Coverage Summary
Medications/Drugs (Outpatient/Part B)

Policy Number: M-005
Products: UnitedHealthcare Medicare Advantage Plans
Original Approval Date: 12/15/2008
Approved by: UnitedHealthcare Medicare Benefit Interpretation Committee
Last Review Date: 08/20/2019

Related Medicare Advantage Policy Guidelines:

- Coverage of Drugs and Biologicals for Label and Off-Label Uses
- Dermal Injections for the Treatment of Facial Lipodystrophy Syndrome (LDS) (NCD 250.5)
- Dimethyl Sulfoxide (DMSO) (NCD 230.12)
- Erythropoiesis Stimulating Agents (ESAs) in Cancer and Related Neoplastic Conditions (NCD 110.21)
- Erythropoietin Stimulating Agent (ESA)
- Halaven® (Eribulin Mesylate)
- Intravenous Immune Globulin (IVIG)
- Intravenous Immune Globulin-for the Treatment of Mucocutaneous Blistering Diseases (NCD 250.3)
- Intravenous Iron Therapy (NCD 110.10)
- Laetrile and Related Substances (NCD 30.7)
- L-Dopa (NCD 160.17)
- Nesiritide for Treatment of Heart Failure Patients (NCD 200.1)
- Self-Administered Drug(s) (SAD)
- Testosterone Replacement Therapy
- Thrombolytic Agents
- Vitamin B12 Injections to Strengthen Tendons, Ligaments, etc., of the Foot (NCD 150.6)
- Xgeva®, Prolia® (Denosumab)

This information is being distributed to you for personal reference. The information belongs to UnitedHealthcare and unauthorized copying, use, and distribution are prohibited. This information is intended to serve only as a general reference resource and is not intended to address every aspect of a clinical situation. Physicians and patients should not rely on this information in making health care decisions. Physicians and patients must exercise their independent clinical discretion and judgment in determining care. Each benefit plan contains its own specific provisions for coverage, limitations, and exclusions as stated in the Member’s Evidence of Coverage (EOC)/Summary of Benefits (SB). If there is a discrepancy between this policy and the member’s EOC/SB, the member’s EOC/SB provision will govern. The information contained in this document is believed to be current as of the date noted.

The benefit information in this Coverage Summary is based on existing national coverage policy, however Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable.

There are instances where this document may direct readers to a UnitedHealthcare Commercial Medical Policy, Medical Benefit Drug Policy, and/or Coverage Determination Guideline (CDG). In the absence of a Medicare National Coverage Determination (NCD), Local Coverage Determination (LCD), or other Medicare coverage guidance, CMS allows a Medicare Advantage Organization (MAO) to create its own coverage determinations, using objective evidence-based rationale relying on authoritative evidence (Medicare IOM Pub. No. 100-16, Ch. 4, §90.5).

INDEX TO COVERAGE SUMMARY

I. COVERAGE
   1. Outpatient Medications/Drugs
      a. Part B Medications/Drugs
      b. Part D Medications/Drugs
      c. Part B vs Part D Medications/Drugs
   2. Unlabeled Use of a Part B Drug
   3. Examples of medications/drugs that are covered under Part B
      a. Durable Medical Equipment (DME) Supply Drugs (e.g., nebulizer inhalation drugs and infusion pump drugs)
      b. Immunosuppressive Drugs
      c. Hemophilia Clotting Factors
<table>
<thead>
<tr>
<th></th>
<th>Medications/Drugs (Outpatient/Part B)</th>
</tr>
</thead>
<tbody>
<tr>
<td>d.</td>
<td>Oral Anti-Cancer Drugs and Oral Anti-Emetics</td>
</tr>
<tr>
<td>e.</td>
<td>Immunizations</td>
</tr>
<tr>
<td>f.</td>
<td>Antigens/Antihistamines</td>
</tr>
<tr>
<td>g.</td>
<td>Parenteral Nutrition</td>
</tr>
<tr>
<td>h.</td>
<td>Intravenous Immune Globulin (IVIG)</td>
</tr>
<tr>
<td>i.</td>
<td>Intravenous Iron Therapy</td>
</tr>
<tr>
<td>j.</td>
<td>Injectable Drugs for the Treatment of Osteoporosis</td>
</tr>
<tr>
<td>k.</td>
<td>L-Dopa (also see #4.d)</td>
</tr>
<tr>
<td>l.</td>
<td>Dimethyl Sulfoxide (DMSO)</td>
</tr>
<tr>
<td>m.</td>
<td>Dermal Injections for the Treatment of Facial Lipodystrophy Syndrome (LDS)</td>
</tr>
<tr>
<td>n.</td>
<td>Drugs Treated as Hospital Outpatient Supplies</td>
</tr>
<tr>
<td>o.</td>
<td>Hereditary Angioedema (HAE) Treatment</td>
</tr>
</tbody>
</table>

