SENTARA COMMUNITY PLAN (MEDICAID)

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-305-2331</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

Gammaked[™] [Immune Globulin Injection

Drug Requested: Check applicable box below. If not checked, authorization could be delayed.

<u>Drug Requested</u>: Immune Globulin Intravenous (IVIG) (Medical) (Miscellaneous Disorders)

	(Human), 10% Liquid] (J1556)		(Human), 10% Caprylate/Chromatography Purified] (J1561)
	Carimune® NF [Nanofiltered, Immune Globulin Intravenous (Human)] (J1566)		Gamunex®-C [Immune Globulin Injection (Human), 10% Caprylate/Chromatography Purified] (J1561)
	Flebogamma® DIF [Human Normal Immunoglobulin (IVIg)] (J1572)		Octagam® [Immune Globulin Intravenous (Human) liquid preparation] (J1568)
	Gammagard® Liquid [Immune Globulin Infusion (Human), 10% Solution, for intravenous and subcutaneous administration] (J1569)		Panzyga [®] [Immune Globulin Intravenous (Human) – ifas 10% Liquid Preparation] (J1576)
	Gammagard® S/D [Immune Globulin Intravenous (Human) Solvent/Detergent Treated (Freeze-Dried Concentrate)] (J1556)		Privigen® [Immune Globulin Intravenous (Human), 10% Liquid] (J1459)
M	EMBER & PRESCRIBER INFORMATION	V: A	Authorization may be delayed if incomplete.
	EMBER & PRESCRIBER INFORMATION mber Name:		· · · · · · · · · · · · · · · · · · ·
Mei	EMBER & PRESCRIBER INFORMATION mber Name: mber Sentara #:		
Mei Mei	mber Name:		Date of Birth:
Me Me Pre	mber Name:mber Sentara #:		Date of Birth:
Mei Mei Pre	mber Name: mber Sentara #: scriber Name:		Date of Birth: Date:
Men Men Pre Pre	mber Name: mber Sentara #: scriber Name: scriber Signature:		Date of Birth: Date:
Men Men Pre Pre Off	mber Name:		Date of Birth: Date:

DRUG INFORMA	ATION: Authorization may be	delayed if incomplete.
Drug Form/Strength:		
Dosing Schedule:		Length of Therapy:
Diagnosis:		ICD Code, if applicable:
Weight:		Date:
Height:	inches	
Dosing should be calcul than his or her ideal bod		if the member's actual body weight is 20% higher
• IBW (kg) for ma	ght = IBW + 0.5 (actual body we ales = $50 + [2.3$ (height in inches nales = $45.5 + [2.3 \times (2.3 \times (2.3 \times 1.3 \times 1$	-60)]
IMPORTANT -] <mark>approval</mark> will be gra	· ·	body weight is not accepted, a PARTIAL
and subsequently stop Γ	VIG therapy if improvement is su	for renewal requests when improvement has occurred ustained with a dose reduction (this does NOT apply tong as immunoglobulin levels are maintained in the
		ne does not jeopardize the life or health of the member d would not subject the member to severe pain.
approval. To support ea		below all that apply. All criteria must be met for on, including lab results, diagnostics, and/or chart
Initial Approval Renewal Approv	immune blistering disorde - 3 months (Dose: 2g/kg divided al - 6 months; renewal is based and disorder is still present.	
	r one of the diagnoses:	
□ Bullous Pemphi		☐ Epidermolysis Bullosa Acquisita
	.a. Cicatricial Pemphigoid)	☐ Pemphigus Foliaceus
□ Pemphigus Vulg		
		<u>. I</u>

