplantation for Genetic Diseases and Aplastic Anemias.

Note: Please see the following documents for additional information:

- MED.00140 Gene Therapy for Beta Thalassemia
- MED.00142 Gene Therapy for Cerebral Adrenoleukodystrophy
- MED.00146 Gene Therapy for Sickle Cell Disease
- MED.00148 Gene Therapy for Metachromatic Leukodystrophy

Clinical Indications

Medically Necessary:

- I. Plasmapheresis or plasma exchange is considered **medically necessary** for any of the following conditions listed in alphabetical order below:

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

- A. Acute inflammatory demyelinating polyradiculoneuropathy / Guillain-Barre syndrome
- B. Anti-glomerular basement membrane disease (Goodpasture's syndrome) when the individual is dialysis-independent or there is evidence of diffuse alveolar hemorrhage (DAH)
- C. Atypical hemolytic uremic syndrome (aHUS) with Factor H autoantibodies or complement gene mutation
- D. Autoimmune hemolytic uremic syndrome- severe cold agglutin disease when there has been an inadequate response to or failure of medical therapy
- E. Catastrophic antiphospholipid syndrome (CAPS)
- F. Chronic inflammatory demyelinating polyneuropathy when **all** of the following criteria are met:
 1. There is muscle weakness or sensory dysfunction caused by neuropathy in more than one limb; **and**
 2. Nerve conduction studies (NCS) or diagnostic criteria confirm evidence of a demyelinating neuropathy; **and**
 3. Other polyneuropathies such as IgM neuropathy, hereditary neuropathy, and diabetic neuropathy have been ruled out
- G. Erythropoietic protoporphyria, liver disease
- H. Cryoglobulinemia, symptomatic/severe, when there has been an inadequate response to or failure of medical therapy
- I. Hyperviscosity syndromes associated with monoclonal gammopathies (such as, multiple myeloma and Waldenström's macroglobulinemia)
- J. Multiple myeloma cast nephropathy (acute renal failure secondary to multiple myeloma) when there has been an inadequate response to or failure of medical therapy
- K. Multiple sclerosis (MS)-acute attack/ relapse when there has been an inadequate response to or failure of medical therapy
- L. Myasthenia gravis, acute short-term treatment
- M. *N*-methyl D-aspartate receptor antibody encephalitis
- N. Neuromyelitis optica spectrum disorders, acute disease/relapse (excluding maintenance therapy) when there has been an inadequate response to or failure of medical therapy
- O. Paraproteinemic demyelinating neuropathies/ chronic acquired demyelinating polyneuropathies; associated with IgA, IgG or IgM monoclonal gammopathy of undetermined significance (MGUS) (excluding multiple myeloma, anti-myelin-associated glycoprotein (MAG) neuropathy or multifocal motor neuropathy (MMN))
- P. Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infection (PANDAS) exacerbation when there has been an inadequate response to or failure of medical therapy
- Q. Systemic lupus erythematosus - for individuals with severe or life-threatening symptoms when conventional therapy has failed to prevent clinical deterioration
- R. Thrombotic microangiopathy secondary to ticlopidine or malignancy
- S. Thrombotic Thrombocytopenic Purpura (TTP)
- T. Transplantation when any of the following are met:
 1. Hematopoietic stem cell transplant
 - a. ABO incompatible; **or**
 - b. Human leukocyte antigen (HLA) incompatibility with haplo-type transplant
 2. Solid organ transplantation for any of the following:
 - a. Heart transplantation recipients who are in the operating room experiencing a hyper-acute rejection episode; **or**

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

- b. Heart transplantation, desensitization/rejection prophylaxis; **or**
 - c. Liver transplantation from a live donor with ABO incompatibility; **or**
 - d. Renal transplant recipients who are post-transplant and experiencing recurrent focal and segmental glomerulosclerosis (FGS) or experiencing a humoral or antibody mediated rejection; **or**
 - e. Renal transplantation in highly sensitive kidney transplant candidates (high PRA protocols) to reduce the number of antibodies reactive against human lymphocyte antigens (HLA); **or**
 - f. Renal transplantation, from a live donor with ABO incompatibility or positive cross-match, where a non-reactive live or cadaveric donor is unavailable; **or**
 - U. Vasculitis anti-neutrophil cytoplasmic antibodies (ANCA) including granulomatosis with polyangiitis, microscopic polyangiitis and renal limited vasculitis, which is associated with rapidly progressive glomerulonephritis and a creatinine level of 5.7 mg/dl or greater, or diffuse alveolar hemorrhage
 - V. Voltage gated potassium channel antibodies when there has been an inadequate response to or failure of medical therapy
 - W. Wilson's disease – fulminant
- II. Low-density lipid (LDL) apheresis or lipoprotein apheresis is considered **medically necessary** for individuals with any of the following:
- A. Homozygous familial hypercholesterolemia;
 - B. Severe, refractory heterozygous familial hypercholesterolemia who have failed a 6-month trial of diet therapy and maximum tolerated combination drug therapy AND who meet either of the following FDA-approved indications:
 - 1. Functional hypercholesterolemic heterozygotes with low-density lipoprotein (LDL) that is greater than 300 mg/dL; **or**
 - 2. Functional hypercholesterolemic heterozygotes with LDL that is greater than 200 mg/dL and documented coronary artery disease
 - C. Lipoprotein (a) hyperlipoproteinemia when there has been an inadequate response to or failure of medical therapy
- III. Cytapheresis:
- A. Erythrocytapheresis (or phlebotomy) is considered **medically necessary** for any of the following indications:
 - 1. Hereditary hemochromatosis; **or**
 - 2. Polycythemia vera; **or**
 - 3. Porphyria cutanea tarda; **or**
 - 4. When used as part of an exchange transfusion process for individuals who are candidates for an ex-vivo gene therapy (for example, lovoitibeglogene autotemcel).
 - B. Leukocytapheresis is considered **medically necessary** for symptomatic hyperleukocytosis
 - C. Thrombocytapheresis is considered **medically necessary** for essential thrombocytosis (symptomatic, when the platelet count is greater than 1,000,000/mm³)
 - D. Red blood cell exchange is considered **medically necessary** for any of the following indications:
 - 1. Babesiosis:
 - a. Severe
-

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

- b. In the high-risk population
- 2. Erythropoietic protoporphyria, liver disease
- 3. Sickle cell disease, acute:
 - a. Acute stroke
 - b. Acute severe, chest syndrome
 - c. Pregnancy
- 4. Sickle cell disease, chronic exchange or non-acute:
 - a. Stroke prophylaxis (primary or secondary)
- 5. When used as part of an exchange transfusion process for individuals who are candidates for an ex-vivo gene therapy (for example, exagamglogene autotemcel).

IV. Immunoadsorption is considered **medically necessary** for individuals with any of the following:

- A. Thrombotic Thrombocytopenic Purpura (TTP); **or**
- B. Acute inflammatory demyelinating polyradiculoneuropathy / Guillain-Barre syndrome
- C. Chronic inflammatory demyelinating polyradiculoneuropathy (CIDP) when **all** of the following criteria are met:
 - 1. Muscle weakness or sensory dysfunction caused by neuropathy is present in more than one limb; **and**
 - 2. Evidence of a demyelinating neuropathy is confirmed by nerve conduction studies (NCS) or diagnostic criteria; **and**
 - 3. Other polyneuropathies (for example IgM neuropathy, hereditary neuropathy or diabetic neuropathy) have been ruled out
- D. Myasthenia gravis, acute short-term treatment
- E. N-methyl D-aspartate receptor antibody encephalitis
- F. Renal transplant recipients who are ABO compatible who are experiencing antibody mediated rejection (AMR) or require desensitization (living donor)
- G. Renal transplant recipients who are ABO incompatible and require desensitization (living donor)
- H. Renal transplant recipients who are post-transplant and experiencing recurrent focal segmental glomerulosclerosis (FSGS)

Not Medically Necessary:

- I. Plasmapheresis or plasma exchange is considered **not medically necessary** when the criteria for plasmapheresis or plasma exchange above are not met and for all other indications.
- II. Low-density lipid apheresis or lipoprotein apheresis is considered **not medically necessary** when the criteria for low-density lipid apheresis or lipoprotein apheresis above are not met and for all other indications.
- III. Selective high-density lipid (HDL) delipidation and therapeutic apheresis is considered **not medically necessary** for all indications.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

- IV. Cytopheresis is considered **not medically necessary** when the criteria for cytopheresis above are not met and for all indications.
- V. Thrombocytopheresis is considered **not medically necessary** when the criteria for thrombocytopheresis above are not met and for all other indications.
- VI. Red blood cell exchange is considered **not medically necessary** when the criteria for red blood cell exchange above are not met and for all other indications.
- VII. Adsorptive cytopheresis is considered **not medically necessary** when the criteria for adsorptive cytopheresis above are not met and for all other indications.
- VIII. Immunoadsorption pheresis is considered **not medically necessary** when the criteria for immunoadsorption pheresis above are not met and for all other indications.

Coding

The following codes for treatments and procedures applicable to this guideline are included below for informational purposes. Inclusion or exclusion of a procedure, diagnosis or device code(s) does not constitute or imply member coverage or provider reimbursement policy. Please refer to the member's contract benefits in effect at the time of service to determine coverage or non-coverage of these services as it applies to an individual member.

Plasmapheresis, plasma exchange

When services are Medically Necessary:

CPT

36514 Therapeutic apheresis; for plasma pheresis

ICD-10 Procedure

6A550Z3 Pheresis of plasma, single
6A551Z3 Pheresis of plasma, multiple

ICD-10 Diagnosis

C88.00-C88.01 Waldenström macroglobulinemia
C90.00-C90.02 Multiple myeloma
G61.0 Guillain-Barre syndrome

When services may be Medically Necessary when criteria are met:

For the procedure codes listed above for the following diagnoses.