4. Examples of medications/drugs that are not covered
   a. Vitamin B12 Injections
   b. Nesiritide for Heart Failure
   c. Laetrile
   d. Outpatient L-Dopa (also see #3.k)
   e. Investigational or Experimental Drugs
   f. Placebos
   g. Outpatient Prescription Drugs
   h. Medications for the Treatment of Sexual Dysfunction
   i. Medications for Elective Enhancement
   j. Drugs Included in the CMS Self-Administered Drug Exclusion List
   k. Off-Label/Unlabeled Drug Use

5. Other Examples of Specific Drugs/Medications
   a. Botulinum Toxin for the Treatment of Migraine Headaches
   b. Crysvita® (burosumab-twza)
   c. Denosumab (Xgeva® , Prolia®)
   d. Erythropoietin
   e. Inflixi
timab (Remicade® , Inflectra™, Renflexis™)
   f. Luxturna™ (voretigene neparvovec)
   g. Onpattro™ (patisiran)
   h. Primacor® (milrinone) – Use in Home Setting
   i. Radicava™ (edaravone)
   j. Rituxan® (rituximab) for Non-Chemotherapeutic Indications
   k. Sodium Hyaluronate Injections for Osteoarthritis of Knee
   l. Soliris® (eculizumab)
   m. Spinraza™ (nusinersen)
   n. Testopel® (testosterone pellet)
   o. Ultomiris® (ravulizumab)
   p. Zolgensma® (Onasemnogene abeparvovec-Xioi)

6. Review at Launch (RAL)
I. COVERAGE

Coverage Statement: Outpatient/Part B medications/drugs are covered when Medicare coverage criteria are met.

DME Face to Face Requirement: Effective July 1, 2013, Section 6407 of the Affordable Care Act (ACA) established a face-to-face encounter requirement for certain items of DME (including implantable infusion pumps; implantable programmable infusion pump; external ambulatory infusion pump and nebulizers). For DME Face to Face Requirement information, refer to the Durable Medical Equipment (DME), Prosthetics, Corrective Appliances/Orthotics (Non-Foot Orthotics) and Medical Supplies Grid.

Guidelines/Notes:
1. Outpatient Medications/Drug
   a. Part B Medications/Drug

      Outpatient (Part B) medications/drugs, in accordance with Medicare coverage criteria, are covered when furnished “incident” to a physician service for drugs that are “not usually self-administered by the patient”. (See Section II for definition of “Not Usually Self-Administered by the Patient”)

      Coverage is usually limited to drugs or biologicals administered by infusion or injection. However, if the injection is generally self-administered (e.g., Immitrex), it is not covered under Part B. Despite the general limitation on coverage for outpatient drugs under Part B, some self-administered medications/drugs are also covered.

      Refer to Guideline 3 and Guideline 4 below for examples of covered and noncovered medications/drugs.