•	Во	th boxes below must be checked.
		Failure, contraindication, or intolerance of conventional therapy with high dose corticosteroids <u>and</u> one immunosuppressant therapies (e.g., azathioprine, cyclophosphamide, mycophenolate, methotrexate)
		AND
		Rapidly progressive disease in which a clinical response cannot be affected quickly enough using conventional drugs
3	If a	agnosis - Immune Thrombocytopenic Purpura (ITP). For ONE (1) treatment. Inother treatment is warranted, PA must be resubmitted. (Dose: 2g/kg divided over 5 s OR 1g/kg once daily for 2 consecutive days in a 28-day cycle)
		Platelet count <30; OR
		Platelet count <50 w/ bleed; AND
		Trial and failure of high dose steroid for 7 days
	Au eit	lagnosis - Acute exacerbation Myasthenia Gravis. Ithorized Approval - ONE month only; submit status report.* (Dose: 1-2g/kg divided as her 0.5g/kg daily x 2 days OR 0.4g/kg daily x 5 days, one course) PLEASE NOTE: IVIG is not covered for maintenance therapy.*
		Member has a positive serologic test for anti-acetylcholine receptor (AchR) antibodies; AND
		Member has an acute exacerbation (within last 60 days) resulting in impending myasthenic crisis (i.e., respiratory compromise, acute respiratory failure, and/or bulbar compromise); AND
		Member is failing on conventional immunosuppressant therapy high dose corticosteroids <u>AND</u> one of the following within the <u>last four (4) months</u> : azathioprine, cyclosporine, mycophenolate, methotrexate, tacrolimus, cyclophosphamide; <u>AND</u>
		Member will be on combination therapy with corticosteroids or other immunosuppressants (e.g., azathioprine, mycophenolate, cyclosporine, methotrexate, tacrolimus, cyclophosphamide)
	In	agnosis - Lambert-Eaton Myasthenic Syndrome. itial Approval - <mark>6 months.</mark> enewal – <mark>12 months</mark> if diagnosis still present.
		Member has a history of failure, contraindication, or intolerance to a high dose corticosteroid; AND
		Member has a history of failure, contraindication, or intolerance to an immunomodulatory monotherapy (e.g., azathioprine, cyclophosphamide, mycophenolate, methotrexate); <u>AND</u>
		Medication will be used with concomitant immunomodulator therapy (e.g., azathioprine, corticosteroids) for long-term management unless contraindicated

Str	agnosis - PANDAS (Pediatric Autoimmune Neuropsychiatric Disorders Associated with reptococcus Infections). athorized Approval – 12 months
Tr	agnosis - Prevention/Treatment of Solid Organ (kidney, liver, lung, heart, and pancreas) cansplant Rejection. athorized Approval - 3 months (Dose: 2g/kg divided over 5 days, 28 days).
	Coverage is provided for one or more of the following (list is not all-inclusive).
	Suppression of panel reactive anti-human leukocyte antigen (HLA) antibodies prior to transplantation
	Treatment of antibody-mediated rejection of solid organ transplantation
	Prevention or treatment of viral infections (e.g., cytomegalovirus, Parvo B-19 virus, and Polyoma BK virus)
	agnosis - Dermatomyositis/Polymyositis. athorized Approval - 6 months (Dose: 2g/kg divided over 2-5 days, 28 days).
	Member has severe active disease.
	DERMATOMYOSITIS – extramuscular manifestations of the antisynthetase syndrome including rash [such as Gottron's papules or a heliotrope rash), polyarthritis, Raynaud phenomenon, and interstitial lung disease, particularly if an antisynthetase antibody (most often anti-Jo-1) is present and marked elevation of muscle enzymes]
	<u>OR</u>
	POLYMYOSITIS
	AND
	Member has proximal weakness in all upper and/or lower limbs; AND
	Diagnosis has been confirmed by muscle biopsy (choose one below) (must submit biopsy results): Dermatomyositis biopsy – evidence of injury to capillaries and perifascicular myofibers and abnormal muscle fibers are usually grouped in one portion of the fascicle, suggestive of microinfarction mediated by blood vessel dysfunction
	<u>OR</u>
	Polymyositis biopsy – cellular infiltrate is predominantly within the fascicle, with inflammatory cells invading individual muscle fibers AND abnormal necrotic and regenerating muscle fibers are scattered throughout the fascicle and are not limited to one portion; AND
	Member has failed a trial (verify through paid claims) of high dose corticosteroids (Prednisone 0.5-1 mg/kg/day for $2-4$ weeks) within the last 4 months and document failure in progress notes; $\underline{\mathbf{AND}}$
	Member has failed a trial (verify through paid claims) of an immunosuppressant (e.g., methotrexate, azathioprine, etc.) within the last 6 months; <u>AND</u>