ICD-10 Diagnosis

B95.0-B95.1 Streptococcus group A/B, as the cause of diseases classified elsewhere [PANDAS]
D47.2 Monoclonal gammopathy

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

D59.12	Cold autoimmune hemolytic anemia
D59.30-D59.39	Hemolytic-uremic syndrome
D68.61	Antiphospholipid syndrome
D89.1	Cryoglobulinemia
D89.89	Other specified disorders involving the immune mechanism, not elsewhere classified [when specified as PANDAS]
E80.0	Hereditary erythropoietic porphyria
E83.01	Wilson's disease
F42.8	Other obsessive-compulsive disorder [specified as PANDAS]
G04.81	Other encephalitis and encephalomyelitis [specified as NMDA receptor antibody encephalitis]
G13.1	Other systemic atrophy primarily affecting central nervous system in neoplastic disease [limbic encephalopathy, voltage gated potassium channel (VGKC) antibody syndrome]
G35.A-G35.D	Multiple sclerosis
G36.0	Neuromyelitis optica [Devic]
G37.0-G37.9	Other demyelinating diseases of central nervous system
G60.8	Other hereditary and idiopathic neuropathies [Morvan's disease, VGKC antibody syndrome]
G61.81-G61.89	Other inflammatory polyneuropathies
G70.00-G70.01	Myasthenia gravis
G71.19	Other specified myotonic disorders [neuromyotonia, VGKC antibody syndrome]
K77	Liver disorders in diseases classified elsewhere
M31.0	Hypersensitivity angitis [Goodpasture's syndrome]
M31.10-M31.19	Thrombotic microangiopathy [includes TTP]
M31.31	Wegener's granulomatosis with renal involvement
M31.7	Microscopic polyangiitis
M32.0-M32.9	Systemic lupus erythematosus (SLE)
N01.0-N01.A	Rapidly progressive nephritic syndrome
N03.0-N03.A	Chronic nephritic syndrome
N04.0-N04.B2	Nephrotic syndrome
N17.0-N17.9	Acute kidney failure
N18.1-N18.9	Chronic kidney disease
N19	Unspecified kidney failure
T86.00-T86.09	Complications of bone marrow transplant
T86.10-T86.19	Complications of kidney transplant
T86.21	Heart transplant rejection
T86.40-T86.49	Complications of liver transplant
T86.5	Complications of stem cell transplant
Z94.0	Kidney transplant status
Z94.1	Heart transplant status
Z94.4	Liver transplant status
Z94.81	Bone marrow transplant status
Z94.84	Stem cells transplant status

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

When services are Not Medically Necessary:

For the procedure and diagnosis codes listed above when criteria are not met or for all other diagnoses not listed.

Low density lipid apheresis, lipoprotein apheresis and immunoabsorption pheresis

When services are Medically Necessary:

CPT

36516 Therapeutic apheresis; with extracorporeal immunoabsorption, selective adsorption or selective filtration and plasma reinfusion

HCPCS

S2120 Low density lipoprotein (LDL) apheresis using heparin-induced extracorporeal LDL precipitation

ICD-10 Diagnosis

E78.010 Homozygous familial hypercholesterolemia [HoFH]

When services may be Medically Necessary when criteria are met:

For the procedure codes listed above for the following diagnoses.

ICD-10 Diagnosis

E78.00 Pure hypercholesterolemia, unspecified
 E78.011-E78.019 Heterozygous familial hypercholesterolemia [HeFH]; familial hypercholesterolemia unspecified
 G04.81 Other encephalitis and encephalomyelitis [specified as NMDA receptor antibody encephalitis]
 G61.0 Guillain-Barre syndrome
 G61.81-G61.9 Other inflammatory polyneuropathies
 G70.00-G70.01 Myasthenia gravis
 M31.10-M31.19 Thrombotic microangiopathy [includes TTP]
 N01.0-N01.A Rapidly progressive nephritic syndrome
 N03.0-N03.A Chronic nephritic syndrome
 N04.0-N04.B2 Nephrotic syndrome
 N17.0-N17.9 Acute kidney failure
 N18.1-N18.9 Chronic kidney disease
 N19 Unspecified kidney failure
 T86.10-T86.19 Complications of kidney transplant
 Z83.42 Family history of familial hypercholesterolemia
 Z94.0 Kidney transplant status

When services are Not Medically Necessary:

For the procedure and diagnosis codes listed above when criteria are not met or for all other diagnoses not listed.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

When services are also Not Medically Necessary:

For the following procedure code; or when the code describes a procedure designated in the Clinical Indications section as not medically necessary.

CPT

0342T Therapeutic apheresis with selective HDL delipidation and plasma reinfusion

ICD-10 Diagnosis

All diagnoses

Cytapheresis

When services may be Medically Necessary when criteria are met:

CPT

36511 Therapeutic apheresis; for white blood cells
 36512 Therapeutic apheresis; for red blood cells [red blood cell exchange]
 36513 Therapeutic apheresis; for platelets

ICD-10 Procedure

6A550Z0 Pheresis of erythrocytes, single
 6A550Z1 Pheresis of leukocytes, single
 6A550Z2 Pheresis of platelets, single
 6A551Z0 Pheresis of erythrocytes, multiple
 6A551Z1 Pheresis of leukocytes, multiple
 6A551Z2 Pheresis of platelets, multiple

ICD-10 Diagnosis

B60.00-B60.09 Babesiosis
 D45 Polycythemia vera
 D47.3 Essential (hemorrhagic) thrombocythemia
 D56.1 Beta thalassemia
 D57.00-D57.819 Sickle-cell disorders
 E80.0 Hereditary erythropoietic porphyria
 E80.1 Porphyria cutanea tarda
 E83.110 Hereditary hemochromatosis
 K77 Liver disorders in diseases classified elsewhere
 O99.011-O99.03 Anemia complicating pregnancy, childbirth and the puerperium

When services are Not Medically Necessary:

For the procedure and diagnosis codes listed above when criteria are not met or for all other diagnoses not listed.

Discussion/General Information

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Description of Pheresis Techniques

The goal of pheresis is the removal of harmful plasma components. Theoretically, decreasing the concentration of the harmful plasma component, will improve the course of the disease. Abnormal components potentially removed with apheresis include toxins, metabolic substances, and plasma components, such as complement or antibodies. Therefore, diseases thought to be caused by these abnormal constituents might best be treated with this form of therapy. Diseases benefiting from these procedures are largely autoimmune or neurological disorders. Pheresis techniques are not intended to be curative treatments for most indications. Rather, they are used to address related symptoms. Depending on the indication, alternative treatments, such as pharmacologic therapy, may be available.

Applications of pheresis can be broadly subdivided into three general categories: acute, self-limited diseases, acute, fulminant exacerbations of chronic diseases, and chronic diseases. In self-limited diseases and acute exacerbations of chronic diseases, therapeutic apheresis is used to acutely lower the circulating pathogenic substance. In chronic diseases, there is ongoing production of pathogenic autoantibodies. Because therapeutic apheresis does not address the underlying pathology, and due to the phenomenon of rebound antibody production, its use in most chronic diseases has been less effective than in acute, self-limiting diseases. For this reason, chronic conditions are not amenable to pheresis treatment. As an example, individuals with chronic progressive or relapsing-remitting multiple sclerosis are unlikely to benefit from pheresis treatment. However, individuals with an acute central nervous system inflammatory demyelinating disease (including the rare proportion of individuals with multiple sclerosis who suffer from this condition) have been found to have a reduction of symptoms.

Pheresis procedures are typically performed in outpatient settings, including blood banks, dialysis centers, hospital clinics and physician's offices. Reinfusion with human plasma may cause anaphylaxis and bleeding complications, and though rare, may require replacement clotting factors. Therefore, pheresis procedures should be performed by appropriately trained clinicians in a setting that can respond to medical emergencies at all times.

Peer-reviewed, published medical literature and medical society guidelines support the clinical effectiveness and safety of therapeutic pheresis modalities for the indications listed in the policy statements. There is evidence for the accepted indications that the use of this procedure can result in an improvement in symptoms, primarily for acute self-limited conditions, and subsequently for an improvement in quality of life. Evidence is limited regarding the role of therapeutic apheresis to remove specific autoantibodies, proteins and complements in the pathogenesis of many other conditions.

The Guidelines on the Use of Therapeutic Apheresis in Clinical Practice (Ninth special issue) were updated by the American Society for Apheresis (ASFA) in 2023. This guideline is typically updated every 3 years with the most recent evidence. Therapeutic apheresis is a general term which includes all apheresis-based procedures used in a therapeutic manner. This involves the individual's blood being passed through an external device which separates blood into components as treatment of a disease. Specific ASFA definitions for the therapeutic modalities addressed in this document include (Connelly-Smith, 2023):

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Erythrocytapheresis: Procedure in which blood of the patient or donor is passed through a medical device which separates RBCs from other components of blood. The RBCs are removed and replaced with crystalloid or colloid solution, when necessary.

Immunoadsorption (IA): A selective method of therapeutic apheresis in which patient plasma, after membrane based or centrifugal separation from blood, is passed through an adsorber column which has a capacity to remove immunoglobulins and Immune complexes by binding them to select ligands (e.g., staphylococcal or recombinant protein A, sheep polyclonal anti-human antibodies, tryptophan, synthetic oligopeptides, monoclonal camel antibody fragments) on the backing matrix surface of membranes or beads.

Leukocytapheresis: A procedure in which blood of the patient or the donor is passed typically through a medical centrifuge which separates out white blood cells (e.g., leukemic blasts or granulocytes), collects the selected cells and returns remainder of the patient's or the donor's blood with or without addition of replacement fluid such as colloid and/or crystalloid solution.

Lipoprotein apheresis (LA): Previously known as low density lipid apheresis. The selective removal of lipoproteins from the blood with the return of the remaining components and without need for replacement fluids such as plasma products, thus avoiding related adverse events.. A variety of methodologies with different physicochemical principles (e.g., filtration, precipitation, or adsorption) are available and include systems treating plasma (e.g., DFPP, HELP-apheresis, polyclonal-sheep-anti-apoB-immunoadsorption, dextran-sulfate plasma adsorption) and whole blood adsorption systems (e.g., using ligands consisting of dextran-sulfate or polyacrylate).

Red blood cell (RBC) Exchange: A therapeutic procedure in which blood of the patient is passed through a medical device which separates RBCs from other components of blood. The patient's RBCs are removed and replaced with donor RBCs.

Therapeutic plasma exchange (TPE): Plasma of the patient is separated from other components of blood, either by membrane filtration (mTPE) or centrifugation (cTPE). The plasma is removed with subsequent substitution of a replacement solution (e.g., human albumin and/or plasma) or a combination of crystalloid/colloid solution.

Thrombocytapheresis: A therapeutic procedure in which blood of the patient is passed through a medical device which separates out platelets, removes the platelets and returns remainder of the patient's blood with or without addition of replacement fluid such as colloid and/or crystalloid solution.