      For Medicare’s detailed coverage criteria for medications/drugs under Part B, refer to the Medicare Benefit Policy Manual, Chapter 15, §50 - Drugs and Biologicals. (Accessed October 8, 2018)

   b. Part D Medications/Drugs

      A Part D covered drug are available only by prescription, approved by the Food and Drug Administration (FDA), used and sold in the United States, and used for a medically accepted indication. A drug for which coverage is available under Part A or Part B, as it is being “prescribed and dispensed or administered” with respect to the individual, is excluded from the definition of a Part D drug and, therefore, cannot be included in Part D basic coverage. CMS interprets this to mean that if payment could be available under Part A or Part B to the individual for such drug, then it will not be covered under Part D.

      Some members may have coverage for Part D drugs under UnitedHealthcare. Refer to the Member’s Pharmacy Booklet or contact the Prescription Solutions Customer Service
Department to determine coverage eligibility for Part D prescription drug plan benefit.

For Medicare’s detailed coverage information for medications/drugs under Part D, refer to the Medicare Prescription Drug Benefit Manual, Chapter 6, §10 - Definition of Part D Drugs. (Accessed October 8, 2018)

c. Part B vs Part D Medications/Drugs

For Part B vs Part D medications/drugs guidelines, refer to the specific medications listed under Guideline 3 below.

2. Unlabeled Use of a Part B Drug

Unlabeled use of a drug may be covered only if a UnitedHealthcare Medical Director or his/her designee determines the use to be medically accepted, taking into consideration the major drug compendia. For the list of the major drug compendia, see the Medicare Benefit Policy Manual, Chapter 15, §50.4.2 - Unlabeled Use of Drug. (Accessed October 8, 2018)

In the case of drugs used in anti-cancer chemotherapeutic regimen, see the Coverage Summary for Chemotherapy, and Associated Drugs and Treatments.

Notes:

- The above information is for determining coverage for the unlabeled use of medication covered under Part B only, not Part D. Refer to the Member’s Pharmacy Booklet or contact the Prescription Solutions Customer Service Department for further information on Part D coverage, if any.

- A prescription drug is a Part D drug only if it is for a medically accepted indication as defined in the Medicaid statute. This definition includes uses supported by a citation included, or approved for inclusion, in one of three compendia. These are:
  - American Hospital Formulary Service Drug Information
  - United States Pharmacopeia-Drug Information (or its successor publications)
  - DRUGDEX Information System

Based on this statutory definition, indications that are supported in peer-reviewed medical literature, but not yet reflected in one of the compendia, are not “medically accepted.” Therefore, the use of a drug for such indications would not meet the definition of a Part D Drug.

See the Medicare Prescription Drug Benefit Manual Chapter 6, §10.6 - Medically Accepted Indication. (Accessed October 8, 2018)

- Definition of Compendium: Effective January 1, 2010, CMS revised the definition of “compendium” in the Medicare Benefit Policy Manual, Chapter 15, §50.4.5, to include this public transparency requirement. In this revised definition, a compendium:
  1. Includes a summary of the pharmacologic characteristics of each drug or biological and may include information on dosage, as well as recommended or endorsed uses in specific diseases;
  2. Is indexed by drug or biological; and
  3. Has a publicly transparent process for evaluating therapies and for identifying potential conflicts of interests.

Refer to the Medicare Benefit Policy Manual, Chapter 15, §50 - Drugs and Biologicals §50.4.5.1.A. (Accessed October 8, 2018)

3. Examples medications/drugs that are covered under Part B include, but not limited to:

a. Durable Medical Equipment (DME) Supply Drugs
Payment may be made for supplies that are necessary for the effective use of durable medical equipment. This includes drugs and biologicals which must be put directly into the equipment in order to achieve the therapeutic benefit of the durable medical equipment or to assure the proper functioning of the equipment. See the Medicare Benefit Policy Manual, Chapter 15, §110.3 - Coverage of Supplies and Accessories. (Accessed October 8, 2018)

Part B vs Part D Guideline:

1) **Nebulizer Inhalation Drugs (e.g., albuterol sulfate, ipratropium bromide):** Certain inhalation drugs are generally covered under Part B when used with a nebulizer in the home. These drugs would not be covered under Part D for use with a nebulizer. However, if these drugs were delivered with a metered dose inhaler or other non-nebulized administration, they would be Part D drugs.