		Must be used as part of combination therapy with other agents; AND
		Member has a documented baseline physical exam and muscular strength/function prior to steroid and post-steroid. Baseline prior to IVIG. Dates taken (must be provided):
		AND
		Serum creatine kinase (CK) levels and dates taken: (must be provided):
		ontinuation of therapy for Dermatomyositis/Polymyositis. Progress notes/lab sessments must be submitted
		Muscle biopsy documents normal (IVIG DISCONTINUE)
		Muscle biopsy documents Dermatomyositis/Polymyositis improving (Approval - 6 months)
		Muscle biopsy documents no improvement with IVIG post-6 months (IVIG DISCONTINUE)
		Muscular strength/function improvement. Dates taken:
		Serum creatine kinase (CK) levels and dates taken: (must be provided):
	Αι	agnosis - Fetal Alloimmune Thrombocytopenia (FAIT). Ithorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until ivery)
	Au del	thorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until
	Au del	thorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until ivery)
	Au del	nthorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until ivery) Previous FAIT pregnancy; OR
	Au del	thorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until ivery) Previous FAIT pregnancy; OR Family history of the disease; OR
	Audelder	thorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until ivery) Previous FAIT pregnancy; OR Family history of the disease; OR Screening reveals platelet alloantibodies agnosis - Neonatal Alloimmune Thrombocytopenia (NAIT).
0	Audelder	Previous FAIT pregnancy; OR Family history of the disease; OR Screening reveals platelet alloantibodies agnosis - Neonatal Alloimmune Thrombocytopenia (NAIT). athorized Approval - One month (1 course) Only; Cannot be renewed. agnosis - Acquired Immune Deficiency secondary to Acute Lymphoblastic cukemia (ALL).
0	Di Au	rthorized Approval until delivery; Cannot be renewed. (Dose: 1g/kg weekly until ivery) Previous FAIT pregnancy; OR Family history of the disease; OR Screening reveals platelet alloantibodies agnosis - Neonatal Alloimmune Thrombocytopenia (NAIT). Ithorized Approval - One month (1 course) Only; Cannot be renewed. agnosis - Acquired Immune Deficiency secondary to Acute Lymphoblastic eukemia (ALL). Ithorized Approval - 4 months (Dose: 400mg/kg every 3-4 weeks)

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Le	iagnosis - Acquired Immune Deficiency secondary to Chronic Lymphocytic eukemia (CLL)† OR Multiple Myeloma (MM)‡
Αl	uthorized Approval - 6 months (Dose: 400mg/kg every 3-4 weeks)
	Member's IgG level is $\leq 200 \text{ mg/dL } \mathbf{OR}$ both of the following:
	☐ Member has history of multiple hard to treat infections as indicated by at least one of the following
	☐ Four or more ear infections within 1 year
	☐ Two or more serious sinus infections within 1 year
	Two or more months of antibiotics with little effect
	☐ Two or more pneumonias within 1 year
	□ Recurrent or deep skin abscesses
	□ Need for intravenous antibiotics to clear infections
	☐ Two or more deep-seated infections including septicemia
	AND
	☐ Member has a deficiency in producing antibodies in response to vaccinations
	<u>AND</u>
	Titers were drawn before challenging with vaccination; AND
	Titers were drawn between 4 and 8 weeks of vaccination
Di	iagnosis - Relapsing-Remitting Multiple Sclerosis*
	Clinical records, labs, x-rays, etc. supporting the diagnosis of RRMS must be provided.
	Current medications and treatment plan with initiation of IVIG, including use of IVIG as monotherapy
	Initiation of immunosuppressive treatment, if necessary
	iagnosis - Kawasaki Syndrome (Pediatric Patients).
Al	uthorized Approval – 1 month; Cannot be renewed. (Dose: 1g/kg to 2g/kg, one course).
	Date of initial onset of symptoms:
	AND
	Current medications and treatment plan with initiation of IVIG:
-	
Αι	iagnosis - Allogeneic Bone Marrow or Stem Cell Transplant. uthorized Approval - 6 months (Dose: 500 mg/kg once weekly x 90 days, then 500 mg/kg ery 3-4 weeks)
	Used for prevention of acute Graft-Versus-Host-Disease (aGVHD) or infection; AND
	Member's BMT was allogeneic; AND
	Member's IgG level is less than 400 mg/dL