The ASFA utilizes categories and grade of recommendation for various indications. The four categories are (Connelly-Smith, 2023):

Category I	Disorders for which apheresis is accepted as first-line therapy, either as a primary standalone treatment or in conjunction with other modes of treatment.
------------	--

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

Category II	Disorders for which apheresis is accepted as second-line therapy, either as a standalone treatment or in conjunction with other modes of treatment.
Category III	Optimum role of apheresis therapy is not established. Decision making should be individualized.
Category IV	Disorders in which published evidence demonstrates or suggests apheresis to be ineffective or harmful. Institutional Review Board (IRB) /Ethics Committee approval is desirable if apheresis treatment is undertaken in these circumstances.

The Grading Recommendations include the following (Connelly-Smith, 2023):

Recommendation	Description	Methodological quality of supporting evidence
Grade 1A	Strong recommendation, high-quality evidence	Randomized controlled trials (RCTs) without important limitations or overwhelming evidence from observational studies
Grade 1B	Strong recommendation, moderate quality evidence	RCTs with important limitations (inconsistent results, methodological flaws, indirect, or imprecise) or exceptionally strong evidence from observational studies
Grade 1C	Strong recommendation, low-quality or very low-quality evidence	Observational studies or case series
Grade 2A	Weak recommendation, high quality evidence	RCTs without important limitations or overwhelming evidence from observational studies
Grade 2B	Weak recommendation, moderate-quality evidence	RCTs with important limitations (inconsistent results, methodological flaws, indirect, or imprecise) or exceptionally strong evidence from observational studies
Grade 2C	Weak recommendation, low-quality or very low-quality evidence	Observational studies or case series

Plasma Pheresis and Plasma Exchange

The ASFA recommends plasmapheresis or plasma exchange as Category I treatment options and the associated levels of evidence range with Grade 1A – 1C recommendations for the following indications (Connelly-Smith, 2023):

- Acute inflammatory demyelinating polyradiculoneuropathy (Guillain-Barre Syndrome)

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

- Anti-glomerular basement membrane disease (Goodpasture's syndrome) when dialysis-independent or evidence of DAH
- Chronic inflammatory demyelinating polyneuropathy
- Focal segmental glomerulosclerosis that is recurrent in a transplanted kidney
- Hyperviscosity syndromes in hypergammaglobulinemia
- Liver transplantation from a live donor with ABO incompatibility, desensitization
- Myasthenia gravis, acute, short-term treatment
- N-methyl D-aspartate receptor antibody encephalitis
- Chronic acquired demyelinating polyneuropathies; IgG/IgA/IgM
- Renal transplantation, desensitization
- Renal transplant recipients with ABO compatibility who are post transplant and experiencing an antibody mediated rejection
- Thrombotic microangiopathy, thrombotic thrombocytopenic purpura (TTP)
- Wilson disease – fulminant

In addition to the above ASFA recommendations, published peer-reviewed literature and/or specialty medical societies support the use therapeutic apheresis when there is evidence that a targeted component(s) of the blood or plasma is readily removed with therapeutic apheresis and there is sufficient improvement in clinical symptoms without causing serious adverse effects.

The ASFA recommends plasmapheresis or plasma exchange as Category II treatment options and the associated levels of evidence range with Grade 1A – 1C recommendations for the following indications (Balogun, 2020; Connelly-Smith, 2023):

- Cardiac Transplantation, desensitization or rejection prophylaxis
- Familial hypercholesterolemia in Homozygotes/ Heterozygotes
- Hematopoietic stem cell transplant – ABO incompatible when specific to the marrow specific hematopoietic progenitor cell (HPC) indication
- Neuromyelitis optica spectrum disorders (NMOSD)-acute attack/relapse
- Multiple sclerosis –acute attack/relapse
- Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infection (PANDAS) exacerbation
- Renal Transplantation, ABO incompatible, antibody mediated rejection
- Voltage-gated potassium channel (VGKC) antibody related diseases

Acute inflammatory demyelinating polyneuropathy (AIDP)/ Guillain-Barre Syndrome (GBS)

The American Academy of Neurology (AAN) (Cortese, 2011) published an updated evidence-based guideline on plasmapheresis in neurologic disorders. Plasmapheresis was recommended in the treatment of “AIDP/GBS severe enough to impair independent walking or to require mechanical ventilation (Level A). Plasmapheresis should be considered in the treatment of milder clinical presentations (Level B).” In a consensus statement by the American Association of Neuromuscular and Electrodiagnostic Medicine (AANEM) Committee (Donofrio, 2009), treatment

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

of GBS in adults, particularly those who require an aid to walk (disability grade ≥ 2) within 2 weeks of the onset of symptoms with IVIG was recommended. The committee noted based on randomized trials, similar functional improvements in individuals treated with IVIG or plasmapheresis. Data did not support the combination of IVIG and plasmapheresis to treat GBS.

Chronic inflammatory demyelinating polyneuropathy (CIDP)

CIDP is a rare disorder which typically presents with symmetrical motor function impairment that may or may not be coupled with sensory disturbances. CIDP generally progresses over the course of at least 2 months but may follow a relapsing/remitting course. CIDP affects approximately 5-7 cases per 100,000 individuals and over 50% of affected individuals will be unable to walk without support during the course of the disease. Approximately 10% of these individuals will become persistently disabled or will die. Although the exact etiology is unknown, CIDP is thought to be the result of an autoimmune attack (humoral and cell-mediated) on the peripheral nerves which damages the myelin sheath. This myelin damage is documented by nerve conduction studies or nerve biopsies. There appears to be similar efficacy between all first-line therapies: steroids, IVIG, or TPE (Van den Bergh, 2021). CIDP is difficult to diagnose, atypical CIDP can account up to 50% of all cases and several neuropathies share a similar clinical presentation with CIDP. Once the diagnosis is made, prompt treatment is needed to halt inflammatory demyelination, prevent secondary axonal degeneration, and minimize the chances of permanent disability.

Complement-mediated thrombotic microangiopathy (TMA)

TMA, previously known as Atypical Hemolytic Uremic Syndrome (aHUS) is a rare genetic and chronic blood disease that can lead to renal failure and is associated with increased risk of stroke and death. Prompt diagnosis is essential, as aHUS is aggressive and treatment can be initiated for affected individuals. According to Loirat (2011), the diagnosis of aHUS relies on: 1) no associated disease; 2) investigations for Shiga toxin E. coli infection at onset of aHUS with no evidence of a Shiga-toxin/EHEC positive test (stool culture and polymerase chain reaction for Shiga-toxins; serology for anti-lipopolysaccharides antibodies); and 3) ADAMTS 13 (A Disintegrin-like And Metalloprotease with ThromboSpondin type 1 motif 13) determination as manifestations of aHUS and TTP may overlap. Frequent causes of aHUS involve complement regulatory abnormalities such as factor H mutations and MCP mutations. Plasma exchange can replace mutated circulating complement factors, specifically factor H. However, plasma exchange does not demonstrate similar efficacy for individuals with isolated MCP mutations, and plasma exchange is not recommended to treat MCP aHUS. Based on expert consensus opinion, plasma therapy has demonstrated efficacy as the first-line treatment for aHUS and should be started as early as possible, typically within 24 hours of presentation (Ariceta, 2009; Lapeyraque, 2011; Loirat, 2011; Taylor 2010). However, since the introduction of eculizumab, which has been shown to be effective as a front-line therapy, the role of plasma therapy has become more limited.

Autoimmune Hemolytic Uremic Syndrome – Severe Cold Agglutinin Disease

Individuals with autoimmune hemolytic anemia may be classified into warm or cold autoantibody types. Those with cold agglutinin disease (CAD) have IgM autoantibodies that target red blood cells when the temperatures are typically between 0–5 degrees Celsius. Avoiding exposure to cold temperatures is the primary treatment. The

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

ASFA notes that most effective current treatment is rituximab, which is recommended as first-line therapy (Connelly-Smith, 2023).

Catastrophic Antiphospholipid Syndrome (CAPS)

ASFA (Connelly-Smith, 2023) defines CAPS as the acute onset of multiple thromboses in at least three organ systems over a period of days or weeks, in individuals with antiphospholipid antibodies. European Forum on Antiphospholipid Antibodies maintains a registry of individuals with CAPS due to the rarity of the disease. Data from the registry documented higher survival rates related to the increased frequency of combination therapy including anticoagulants, glucocorticoids and plasma exchange and/or intravenous immune globulin (Cervera, 2012). Specialty input recommends use of plasmapheresis or plasma exchange to treat CAPS.

Cryoglobulinemia

Cryoglobulins, single or mixed immunoglobulins may precipitate at low temperatures and can deposit in the vessels and tissues of the body. These cryoglobulins can increase the viscosity of the circulating blood and may cause vasculitis and obstruct vessels causing organ damage. Cryoglobulinemia involves the presence of cryoglobulins in the blood and typically occurs concomitantly with other conditions such as viral hepatitis, lymphoproliferative and autoimmune disorders. Treatment is based on the severity of disease and addressing the underlying disorder. Individuals who are asymptomatic do not require treatment. Mild symptoms can be treated with cold avoidance and analgesics. People with more severe cases are treated with immunosuppressive therapy. The AFSA recommendation for plasmapheresis or plasma exchange in severe or symptomatic is a category II with an evidence level of 2A (Connelly-Smith, 2019).

Hyperviscosity syndromes associated with monoclonal gammopathies

The National Comprehensive Cancer Network® Clinical Practice Guidelines in Oncology (NCCN Guidelines® V1.2025) on multiple myeloma recommend plasmapheresis as an adjunctive therapy for symptomatic hyperviscosity. The guideline also notes, “Institutions differ in their use of plasmapheresis for adjunctive treatment of renal dysfunction.” The recommendation to improve renal function by utilizing plasmapheresis is a category 2B recommendation.

Lipoprotein (a) hyperlipoproteinemia

Lipoprotein (a) is recognized as an independent and causal risk factor for atherosclerotic cardiovascular disease (ASCVD). While its physiological function is still unknown, lipoprotein (a) exerts all the atherogenic effects of an LDL-particle. As elevated lipoprotein (a) levels are prevalent in the population, treatment is limited to those with additional comorbidities. While lipoprotein (a) levels are not affected by changes in diet or lifestyle, some existing lipid lowering drugs can reduce lipoprotein (a) levels.