In the case of a member in a hospital, or a SNF bed, (1) who does not have Part A coverage, (2) whose Part A coverage for the stay has run out or (3) whose stay is noncovered - infusible DME supply drugs are not covered under Part B because the law limits coverage under Part B’s DME benefit to those items that are furnished for use in a patient’s home, and specifies that a hospital or SNF cannot be considered the member’s “home” for this purpose. In this case, coverage for the drugs would be available under Part D.

In addition to a hospital, a SNF or a distinct part SNF, the following facilities cannot be considered a home for purposes of receiving the Medicare DME benefit:

- a nursing home that is dually-certified as both a Medicare SNF and a Medicaid nursing facility (NF);
- a Medicaid-only NF that primarily furnishes skilled care;
- a non-participating nursing home (i.e., neither Medicare or Medicaid) that provides primarily skilled care; and
- an institution which has a distinct part SNF and which also primarily furnishes skilled care.

For the list of nebulizer drugs covered under Part B, refer to the DME MAC LCD for Nebulizers (L33370). Compliance with these policies is required where applicable. (Accessed July 2, 2019)

2) **Infusion Pump Medications (e.g., some chemotherapeutic agents):**

In general, the supplier would bill Part B if the drug was administered using an infusion pump and bill the Part D plan for infusion using other methods (e.g., IV push). While professional services and supplies related to the administration of the infused drug are not payable under Part D, some coverage may be available under Part A or B home health benefits, under Medicaid, or from secondary commercial health benefits.

As a rule, drugs infused using an implantable pump would be covered under Part B. Drugs infused in the home using an external pump are covered under Part B if they are included under the local coverage policy of the applicable Medicare DME MAC.

In the case of a member in a hospital, or a SNF bed, (1) who does not have Part A coverage, (2) whose Part A coverage for the stay has run out or (3) whose stay is noncovered infusible DME supply drugs are not covered under Part B because the law limits coverage under Part B’s DME benefit to those items that are furnished for use in a patient’s home, and specifies that a hospital or SNF cannot be considered the member’s “home” for this purpose. In this
case, coverage for the drugs would be available under Part D.

In addition to a hospital, a SNF or a distinct part SNF, the following facilities cannot be considered a home for purposes of receiving the Medicare DME benefit:

- a nursing home that is dually-certified as both a Medicare SNF and a Medicaid nursing facility (NF);
- a Medicaid-only NF that primarily furnishes skilled care;
- a non-participating nursing home (i.e., neither Medicare or Medicaid) that provides primarily skilled care; and
- an institution which has a distinct part SNF and which also primarily furnishes skilled care.


b. Immunosuppressive Drugs

Imunosuppressive drug therapy following a Medicare covered organ transplant is covered.

Note: If a supplier of the immunosuppressive drugs has not determined (or does not have documentation on file to support a determination) that either the member beneficiary did not receive an organ transplant or that the beneficiary was not enrolled in Medicare Part A as of the date of the transplant, then the supplier may not, with respect to furnishing an immunosuppressive drug: 1) bill Medicare, 2) bill or collect any amount from the beneficiary, or 3) issue an Advance Beneficiary Notice (ABN) to the beneficiary. See the Medicare Claims Processing Manual, Chapter 17, §80.3 - Billing for Immunosuppressive Drugs. (Accessed October 8, 2018)

Covered drugs include those immunosuppressive drugs that have been specifically labeled as such and approved for marketing by the FDA. (This is an exception to the standing drug policy which permits coverage of FDA approved drugs for nonlabeled uses, where such uses are found to be reasonable and necessary in an individual case.)

Immunosuppressive drugs are substances that suppress or interfere with normal immune responses. They are used in controlling autoimmune diseases and in enhancing the chances for survival of foreign-tissue grafts and transplants.