Di	agnosis - Hyperbilirubinemia in the newborn
If	agnosis - Guillain-Barre Syndrome. For ONE (1) treatment. another treatment is warranted, PA must be resubmitted. aximum of 2 treatments. (Initial Dose: 2g/kg divided over 5 days, 2 courses)
	Defined by the following:
	□ Bilateral & flaccid weakness of the limbs; <u>AND</u>
	□ Decreased or absent deep tendon reflexes in weak limbs; <u>AND</u>
	☐ Monophasic illness pattern and interval between onset and nadir of weakness between 12h and 28 days and subsequent clinical plateau; AND
	□ Electrophysiological findings consistent with GBS; <u>AND</u>
	□ Cytoalbuminologic dissociation (elevation of CSF protein level above laboratory normal value &/or CSF total white count <50 cells/µL; AND
	☐ Member is non-ambulatory and 4 weeks or less have elapsed since onset of symptoms
	agnosis - HIV Infection/children. oproval – 4 months. (Dose: 400 mg/kg every 2-4 weeks)
	In conjunction w/ AZT or other antiretroviral, to prevent mild to severe bacterial infection w/CD4+ counts $\!<\!200/\!\text{uL}$
	In conjunction w/ AZT, to prevent maternal transmission of HIV infection
	HIV-positive children exposed to measles or live in a high-prevalence measles area
	HIV-related ITP
	agnosis - Stiff-Person Syndrome. oproval - 6 months (Dose: 2g/kg divided over 5 days, 28-days)
	Member has anti-glutamic acid decarboxylase (GAD) antibodies; AND
	Member failed at least 2 of the following treatments within the last 9 months from request of diagnosis: benzodiazepines, baclofen, gabapentin, valproate, tiagabine, or levetiracetam; <u>AND</u>
	Member has documented baseline on physical exam
	Reauthorization for Stiff-Person Syndrome:
	□ Positive anti-glutamic acid decarboxylase antibodies; <u>AND</u>
	Baseline physical exam document improvement: member must meet <u>one</u> of the criteria below:
	□ Negative signs – discontinue IVIG
	OR □ Positive signs – continue and TAPER down IVIG
	- 1 Usitive signs - continue and IAI EIX down IVIO

		agnosis - Toxic Shock Syndrome.
	A	oproval – 4 months; Cannot be renewed (Dose: 2g/kg divided over 5 days, one course)
		Infection is refractory to aggressive treatment. Include therapies tried.
		Presence of an undrainable focus
		Persistent oliguria with pulmonary edema
M	edi	cation being provided by (check box below that applies):
M	edi	Location/site of drug administration:
M		
M		Location/site of drug administration:
M		Location/site of drug administration:

For urgent reviews: Practitioner should call Sentara Health Plans Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. Sentara Health Plan's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

Use of samples to initiate therapy does not meet step edit/preauthorization criteria.

Previous therapies will be verified through pharmacy paid claims or submitted chart notes.