Myeloma Cast Nephropathy

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

The formation of casts in the distal tubules in the kidneys of some individuals with multiple myeloma is thought to be due to an increase in light chain concentrations typically seen with tumor progression (Connelly-Smith, 2019). Primary therapy typically involves chemotherapy and intravenous fluid to increase and the urine's alkalinity to dissolve the light chains. The ASFA recommendation was a category II with a level of evidence 2B.

Multiple Sclerosis (MS) – Acute CNS inflammatory demyelinating disease

The ASFA recommends the use of plasmapheresis or plasma exchange to treat acute CNS inflammatory demyelinating disease associated with multiple sclerosis (Connelly-Smith, 2019). The AAN has a Level B recommendation for plasmapheresis as adjunctive treatment of exacerbations in relapsing forms of MS. Based on Level A evidence, the AAN states, "Plasmapheresis should not be offered for chronic progressive or secondary progressive MS" (Cortese, 2011).

Neuromyelitis optica spectrum disorders (NMOSD)

NMOSD, also known as Devic's disease, is an inflammatory CNS syndrome, which is distinct from multiple sclerosis. NMOSD targets optic nerves and the spinal cord, causing immune-mediated demyelination and axonal injury. High-dose intravenous methylprednisolone followed by oral dosing is considered first-line therapy. Plasmapheresis therapy is usually limited to those who do not respond adequately to first line therapy (Connelly-Smith, 2023). In addition to methylprednisolone therapy, other pharmacological treatments have shown promise in the treatment of NMOSD (Cree, 2019; Damato, 2016; Pittcock, 2019).

PANDAS

Although ASFA (Connelly-Smith, 2023) provides a Category II recommendation with grade 1B level of evidence for plasmapheresis or plasma exchange as a treatment for PANDAS, there is still conflicting information in the published literature. The AAN guideline on plasmapheresis in neurologic disorders (Cortese, 2011) stated there is "insufficient evidence to support or refute the use of plasmapheresis in the treatment of acute obsessive compulsive disorders (OCD) and tic symptoms in the setting of PANDAS." Swedo and colleagues (2012) proposed a set of diagnostic criteria for Pediatric Acute-onset Neuropsychiatric Syndrome (PANS) as modified from PANDAS criteria. The authors noted this new set of criteria would need to be validated in large trials. In a retrospective review in a large metropolitan area, Gabbay (2008) reported significant over diagnosis of PANDAS and subsequent therapies that included plasma exchange. The AFSA noted that initial treatment includes cognitive behavioral therapy and/or anti-obsessional medications (Connelly-Smith, 2023).

Sydenham's chorea

ASFA (Connelly-Smith, 2023) provides a Category III recommendation grade 2B level of evidence for plasmapheresis or plasma exchange as a treatment for Sydenham's chorea, there is still conflicting information in the published literature. Sydenham's chorea is a pediatric post-infectious autoimmune neuropsychiatric disorder that manifests after an acute bout of rheumatic fever. Symptoms include rapid and jerky, involuntary movements that may affect the face, trunk and extremities, which may prevent independent activities of daily living. Chorea is usually treated with neuroleptics, valproic acid and corticosteroids. Sydenham's chorea is thought to be an

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

autoimmune disorder based on the presence of antibodies that react with neuronal tissue that control motor activity (Garvey, 2005; National Institute of Neurological Disorders and Stroke [NINDS], 2007). The AAN guideline (Cortese, 2011) noted there is “Insufficient evidence to support or refute the use of plasmapheresis in the treatment of Sydenham chorea.” The NINDS notes there is no specific treatment for Sydenham’s chorea.

Systemic Lupus Erythematosus (SLE) – Severe

Circulating autoantibodies and immune complexes along with complement deposition results in injury to cells and tissues of individuals with SLE. Immunosuppressive agents and biologic therapies are used to treat SLE. Specialty input recommends therapeutic plasmapheresis or plasma exchange for those with severe or life-threatening symptoms when conventional therapies have failed to prevent clinical deterioration.

Thrombotic Microangiopathy (TMA)

TMA is a rare medical condition involving small blood vessel damage from blood clots, usually as a result of other complex and serious illnesses. There are various syndromes such as HUS and TTP that involve TMA as a component of the specific syndrome but are classified separately based on additional clinical indications (Loriat, 2011).

Transplantation

Approximately 20% of individuals waiting for cadaveric renal transplantation have high cytotoxic antibody titers rendering them at high risk for hyperacute and acute allograft rejection. Plasma exchange and immunoadsorption have been used to significantly reduce levels of these antibodies prior to transplantation. When used in individuals with recurrent FSGS, pheresis induced a decrease in urinary protein. Additionally, randomized controlled trials have been conducted establishing the efficacy of plasma exchange in the treatment of biopsy proven acute antibody mediated renal allograft rejection. The literature on pheresis for the treatment of chronic rejection is limited to a few uncontrolled studies with modest and transient results.

Voltage-gated potassium channel (VGKC) antibody related diseases

Dysfunction of the VGKC antibodies, which are important in the control of membrane excitability in the nervous system, is associated with a variety of neurological conditions, including limbic encephalitis, acquired neuromyotonia (NMT), LGI1/CASPR2- antibody mediated encephalitis (AME), and Morvan’s syndrome (MVS). Affected individuals present with a variety of symptoms, delaying diagnosis in many individuals. Immunotherapy, including steroids and immunosuppressive drugs, as well as symptomatic treatment, are typical first-line treatments. The AFSA notes that due to a high variability in presentation and treatment response associated with VGKC antibody related disease, individualized treatment is warranted.

Other Indications

Additional ASFA indications for plasma pheresis or plasma exchange were of lower levels of evidence and/or the optimum role of therapy could not be established; or the evidence demonstrated the therapy could be ineffective or

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

harmful. These indications are listed by categories:

Plasmapheresis or plasma exchange treatment options with Category I-II recommendation and levels of evidence ranging from Grade 2A – 2C include the following indications (Connelly-Smith, 2023):

- Acute disseminated encephalomyelitis (ADEM); steroid refractory
- Autoimmune hemolytic anemia, severe cold agglutinin disease
- Catastrophic antiphospholipid syndrome
- Cryoglobulinemia severe/symptomatic
- Erythropoietic protoporphyria, liver disease
- Hemapoietic stem cell transplant, major ABO incompatible, HPC(A)
- Lambert-Eaton myasthenic syndrome
- Myasthenia gravis; long-term management
- Myeloma cast nephropathy
- Overdose of drugs or poisoning – mushroom
- Phytanic acid storage disease (Refsum's disease)
- Steroid-responsive encephalopathy associated with autoimmune thyroiditis (Hashimoto's Encephalopathy)
- Systemic lupus erythematosus (SLE)
- Thrombotic microangiopathy, factor H autoantibody mediated or associated with Ticlopidine
- Thyroid storm
- Vasculitis, Hepatitis B polyarteritis nodosa

Plasmapheresis or plasma exchange treatment options with Category III recommendations and levels of evidence ranging from Grade 1B – 2C were provided for the following indications (Connelly-Smith, 2023):

- Acute liver failure
- Alzheimer's disease
- Anti-glomerular basement membrane disease (Goodpasture's syndrome) when the individual is dialysis-dependent and there is no evidence of DAH
- Atopic dermatitis, recalcitrant
- Autoimmune dysautonomia
- Autoimmune hemolytic anemia, warm (WAHA), severe
- Burn shock resuscitation
- Cardiac neonatal lupus
- Chronic acquired demyelinating polyneuropathies; anti-myelin-associated glycoprotein or = chronic ataxic neuropathy, ophthalmoplegia, immunoglobulin M paraprotein, cold agglutinins and disialosyl antibodies syndrome (CANOMAD) / chronic ataxic neuropathy with disialosyl antibodies syndrome (CANDA)
- Complex regional pain syndrome, chronic
- Dilated cardiomyopathy, idiopathic
- Focal segmental glomerulosclerosis (FSGS); steroid resistant in native kidney

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

- Heart (cardiac) transplantation; antibody mediated rejection (AMR)
- Hematopoietic stem cell transplantation, ABO incompatible with pure red blood cell aplasia or HLA desensitization
- Hemophagocytic lymphohistiocytosis; hemophagocytic syndrome; macrophage activating syndrome
- Heparin induced thrombocytopenia and thrombosis
- Hypertriglyceridemic (HTG) pancreatitis
- Idiopathic inflammatory myopathies
- Immune checkpoint inhibitors, immune-related adverse events
- Immune thrombocytopenia, refractory
- Immunoglobulin A nephropathy
- Intestine transplantation
- Liver transplantation – desensitization from a deceased donor or antibody mediated rejection
- Lung transplantation – AMR or desensitization
- Multiple sclerosis; chronic primary or secondary progressive
- Neuromyelitis optica spectrum disorders (NMOSD); maintenance treatment
- Nephrogenic systemic fibrosis
- Overdose of drugs or poisoning – envenomation
- Paraneoplastic syndromes (autoimmune retinopathies and neurological)
- Pediatric autoimmune neuropsychiatric disorders; severe Sydenham's chorea
- Pemphigus vulgaris, severe
- Post transfusion purpura
- Progressive multifocal leukoencephalopathy (PML); natalizumab associated
- Pruritus related to hepatobiliary disease; treatment resistant
- Psoriasis, disseminated pustular
- Red cell alloimmunization in pregnancy, prevention and treatment
- Systemic sclerosis
- Sepsis with multi-organ failure
- Stiff-person syndrome
- Sudden sensorineural hearing loss
- Thrombotic microangiopathy
 - coagulation mediated
 - complement factor gene mutations mediated
 - clopidogrel associated
 - infection associated
 - pregnancy associated
 - Shiga toxin-producing Escherichia coli hemolytic uremic syndrome (STEC-HUS)
 - transplantation associated
- Toxic epidermal necrolysis, refractory
- Vaccine-induced immune thrombotic thrombocytopenia
- Vasculitis
 - ANCA associated

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

- IgA
- Kawasaki disease
- Multisystem inflammatory syndrome in children

Plasmapheresis or plasma exchange treatment options with Category IV recommendations were provided for the following indications (Connelly-Smith, 2023):

- Psoriasis
- Thrombotic microangiopathy, drug (gemcitabine or quinine) related

Immune thrombocytopenia

Immune thrombocytopenia has a widely accepted abbreviation, ITP, and is also known as “immune thrombocytopenic purpura and idiopathic thrombocytopenic purpura.” The most recent name “immune thrombocytopenia” is based on an international work group to standardize the terminology and definitions of ITP (Neunert, 2019; Rodeghiero, 2009). The 2019 evidence-based practice guideline for ITP does not include any form of therapeutic apheresis as a recommended treatment (Neunert, 2019).