Examples of FDA-approved immunosuppressive drugs include, but are not limited to:
- Sandimmune (cyclosporine), Sandoz Pharmaceutical;
- Imuran (azathioprine), Burroughs Wellcome;
- Atgam (antithymocyte globulin), Upjohn;
- Orthoclone OKT3 (Muromonab-CD3), Ortho Pharmaceutical;
- Prograf (tacrolimus), Fujisawa USA, Inc;
- Celicept (mycophenolate mofetil), Roche Laboratories;
- Daclizumab (Zenapax);
- Cyclophosphamide (Cytoxan);
- Prednisone and Prednisolone

Notes:
- Prescription drugs, such as prednisone, used in conjunction with immunosuppressive drugs as part of a therapeutic regimen are covered as reflected in FDA approved labeling for
Immunosuppressive drugs. Therapeutic regimen is a combination of drugs which has been clinically recognized for the treatment of a specific type of disorder or to treat toxicities or side effects of drugs which are used at different times following an approved transplant.

- Immunosuppressive drugs for organ transplants are covered under Part B coverage except when furnished during an inpatient stay or upon discharge from the hospital, then the drugs are covered as Part A.
- CMS expects contractors to keep informed of FDA additions to the list of the immunosuppressive drugs.
- Members may have additional coverage for immunosuppressive drugs under the Part D Prescription Drug Plan which are not covered in this benefit interpretation policy. Refer to the Member’s Pharmacy Booklet or contact the Prescription Solutions Customer Services Department to determine coverage eligibility for prescription drug plan benefit.

See the Medicare Benefit Policy Manual, Chapter 15, §50.5.1 - Immunosuppressive Drugs. (Accessed October 8, 2018)

**Part B vs Part D Guideline:**

Part B would be billed if the individual had a Medicare-covered transplant; otherwise, the Part D plan would be billed.

Pharmacists would bill Part B or the individual’s Part D plan based on information received from the individual or the Part D plan. Part B would be billed if the individual had a Medicare-covered transplant; otherwise, the Part D plan would be billed. (Part D plan eligibility systems could contain a marker for members who had a Medicare covered transplant. This information could come from a question included on the Part D sponsor’s enrollment or coordination of benefit (COB) survey form.

In determining whether to pay for an immunosuppressive drug under Part D, it would not be appropriate for a Part D sponsor to institute a general policy of requiring a Part B claim rejection, as a substitute for maintaining information on transplant status and paying claims based on that information. Such a policy would be disruptive to beneficiaries and pharmacies and would unnecessarily increase Part B contractor costs. Instead a prior authorization requirement would be appropriate.


Also see the Coverage Summary for Transplants: Organ and Tissue Transplants.

c. **Hemophilia Clotting Factors**

Hemophilia clotting factors are covered for hemophilia patients competent to use such factors to control bleeding without medical supervision of factors and items related to the administration of such factors. See the Coverage Summary for Blood, Blood Products and Related Procedures and Drugs.

**Part B vs Part D Guideline:** Hemophilia blood clotting factors would not be a Part D benefit because of the Part B coverage. See the Medicare Prescription Drug Benefit Manual, Chapter 6, Appendix C - Medicare Part B versus Part D Coverage Issues. (Accessed October 8, 2018)

d. **Oral Anti-Cancer Drugs and Oral Anti-Emetics**

Oral anti-cancer drugs and oral anti-nausea (anti-emetic) drugs are covered when criteria are met.
See the Coverage Summary for Chemotherapy, and Associated Drugs and Treatments.

Part B vs Part D Guideline:

Certain oral chemotherapy agents used in cancer treatment for which there is an infusible version of the drug.

- Pharmacists would need to determine the reason for treatment. If related to cancer treatment, Part B would be billed; otherwise, the Part D plan should be billed.
- To the extent that a Part B-covered oral anti-cancer drug has no other medically accepted indication besides cancer treatment, Part D sponsors should not include these drugs on their formularies because of Part B coverage. For the drugs that have other medically accepted indications, prior authorization programs or other mechanisms to obtain diagnostic information could be used to ensure appropriate payment.