Lupus Nephritis

The ASFA lists the use of plasmapheresis or plasma exchange treatment options with a Category IV (treatment is ineffective and potentially harmful) recommendation with a level of evidence Grade 1B. This recommendation was based on a controlled study that showed no benefit of adding therapeutic plasmapheresis or plasma exchange to prednisone and cyclophosphamide for individuals with severe lupus nephritis, these findings were supported by smaller, later trials. (Schwartz, 2016). While a 2010 RCT suggested that adjunctive IA and TPE were equally effective in reducing systemic lupus erythematosus Disease Activity Index (SLEDAI) scores, the ASFA did not change their recommendation. American College of Rheumatology (ACR) guidelines for screening, treatment and management of lupus nephritis do not include the use of plasmapheresis or plasma exchange (Hahn, 2012). Similarly, the Joint European League Against Rheumatism and European Renal Association-European dialysis and Transplant Association (EULAR/ERA-EDTA) (Bertsias, 2012) do not include the use of plasmapheresis or plasma exchange as a recommended treatment for the management of adult or pediatric lupus nephritis.

Phytanic Acid Storage Disease (Refsum's disease)

Refsum Disease, or phytanic acid storage disease, is a rare genetic disease where individuals lack the enzyme to break down phytanic acid found in certain foods. The excess phytanic acids accumulate in the brain, blood and other tissues, which can cause blindness and arrhythmias in the heart. Avoidance of foods such as dairy products, beef, lamb, and fatty fish (e.g., tuna, cod and haddock) is the primary treatment. When there is excessive buildup of phytanic acid, plasma exchange has been utilized with some improvement in symptoms, but vision and hearing problems persisted (NINDS, 2011).

Stiff Person (Man) Syndrome

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Also known as Stiff-Man Syndrome, the classic features of this syndrome include painful muscular spasms associated with muscular stiffness, often times impairing the ability to walk. This syndrome is a rare, chronic but usually not progressive disorder. Published literature includes case reports with mixed results from the use of plasma exchange to treat this condition (Connelly-Smith, 2023).

Lipoprotein apheresis

Low-density apheresis

Low-density lipoprotein (LDL) apheresis describes a variety of technologies used to acutely remove lipoprotein LDL from the plasma. The individual initially undergoes an apheresis procedure to isolate the plasma. The LDLs are then selectively removed from the plasma by either immunoadsorption, heparin-induced extracorporeal LDL precipitation (also referred to as HELP), or dextran sulfate adsorption. In immunoadsorption, polyclonal antihuman apoB antibodies from sheep selectively bind and remove LDL. (ApoB is the protein moiety of low-density lipoprotein.) In HELP, LDL and other particles containing ApoB are precipitated by heparin at an acidic pH. Dextran sulfate adsorption removes LDL by binding the positively charged apoB to dextran sulfate particles bound to cellulose. LDL apheresis is a selective procedure in which only pathogenic low-density lipoproteins are removed. The plasma is then returned to the individual. Examples of two LDL-apheresis systems currently FDA-approved for use in the United States are the Plasmat[®] Futura Apheresis system (B. Braun Avitum AG., Bethlehem, PA) and the LipoSorber[®] system (Kaneka Medical America LLC, New York, NY).

The ASFA recommends lipoprotein apheresis as Categories I-II treatment options and the associated level of Grade 1A (strong recommendation, high-quality evidence) (Connelly-Smith, 2023). A single session of lipoprotein apheresis has been shown to decrease cholesterol levels by 60-70% (Connelly-Smith, 2023). There are various recommendations for individual selection criteria, however, there is a lack of evidence to support the initiation of treatment at specific LDL levels.

Familial hypercholesterolemia is an autosomal, co-dominant inherited disorder of lipoprotein metabolism that features mutations in the LDL-receptor gene. Due to reduced or absent LDL receptors, low-density lipoprotein cholesterol (LDLc) is not effectively cleared by the liver, resulting in very high plasma concentration in the circulating blood. LDLc deposits can be found in the tendons (tendon xanthomas) and arterial walls and may be associated with an increased risk of early onset of atherosclerosis and premature coronary heart disease (CHD.) The estimated frequency is 1:500 for heterozygotes, and 1:1,000,000 for homozygotes. Recommendations noting individual selection criteria regarding when treatment should be initiated vary between sources.

Individuals with homozygous hypercholesterolemia are treated aggressively to prevent or slow the progression of cardiovascular disease (CVD) due to the prolonged high lipid level exposure since birth (de Ferranti, 2019; Robinson, 2013). Statins and other agents along with diet and lifestyle modifications are recommended to lower TC. In addition, a new class of drugs, proprotein convertase subtilisin kexin 9 (PCSK9) inhibitors were approved by the FDA to treat familial hypercholesterolemia has been shown to aggressively lower LDL levels. LDL apheresis may be used to selectively lower LDLc to optimal levels.

High-density lipid (HDL) apheresis

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Autologous apheresis with selective HDL delipidation and infusion of selected pre β -HDL is being evaluated as a method to reduce atherosclerosis in individuals at high-risk for cardiovascular disease (CVD). The selective removal of cholesterol from HDL (α -HDL), also called delipidation, involves exposing the plasma to organic solvents in a proprietary collection device and returning the resulting small form of HDL (β -HDL). A randomized, placebo-controlled study examined the safety of a device called LS PDS-2 (Lipid Sciences Plasma Delipidation System-2; Life Sciences, CA) that does not have U.S. FDA approval at this time. A total of 28 individuals with acute coronary syndrome (ACS) who were scheduled for diagnostic cardiac catheterization were enrolled and 26 participants completed all of the treatments and visits. Treatment involved HDL selected apheresis once every 7 days, for a total of 7 treatments. Fourteen individuals randomized to the treatment arm completed the 7-week selective HDL delipidation treatment period and 12 individuals assigned to the control arm completed the placebo treatment period. Pre β -HDL was evaluated by electrophoresis and quantitative analysis. On average, pre β -HDL was increased by 28 times in post-delipidated plasma compared to baseline volume. Intravascular ultrasound (IVUS) of the target vessel was performed and a non-significant reduction in the volume of the atheroma was reported in the delipidated group versus the placebo group. Fifteen of the participants had 1 or more adverse events, with a total of 38 reported adverse events. Hypotension was the most common adverse event. The authors noted limitations to the study that included a small sample size and confounding by the fact that the majority of the participants on the trial were also on statin therapy. Additionally, the authors noted “It is not clear whether acute regression of atherosclerotic burden will be associated with decreased clinical cardiovascular events.” (Waksman, 2010). The evidence does not support that there are improved net clinical health outcomes in individuals treated with therapeutic apheresis combined with selective HDL delipidation. Subedi and associates (2014) notes that further evaluation regarding the association between HDL and ASCVD is needed. The 2019 ASFA guidelines do not address HDL lipid apheresis.

Cytapheresis

Erythrocytapheresis

The ASFA recommends erythrocytapheresis with Category I and the associated Grade 1B level of evidence (strong recommendations, moderate quality evidence) treatment options were listed for the following indications (Connelly-Smith, 2023):

- Hereditary hemochromatosis
- Polycythemia vera

The ASFA notes management of individuals with low-risk polycythemia vera (PV) include maintaining hematocrit $\leq 45\%$ with phlebotomy and low dose aspirin. Individuals with high risk are treated with phlebotomy, aspirin and other cytoreductive agents. Cytapheresis is recommended to correct hyperviscosity as an alternative to emergent large-volume phlebotomy (Connelly-Smith, 2023).

Specialty input recommends use of erythrocytapheresis to treat symptomatic secondary polycythemia vera and porphyria cutanea tarda. Porphyria is a group of rare disorders in which the process of heme production is defective. Porphyria cutanea tarda is the most common type of porphyria.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Ex vivo gene therapy

The development of ex vivo gene therapy and autologous cellular immunotherapy products requires the extraction of hematopoietic stem and progenitor cells (HSPCs). These HSPCs are acquired either through bone marrow aspiration or by mobilizing HSPCs followed by apheresis (Canarutto, 2023). Peripheral stem cell collection requires additional pheresis procedures to ensure a sufficient supply of viable stem cells.

In November 2023, the FDA approved the first cell-based gene therapy as a treatment of sickle cell disease - exagamglogene autotemcel (Casgevy™, Vertex Pharmaceuticals Inc., Boston, MA). In December 2023, lovotibeglogene autotemcel (Lyfgenia®, bluebird bio, Inc., Somerville, MA) was also approved by the FDA as a cell-based gene therapy for the treatment of sickle cell disease. Both therapies are autologous hematopoietic treatments which require a sufficient number of viable, non-sickled red blood cells prior to mobilization and apheresis in order to minimize the risk of potential complications and enhance the collection of viable cells. Erythrocytapheresis or red blood cell exchange is performed approximately 2 months prior to mobilization in order to reach a target of less than 30% of the abnormal hemoglobin variant (hemoglobin S).

Leukocytapheresis

Symptomatic hyperleukocytosis manifests itself with a variety of symptoms including confusion, somnolence, dizziness, headache, coma, parenchymal hemorrhage, hypoxemia, diffuse alveolar hemorrhage, and respiratory failure. Induction chemotherapy with aggressive supportive care is considered the definitive treatment. The ASFA provides category III recommendations for leukocytapheresis for the prophylactic or secondary treatment of hyperleukocytosis (Connelly-Smith, 2023).