Oral anti-emetics used in cancer treatment as a full replacement for intravenous treatment.

- Pharmacists would need to determine the reason for treatment. If both related to cancer treatment and a full replacement for intravenous administration within 48 hours of cancer treatment, Part B would be billed; otherwise, the Part D plan should be billed. NOTE: In order to receive Part B payment, CMS currently requires that the prescribing physician indicate on the prescription that the oral anti-emetic is being used “as a full therapeutic replacement for an intravenous anti-emetic drug as part of a cancer chemotherapeutic regimen.”
- If based on a prior authorization program or other mechanism to obtain diagnostic information, a Part D sponsor determined that a) a Part B-covered oral anti-emetic was being billed, and b) the drug was being furnished in the context of cancer treatment for use within 48 hours of cancer treatment, the Part D sponsor should deny payment. Such drugs dispensed for use after the 48-hour period, or any oral anti-emetic prescribed for conditions other than the effects of cancer treatment, would be Part D drugs.


e. Immunizations

Immunizations (e.g., pneumococcal vaccine, Hepatitis B vaccine, and influenza vaccine) are covered when criteria are met.

Refer to the Coverage Summary for Preventive Health Services and Procedures for coverage criteria.

Part B vs Part D Guideline:

For Hepatitis B vaccine, physicians would need to determine the level of risk of the individual. If the individual is at high or intermediate risk, Part B would be billed. For all other individuals, prior authorization programs could be used to ensure appropriate level of risk.

Pneumococcal and influenza vaccines would not be covered under Part D because of Part B coverage.


f. Antigens/Antihistamines

Antigens/antihistamines are covered when criteria are met. These are prepared by a physician (usually an allergist) for a specific patient. The physician or physician’s nurse generally
administrers them in the physician’s office. In some cases the physician prepares antigens and furnishes them to a patient who has been taught to self-administer them at home. Refer to the Coverage Summary for Allergy Testing and Allergy Immunotherapy.

**Part B vs Part D Guideline:** Antigens would not be a Part D benefit because of the Part B coverage. See the Medicare Prescription Drug Benefit Manual, Chapter 6, Appendix C - Medicare Part B versus Part D Coverage Issues. (Accessed October 8, 2018)

g. **Parenteral Nutrition**

Parenteral nutrition, including Intralaytic Parenteral Nutrition (IDPN), is covered under the prosthetic benefit when criteria are met. See the Coverage Summary for Nutritional Therapy: Enteral and Parenteral Nutritional Therapy for coverage criteria.

**Part B vs Part D Guideline:** If the therapy was being provided because of a non-functioning digestive tract, Part B would be billed; if not, this would be a Part D drug. See the Medicare Prescription Drug Benefit Manual, Chapter 6, Appendix C - Medicare Part B versus Part D Coverage Issues. (Accessed October 4, 2018)

h. **Intravenous Immune Globulin (IVIG)**

1) **Intravenous immune globulin (IVIG) in the Home**

Intravenous immune globulin (IVIG) for the treatment of primary immune deficiency diseases is covered in the home under Part B if all of the following criteria are met:

- It is an approved pooled plasma derivative for the treatment of primary immune deficiency disease; and
- The patient has a diagnosis of primary immune deficiency disease (ICD-9 codes 279.04, 279.05, 279.06, 279.12, and 279.2 or ICD-10-CM codes D80.0, D80.5, D81.0, D81.1, D81.2, D81.6, D81.7, D81.89, D81.9, D82.0, D83.0, D83.2, D83.8, or D83.9 if only an unspecified diagnosis is necessary); and
- The IVIG is administered in the home; and
- The treating physician has determined that administration of the IVIG in the patient’s home is medically appropriate.