RBC exchange

The ASFA has category I-II recommendations for RBC exchange with Grade 1A -2C level of evidence. Treatment options were listed for the following indications (Connelly-Smith, 2023):

- Sickle cell disease, acute:
 - Acute stroke
 - Acute severe, chest syndrome
- Sickle cell disease, non-acute:
 - Stroke prophylaxis
 - Pregnancy
 - Recurrent vaso-occlusive pain crisis

RBC exchange with Category III and an associated Grade 2B or 2C level of evidence was listed for the following indications (Connelly-Smith, 2023):

- Babesiosis, severe
- Malaria, severe
- Red cell alloimmunization, prevention and treatment; Exposure to RhD⁺ RBCs

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

- Transplantation, hematopoietic stem cell, minor ABO incompatible; hematopoietic progenitor cell, apheresis

Additional indications were recommended by specialty input using RBC exchange for the following indications:

- Babesiosis in the high-risk population
- Severe malaria
- Sickle cell disease
 - Pregnancy
 - Primary or secondary stroke prophylaxis

Thrombocytapheresis

Thrombocytapheresis with Category II with “weak recommendation, low-quality or very low-quality evidence” or Category III recommendations were provided for the following indications (Connelly-Smith, 2023):

- Thrombocytosis – symptomatic, prophylactic or secondary

Immunoadsorption Pheresis

Immunoadsorption pheresis with Category I or Category II 1B or 2B recommendations were provided for the following indications (Connelly-Smith, 2023):

- Acute inflammatory demyelinating polyradiculoneuropathy
- Chronic inflammatory demyelinating polyradiculoneuropathy
- Cryoglobulinemia, symptomatic or severe
- Dilated cardiomyopathy, idiopathic; New York functional classification II-IV
- Multiple sclerosis, acute attack/relapse
- Myasthenia gravis, acute short-term or long-term treatment
- Neuromyelitis optical spectrum disorder
- N-methyl-D-aspartate receptor antibody encephalitis
- Transplantation, kidney
- Voltage-gated potassium channel antibody-related diseases

Immunoadsorption pheresis with Category III recommendations were provided for the following indications (Connelly-Smith, 2023):

- Atopic (neuro-) dermatitis (atopic eczema), recalcitrant
- Chronic focal encephalitis
- Coagulation factor inhibitors
- Heparin-induced thrombocytopenia and thrombosis; pre-procedure
- Immune thrombocytopenia, refractory
- Multiple sclerosis, chronic primary or secondary progressive
- Paraneoplastic neurological syndromes

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

- Pemphigus vulgaris
- Thrombotic microangiopathy, infection associated

The ACR 2002 Update of the Rheumatoid Arthritis Guidelines note use of Staphylococcal protein A immunoadsorption column, “should be considered only for patients with refractory rheumatoid arthritis in whom treatment with several disease modifying anti-rheumatic drugs (DMARDs) has failed.” This recommendation was based on the improvement in 31.9% of the participants treated with the immunoadsorption column compared to 11.4% who received sham treatment in a randomized, multicenter trial (Felson, 1999). More recent ACR guidelines (2015, 2021) do not mention IA for the treatment of RA. There have been a few other older studies which support use of IA in the treatment of RA (Furst, 2000; Poullin, 2005).

The peer-reviewed medical literature supports the clinical effectiveness and safety of immunoadsorption for the indications listed in the policy statements. Most studies to date have supported the use of this procedure for individuals with hemolytic uremic syndrome and TTP resulting in a reduction of symptoms. Furthermore, the safety and efficacy of immunoadsorption has been well established in individuals with HUS, TTP, in post kidney transplant recipients with recurrent focal and segmental glomerulosclerosis (FGS). The current research does not support the benefits of immunoadsorption in other immune-related disorders.

Based on the data from the published studies and specialty consensus input, therapeutic apheresis modalities are considered medically necessary for selected indications. Conditions which have been evaluated by the AFSA and have been found to be lacking data showing the efficacy or clinical utility of apheresis or conditions which have not been evaluated by the AFSA and there is a lack of published data showing improved clinical outcomes associated with apheresis when compared to alternative treatments are considered not medically necessary.

Definitions

Antibodies: Immunoglobulins (a specialized immune protein) produced as a result of the introduction of an antigen into the body.

Autoimmune disease: An illness occurring when the body tissues are attacked by its own immune system; as a result, individuals with these diseases frequently have unusual antibodies circulating in their blood that target their own body tissues.

Bullous pemphigoid: A disease characterized by tense blistering eruptions of the skin, generally caused by antibodies abnormally accumulating in a layer of the skin.

Cerebritis: Inflammation of the brain.

Cryoglobulinemia: The presence of abnormal proteins called cryoglobulins that, by definition, have the unusual properties of precipitating from the blood serum when it is chilled and re-dissolving upon rewarming.

Cytapheresis: Subtype of therapeutic apheresis in which the white blood cells are isolated and retained (leukapheresis or lymphocytophoresis) or red blood cells are isolated and retained (erythrocytapheresis).

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

Guillain-Barre: A condition that usually occurs after an infection; the signs and symptoms include loss of sensation in the arms and legs and increasing weakness.

Heterozygotes: A person possessing two different forms of a particular gene, one inherited from each parent.

Homozygotes: A person who has two identical forms of a particular gene, one inherited from each parent.

Immune complex: A combination of an antibody (immunoglobulin), and an antigen (the target that the antibody is attacking).

Immune thrombocytopenic purpura: A condition in which antibodies destroy the cells in the body that is responsible for blood clotting (platelets).

Immunoglobulin: A protein produced by plasma cells and lymphocytes; immunoglobulins are an essential part of the body's immune system which attach to foreign substances, such as bacteria, and assist in destroying them.

Myocarditis: Inflammation of the heart muscle.

Nephritis: Inflammation of the kidney.

Pemphigus vulgaris: An autoimmune disease of the skin, with blistering.

Polymyositis: A chronic inflammatory disease of muscle that begins when white blood cells spontaneously invade muscles, which may result in severe muscle pain, tenderness and weakness.

PRA: Panel reactive antibodies.

Pure red cell aplasia: A condition where an individual has an inability to produce red blood cells.

Regional enteritis: Also called Crohn's disease, a chronic inflammatory disease of the intestine primarily in the small and large intestines but which can occur anywhere in the digestive system between the mouth and the anus.

Scleroderma: A disease of connective tissue resulting in formation of scar tissue in the skin and at times other organs of the body.

Segmental glomerulosclerosis: An illness that occurs when scar tissue forms in some of the glomeruli (structures involved in the filtration of blood) of the kidney.

Thymectomy: Removal of the thymus gland.

Vasculitis: A general term for a group of uncommon diseases characterized by inflammation of the blood vessels.

Waldenström's macroglobulinemia: A disease where abnormal white blood cells produce excessive amounts of antibodies; bleeding and enlarged liver and spleen may be seen.

References

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Peer Reviewed Publications:

1. Allen ES, Stroncek DF, Ren J, et al. Autologous lymphapheresis for the production of chimeric antigen receptor T cells. *Transfusion*. 2017; 57(5):1133-1141.
2. Ariceta G, Besbas N, Johnson S, et al. Guideline for the investigation and initial therapy of diarrhea-negative hemolytic uremic syndrome. *Pediatr Nephrol*. 2009; 24(4):687-696.
3. Balogun RA, Sanchez AP, Klingel R, et al. Update to the ASFA guidelines on the use of therapeutic apheresis in ANCA-associated vasculitis. *J Clin Apher*. 2020; 35: 493–499.
4. Bayraktaroglu Z, Demirci F, Balat O, et al. Plasma exchange therapy in HELLP syndrome: a single-center experience. *Turk J Gastroenterol*. 2006; 17(2):99-102.
5. Bunchorntavakul C, Reddy KR. Pruritus in chronic cholestatic liver disease. *Clin Liver Dis*. 2012; 16(2):331-346.
6. Canarutto D, Omer Javed A, Pedrazzani G, et al. Mobilization-based engraftment of haematopoietic stem cells: a new perspective for chemotherapy-free gene therapy and transplantation. *Br Med Bull*. 2023; 147(1):108-120.
7. Carr RA, Rejowski BJ, Cote GA, et al. Systematic review of hypertriglyceridemia-induced acute pancreatitis: A more virulent etiology? *Pancreatol*. 2016; 16(4):469-476.
8. Cervera R. 8th International Congress on Autoimmunity: new perspectives for refractory catastrophic antiphospholipid syndrome. *Expert Rev Clin Immunol*. 2012; 8(7):617-619.
9. Cervera R, Espinosa G. Update on the catastrophic antiphospholipid syndrome and the “CAPS Registry.” *Semin Thromb Hemost*. 2012; 38(4):333-338.
10. Clark WF. Plasma exchange for renal disease: evidence and use 2011. *J Clin Apher*. 2012; 27(3):112-116.
11. Clark WF. Thrombotic microangiopathy: current knowledge and outcomes with plasma exchange. *Semin Dial*. 2012; 25(2):214-219.
12. Cree BAC, Bennett JL, Kim HJ, et al. Inebilizumab for the treatment of neuromyelitis optica spectrum disorder (N-MOMentum): a double-blind, randomised placebo-controlled phase 2/3 trial. *Lancet*. 2019; 394(10206):1352-1363.
13. Espiritu AI, Pasco PMD. Efficacy and tolerability of azathioprine for neuromyelitis optica spectrum disorder: A systematic review and meta-analysis. *Mult Scler Relat Disord*. 2019; 33:22-32.
14. Donofrio PD, Berger A, Brannagan TH 3rd, et al. Consensus statement: the use of intravenous immunoglobulin in the treatment of neuromuscular conditions report of the AANEM AD HOC committee. American Association of Neuromuscular and Electrodiagnostic Medicine (AANEM). *Muscle Nerve*. 2009; 40(5):890–900.
15. Eser B, Guven M, Unal A, et al. The role of plasma exchange in HELLP syndrome. *Clin Appl Thromb Hemost*. 2005; 11(2):211-217.
16. Felson DT, LaValley MP, Baldassare AR, et al. The Prosorba column for treatment of refractory rheumatoid arthritis: a randomized, double-blind, sham-controlled trial. *Arthritis Rheum*. 1999; 42(10):2153-2159.
17. Gabbay V, Coffey BJ, Babb JS, et al. Pediatric autoimmune neuropsychiatric disorders associated with streptococcus: comparison of diagnosis and treatment in the community and at a specialty clinic. *Pediatrics*. 2008; 122(2):273-278.
18. Gendreau RM; Prosorba Clinical Trial Group. A randomized double-blind sham-controlled trial of the Prosorba column for treatment of refractory rheumatoid arthritis. *Ther Apher*. 2001; 5(2):79-83.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

19. Habermalz B, Sauerland S. Clinical effectiveness of selective granulocyte, monocyte adsorptive apheresis with the Adacolumn device in ulcerative colitis. *Dig Dis Sci*. 2010; 55(5):1421-1428.
20. Jaben EA, Winters JL. Plasma exchange as a therapeutic option in patients with neurologic symptoms due to antibodies to voltage-gated potassium channels: a report of five cases and review of the literature. *J Clin Apher*. 2012; 27(5):267-273.
21. Kohler W, Bucka C, Klingel R. A randomized and controlled study comparing immunoadsorption and plasma exchange in myasthenic crisis. *J Clin Apher*. 2011; 26(6):347-355.
22. Lapeyraque AL, Malina M, Fremaux-Bacchi V, et al. Eculizumab in severe Shiga-toxin-associated HUS. *N Engl J Med*. 2011; 364(26):2561-2563.
23. Loirat C, Frémeaux-Bacchi V. Atypical hemolytic uremic syndrome. *Orphanet J Rare Dis*. 2011; 6:60.
24. Morgan RA, Gray D, Lomova A, Kohn DB. Hematopoietic stem cell gene therapy: Progress and lessons learned. *Cell Stem Cell*. 2017; 21(5):574-590.
25. Morgan SM, Shaz BH, Pavenski K, et al. The top clinical trial opportunities in therapeutic apheresis and neurology. *J Clin Apher*. 2014; 29(6):331-335.
26. Pittock SJ, Berthele A, Fujihara K, et al. Eculizumab in Aquaporin-4-Positive Neuromyelitis Optica Spectrum Disorder. *N Engl J Med*. 2019; 381(7):614-625.
27. Ramos-Casals M, Stone JH, Cid MC, Bosch X. The cryoglobulinaemias. *Lancet*. 2012; 379(9813):348-360.
28. Sander HW, Latov N. Research criteria for defining patients with CIDP. *Neurology*. 2003; 60(8 Suppl 3):S8-S15.
29. Subedi BH, Joshi PH, Jones SR, et al. Current guidelines for high-density lipoprotein cholesterol in therapy and future directions. *Vasc Health Risk Manag*. 2014; 10:205-216.
30. Swedo SE, Leckman JF, Rose NR. From research subgroup to clinical syndrome: modifying the PANDAS criteria to describe PANS (Pediatric Acute-onset Neuropsychiatric Syndrome). *Pediatr Therapeut*. 2012; 2(2):113.
31. Tan J, Smith CH, Goldman, RD. Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections. *Can Fam Physician*. 2012; 58(9):957-959.
32. Taylor CM, Machin S, Wigmore SJ, Goodship TH. Clinical practice guidelines for the management of atypical haemolytic uraemic syndrome in the United Kingdom. *Br J Haematol*. 2010; 148(1):37-47.
33. Uslu-Gökçeoğlu A, Doğan CS, Comak E, et al. Atypical hemolytic uremic syndrome due to factor H antibody. *Turk J Pediatr*. 2013; 55(1):86-89.
34. Waksman R, Torguson R, Kent KM, et al. A first-in-man, randomized, placebo-controlled study to evaluate the safety and feasibility of autologous delipidated high-density lipoprotein plasma infusions in patients with acute coronary syndrome. *J Am Coll Cardiol*. 2010; 55(24):2727-2735.
35. Walsh M, Merkel PA, Peh CA, et al.; PEXIVAS Investigators. Plasma exchange and glucocorticoids in severe ANCA-associated vasculitis. *N Engl J Med*. 2020; 382(7):622-631.
36. Weiss PF, Klink AJ, Friedman DF, Feudtner C. Pediatric therapeutic plasma exchange indications and patterns of use in U.S. children's hospitals. *J Clin Apher*. 2012; 27(6):287-294.
37. Yu X, Gan L, Wang Z, et al. Chemotherapy with or without plasmapheresis in acute renal failure due to multiple myeloma: a meta-analysis. *Int J Clin Pharmacol Ther*. 2015; 53(5):391-397.

Government Agency, Medical Society, and Other Authoritative Publications:

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

1. Ariceta G, Besbas N, Johnson S, et al.; European Paediatric Study Group for HUS. Guideline for the investigation and initial therapy of diarrhea-negative hemolytic uremic syndrome. *Pediatr Nephrol.* 2009; 24(4):687-696.
2. Carless PA, Rubens FD, Anthony DM, et al. Platelet-rich-plasmapheresis for minimising peri-operative allogeneic blood transfusion. *Cochrane Database Syst Rev.* 2011; (3):CD004172.
3. Centers for Medicare and Medicaid Services. National Coverage Determination for Apheresis (Therapeutic Pheresis). NCD#110.14 Effective July 30, 1992. Available at: <https://www.cms.gov/medicare-coverage-database/new-search/search.aspx>. Accessed on January 2, 2025.
4. Chevret S, Hughes RA, Annane D. Plasma exchange for Guillain-Barré syndrome. *Cochrane Database Syst Rev.* 2017; 2:CD001798.
5. Colvin MM, Cook JL, Chang P, et al.; American Heart Association Heart Failure and Transplantation Committee of the Council on Clinical Cardiology; American Heart Association Heart Failure and Transplantation Committee of the Council on Cardiopulmonary Critical Care, Perioperative and Resuscitation; American Heart Association Heart Failure and Transplantation Committee of the Council on Cardiovascular Disease in the Young; American Heart Association Heart Failure and Transplantation Committee of the Council on Clinical Cardiology, Council on Cardiovascular and Stroke Nursing; American Heart Association Heart Failure and Transplantation Committee of the Council on Cardiovascular Radiology and Intervention; American Heart Association Heart Failure and Transplantation Committee of the Council on Cardiovascular Surgery and Anesthesia. Antibody-mediated rejection in cardiac transplantation: emerging knowledge in diagnosis and management: a scientific statement from the American Heart Association. *Circulation.* 2015; 131(18):1608-1639.
6. Connelly-Smith L, Alquist CR, Aqui NA, et al. Guidelines on the use of therapeutic apheresis in clinical practice - evidence-based approach from the Writing Committee of the American Society for Apheresis: The Ninth Special Issue. *J Clin Apher.* 2023; 38(2):77-278.
7. Cortese I, Chaudhry V, So YT, et al. Evidence-based guideline update: plasmapheresis in neurologic disorders: report of the Therapeutics and Technology Assessment Subcommittee of the American Academy of Neurology. *Neurology.* 2011; 76(3):294-300.
8. Costanzo MR, Dipchand A, Starling R, et al. The International Society of Heart and Lung Transplantation guidelines for the care of heart transplant recipients. *J Heart Lung Transplant.* 2010; 29(8):914-956.
9. de Ferranti SD, Steinberger J, Ameduri R, et al. Cardiovascular risk reduction in high-risk pediatric patients: a scientific statement from the American Heart Association. *Circulation.* 2019; 139(13):e603-e634.
10. U.S. Food and Drug Administration (FDA). Product Label. Accessed on January 2, 2025.
 - Casgevy™ highlights of prescribing information. Revised on December 2023. Available at: <https://www.fda.gov/media/174615/download?attachment>.
 - Lyfgenia® highlights of prescribing information. Revised on December 2023. Available at: <https://www.fda.gov/media/174610/download>.
11. Fraenkel L, Bathon JM, England BR, et al. 2021 American College of Rheumatology guideline for the treatment of rheumatoid arthritis. *Arthritis Care Res (Hoboken).* 2021; 73(7):924-939.
12. Furst D, Felson D, Thoren G, Gendreau RM. Immunoabsorption for the treatment of rheumatoid arthritis: final results of a randomized trial. *Prosorba Trial Investigators. Ther Apher.* 2000; 4(5):363-373.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

13. Gabbay V, Coffey BJ, Babb JS, et al. Pediatric autoimmune neuropsychiatric disorders associated with streptococcus: comparison of diagnosis and treatment in the community and at a specialty clinic. *Pediatrics*. 2008; 122(2):273-278.
14. Gordon PA, Winer JB, Hoogendijk JE, Choy EH. Immunosuppressant and immunomodulatory treatment for dermatomyositis and Polymyositis. *Cochrane Database Syst Rev*. 2012; (8):CD003643.
15. Hahn BH, McMahon MA, Wilkinson A, et al. American College of Rheumatology Guidelines for screening, treatment, and management of lupus nephritis. *Arthritis Care Res (Hoboken)*. 2012; 64(6):797-808.
16. Hughes RA, Bouche P, Cornblath DR, et al. European Federation of Neurological Societies/Peripheral Nerve Society guideline on management of chronic inflammatory demyelinating polyradiculoneuropathy: report of a joint task force of the European Federation of Neurological Societies and the Peripheral Nerve Society. *Eur J Neurol*. 2006; 13(4):326-332.
17. Ito MK, McGowan MP, Moriarty PM; National Lipid Association Expert Panel on Familial Hypercholesterolemia. Management of familial hypercholesterolemias in adult patients: recommendations from the National Lipid Association Expert Panel on familial hypercholesterolemia. *J Clin Lipidol*. 2011; 5(3 Suppl):S38-S45.
18. Jacobson TA, Maki KC, Orringer CE, et al.; NLA Expert Panel. National Lipid Association Recommendations for Patient-Centered Management of Dyslipidemia: Part 2. *J Clin Lipidol*. 2015; 9(6 Suppl):S1-S122.e1.
19. Medical Advisory Secretariat. Low density lipoprotein apheresis: an evidence-based analysis. *Ontario Health Technology Assessment Series*. 2007; 7(5):1-101.
20. Mehndiratta MM, Hughes RA, Pritchard J. Plasma exchange for chronic inflammatory demyelinating polyradiculoneuropathy. *Cochrane Database Syst Rev*. 2015; (8):CD003906.
21. National Heart Lung and Blood Institute (NHLBI). Integrated guidelines for cardiovascular health and risk reduction in children and adolescents. January 2013. Available at: http://www.nhlbi.nih.gov/guidelines/cvd_ped/index.htm. Accessed on January 3, 2025.
22. NCCN Clinical Practice Guidelines in Oncology® (NCCN). © 2025 National Comprehensive Cancer Network, Inc. For additional information visit the NCCN website: <http://www.nccn.org>. Accessed on January 30, 2025.
 - Acute Myeloid Leukemia (V2.2025). Revised January 27, 2025.
 - Multiple Myeloma (V1.2025). Revised September 17, 2024.
 - Waldenstrom Macroglobulinemia/Lymphoplasmacytic Lymphoma (V2.2025). Revised December 18, 2024.
23. Neunert C, Lim W, Crowther M, et al. The American Society of Hematology 2011 evidence-based practice guideline for immune thrombocytopenia. *Blood*. 2011; 117(16):4190-4207.
24. Neunert C, Terrell DR, Arnold DM, et al. American Society of Hematology 2019 guidelines for immune thrombocytopenia. *Blood Adv*. 2019; 3(23):3829-3866.
25. Nordestgaard BG, Chapman MJ, Humphries SE, et al.; European Atherosclerosis Society Consensus Panel. Familial hypercholesterolaemia is underdiagnosed and undertreated in the general population: guidance for clinicians to prevent coronary heart disease: consensus statement of the European Atherosclerosis Society. *Eur Heart J*. 2013; 34(45):3478-3490.
26. Padmanabhan A, Connelly-Smith L, Aqui N, et al. Guidelines on the use of therapeutic apheresis in clinical practice - evidence-based approach from the Writing Committee of the American Society for Apheresis: the eighth special issue. *J Clin Apher*. 2019; 34(3):171-354.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

27. Poullin P, Announ N, Mugnier B, et al. Protein A-immunoabsorption (Prosorba column) in the treatment of rheumatoid arthritis. *Joint Bone Spine*. 2005; 72(2):101-103.
28. Robinson JG. Management of familial hypercholesterolemia: a review of the recommendations from the National Lipid Association Expert Panel on Familial Hypercholesterolemia. *J Manag Care Pharm*. 2013; 19(2):139-149.
29. Schwartz J, Padmanabhan A, Aqui N, et al. Guidelines on the use of therapeutic apheresis in clinical practice—evidence-based approach from the Writing Committee of the American Society for Apheresis: the seventh special issue. *J Clin Apher*. 2016; 31(3):149-162.
30. Singh JA, Saag KG, Bridges SL Jr, et al.; American College of Rheumatology. 2015 American College of Rheumatology Guideline for the Treatment of Rheumatoid Arthritis. *Arthritis Care Res (Hoboken)*. 2016; 68(1):1-25.
31. Smith JW, Weinstein R, Hillyer KL; AABB Hemapheresis Committee; American Society for Apheresis. Therapeutic apheresis: a summary of current indication categories endorsed by the AABB and the American Society for Apheresis. *Transfusion*. 2003; 43(6):820-822.
32. Van den Bergh PY, Hadden RD, Bouche P, et al.; European Federation of Neurological Societies; Peripheral Nerve Society. European Federation of Neurological Societies/Peripheral Nerve Society guideline on management of chronic inflammatory demyelinating polyradiculoneuropathy: report of a joint task force of the European Federation of Neurological Societies and the Peripheral Nerve Society - first revision. *Eur J Neurol*. 2010; 17(3):356-363.
33. Van den Bergh PYK, van Doorn PA, Hadden RDM, et al. European Academy of Neurology/Peripheral Nerve Society guideline on diagnosis and treatment of chronic inflammatory demyelinating polyradiculoneuropathy: Report of a joint Task Force-Second revision. *Eur J Neurol*. 2021; 28(11):3556-3583.
34. Velleca A, Shullo MA, Dhital K, et al. The International Society for Heart and Lung Transplantation (ISHLT) guidelines for the care of heart transplant recipients. *J Heart Lung Transplant*. 2023; 42(5):e1-e141.

Websites for Additional Information

1. National Institute of Neurological Disorders and Stroke (NINDS). NINDS Disease Information. Available at: <https://www.ninds.nih.gov/Disorders/All-Disorders>. Accessed on January 3, 2025.
 - Polymyositis Information Page
 - Refsum Disease Information Page
 - Sydenham Chorea Information Page
2. National Library of Medicine. MedlinePlus. Exchange Transfusion. Last reviewed November 6, 2023. Available at: <https://medlineplus.gov/ency/article/002923.htm>. Accessed on January 3, 2025.
3. National Organization for Rare Disorders (NORD). Chronic Inflammatory Demyelinating Polyneuropathy. Last updated 10/1/2024. Available at: <https://rarediseases.org/rare-diseases/chronic-inflammatory-demyelinating-polyneuropathy/>. Accessed on January 3, 2025.

Index

Adacolumn
Apheresis, therapeutic

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

Cellsorba®
 Cytapheresis
 Immunoabsorption
 LDL apheresis
 Lipsorb®
 Pheresis
 Plasma Exchange
 Plasmapheresis
 Prosorba®
 Protein A column
 Selective HDL delipidation

The use of specific product names is illustrative only. It is not intended to be a recommendation of one product over another, and is not intended to represent a complete listing of all products available.

History

Status	Date	Action
	10/01/2025	Updated Coding section with 10/01/2025 ICD-10-CM changes, added G35.A-G35.D replacing G35, E78.010 and E78.011-E78.019 replacing E78.01, and N04.B2 to end of range.
Revised	02/20/2025	Medical Policy & Technology Assessment Committee (MPTAC) review. Added <i>is</i> to the vasculitis anti-neutrophil cytoplasmic antibodies MN indication for plasmapheresis. Revised Description, Discussion, References, and Websites sections.
	10/01/2024	Updated Coding section with 10/01/2024 ICD-10-CM changes, added C88.00-C88.01 replacing C88.0.
Revised	05/09/2024	MPTAC review. Added MN indications to the cytapheresis section regarding erythrocytapheresis and red blood cell exchange for when used as part of the development of an FDA-approved ex vivo gene therapy. Updated Description, Discussion and References sections. Updated coding section, added ICD-10-CM dx code D56.1.
Revised	02/15/2024	MPTAC review. Added <i>erythropoietic protoporphyria, liver disease</i> to the plasmapheresis and RBC exchange and <i>heart transplantation, desensitization/rejection prophylaxis</i> to the plasmapheresis medically necessary sections. Updated Discussion and References sections. . Updated Coding section, added ICD-10-CM codes E80.0, K77 for plasmapheresis, cytapheresis.
Reviewed	02/16/2023	MPTAC review. Updated Description, Discussion, References and Websites sections.
	09/28/2022	Updated Coding section with 10/01/2022 ICD-10-CM changes; added D59.30-D59.39 replacing D59.3.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

	08/24/2022	Updated the Description section adding <i>ex vivo gene therapy</i> and an example to the Note regarding apheresis therapy which is not addressed in the document.
Reviewed	02/17/2022	MPTAC review. Updated Discussion, References and Websites sections.
	10/01/2021	Updated Coding section with 10/01/2021 ICD-10-CM changes; added M31.10-M31.19 replacing M31.1 deleted 09/30/2021.
Revised	02/11/2021	MPTAC review. Removed medically necessary indication for lymphapheresis as a component of CAR-T therapy. Updated Description, Discussion and References sections. Reformatted Coding section and removed CAR-T therapy related diagnosis codes.
	10/01/2020	Updated Coding section with 10/01/2020 ICD-10-CM changes; added B60.00-B60.09 replacing B60.0, D59.12 replacing D59.0-D59.1 and N01.A, N03.A, N04.A.
Revised	05/14/2020	MPTAC review. Added requirement that medical treatments must have been tried and failed prior to therapeutic apheresis to medically necessary category II indications, with the exception of symptomatic hyperleukocytosis. Removed the following medically necessary indications: H.E.L.L.P Syndrome of pregnancy, progressive multifocal leukoencephalopathy (natalizumab associated), Sydenham's chorea, symptomatic secondary polycythemia, severe malaria, multi organ failure syndrome, pre-procedure preparation for surgery or hematopoietic stem cell transplant, iron overload and moderate to severe rheumatoid arthritis. Removed examples from the not medically necessary statements. Updated Discussion, References, and Coding section.
Revised	11/07/2019	MPTAC review. Added diagnostic criteria to the condition chronic inflammatory demyelinating polyradiculoneuropathy (CIDP) when treated by plasmapheresis or immunoadsorption. Updated Discussion/General Information and References sections.
Revised	08/22/2019	MPTAC review. Added medically necessary indications for acute inflammatory demyelinating polyradiculoneuropathy/Guillain-Barre syndrome, chronic inflammatory demyelinating polyradiculoneuropathy, myasthenia gravis, N-methyl D-aspartate receptor antibody encephalitis and renal transplant recipients. Revised the medically necessary indication for anti-neutrophil cytoplasmic antibodies vasculitis. Added not medically necessary indications for red blood cell exchange and adsorptive cytappheresis when criteria are not met. Clarified not medically necessary statements. Removed clarifying not medically necessary statement regarding immunoadsorption. Clarified term Low-density lipid (LDL) apheresis as also lipoprotein apheresis. Clarified several medically necessary indications based on 2019 ASFA indications. Updated Description, References, Websites and Coding sections.
Reviewed	11/08/2018	MPTAC review.
Reviewed	10/31/2018	Hematology/Oncology Subcommittee review. Updated Discussion/General Information and References sections.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.

Therapeutic Apheresis

Reviewed	03/22/2018	MPTAC review. Updated References section.
New	11/02/2017	MPTAC review. Initial document development.
New	11/01/2017	Hematology/Oncology Subcommittee review. Initial document development. Moved content of MED.00113 Therapeutic Apheresis to new clinical utilization management guideline document with the same title. Added medically necessary indications for N-methyl D-aspartate receptor antibody encephalitis, progressive multifocal leukoencephalopathy and apheresis as a component of CAR-T therapy. Revised medically necessary indications for polyneuropathy, glomerulonephritis, neuromyelitis optica disorders and paraproteinemic demyelinating neuropathies. Updated Coding section with 01/01/2018 CPT descriptor change for code 36516 and removed code 36515 deleted 12/31/2017.

Federal and State law, as well as contract language including definitions and specific coverage provisions/exclusions, and Medical Policy take precedence over Clinical UM Guidelines and must be considered first in determining eligibility for coverage. The member's contract benefits in effect on the date that services are rendered must be used. Clinical UM Guidelines, which address medical efficacy, should be considered before utilizing medical opinion in adjudication. Medical technology is constantly evolving, and we reserve the right to review and update Clinical UM Guidelines periodically. Clinical UM guidelines are used when the plan performs utilization review for the subject. Due to variances in utilization patterns, each plan may choose whether or not to adopt a particular Clinical UM Guideline. To determine if review is required for this Clinical UM Guideline, please contact the customer service number on the back of the member's card.

No part of this publication may be reproduced, stored in a retrieval system or transmitted, in any form or by any means, electronic, mechanical, photocopying, or otherwise, without permission from the health plan.