**Part B vs Part D Guideline:** Part B coverage for IVIG in the home is for individuals whose diagnosis is primary immune deficiency disease. Part D would provide coverage for IVIG in the home for all other medically accepted indications. Prior authorization requirements could be used to ensure appropriate payment in accordance with the Part D sponsor’s medical necessity criteria. It would not be appropriate to routinely require a rejection of a claim under Part B before processing a Part D claim. Such a policy would be disruptive to beneficiaries and pharmacies and would unnecessarily increase Part B contractor cost.

The supplier would bill Part B if the diagnosis is primary immune deficiency disease. IVIG provided in the home for other diagnoses would be a Part D benefit. As discussed above, it would not be appropriate, as a general rule, for Part D sponsors to require a rejection of a claim under Part B before processing a Part D claim. Prior authorization programs could be used to ensure medical necessity in accordance with the Part D sponsor’s policy.
2) Treatment of Autoimmune Mucocutaneous Blistering Diseases

IVIg is covered for the treatment of biopsy-proven: (1) Pemphigus Vulgaris, (2) Pemphigus Foliaceus, (3) Bullous Pemphigoid, (4) Mucous Membrane Pemphigoid (a.k.a., Cicatricial Pemphigoid), and (5) Epidermolysis Bullosa Acquisita.

For more specific coverage guideline, see the NCD for Intravenous Immune Globulin for the Treatment of Autoimmune Mucocutaneous Blistering Diseases (250.3). (Accessed October 4, 2018)

Local Coverage Determinations (LCDs)/Local Coverage Articles (LCAs) exist for IVIG and compliance with these policies is required. For state-specific LCDs/LCAs, see the LCD Availability Grid (Attachment B).

3) Other Indications

- Medicare does not have an NCD for other indications other than the ones outlined on #1 and #2 above.
- Local Coverage Determinations (LCDs)/Local Coverage Article (LCAs) exist for all 50 states and compliance with these policies is required where applicable. For state-specific LCDs/LCAs, refer to the LCD Availability Grid (Attachment B).
- Committee approval date: October 16, 2018
- Accessed August 12, 2019

i. Intravenous Iron Therapy

Intravenous iron therapy (sodium ferric gluconate complex in sucrose injection) is covered as a first line treatment of iron deficiency anemia when furnished intravenously to patients undergoing chronic hemodialysis who are receiving supplemental erythropoietin therapy. For more specific information; see the NCD for Intravenous Iron Therapy (110.10). (Accessed October 4, 2018)

j. Injectable Drugs for the Treatment of Osteoporosis

Injectable drugs for the treatment of osteoporosis when provided by the home health agency and the following criteria are met:

- The member is unable to learn the skills needed to self-administer the drug, or is otherwise physically or mentally incapable of administering the drug, and that her family or caregiver are unable or unwilling to administered the drug, as documented by the home health agency, and
- The member sustained a bone fracture that a physician certifies was related to (post-menopausal) osteoporosis; and
- The member is unable to learn the skills needed to self-administer the drug, or is otherwise physically or mentally incapable of administering the drug, and that her family or caregiver are unable or unwilling to administered the drug, as documented by the home health agency,
and

- The member is homebound as defined in the Section II (Definitions) below.

See the Medicare Benefit Policy Manual Chapter 7, §50.4.3 - Covered Osteoporosis Drugs. (Accessed October 4, 2018)

Also see the Coverage Summary for Home Health Services and Home Health Visits.

Also see the CMS Tips to Partners on Osteoporosis Drugs at http://www.cms.gov/Outreach-and-Education/Outreach/Partnerships/downloads/11315-P.pdf. (Accessed October 4, 2018)

k. L-Dopa

L-Dopa in the inpatient setting (hospital or SNF) is covered for patients with Parkinsonism that have concurrent disease(s), e.g., cardiovascular, gastrointestinal or neuropsychological.

For more specific information; see the NCD for L-Dopa (160.17). (Accessed October 4, 2018)

Also see Guideline 4.d.

l. Dimethyl Sulfoxide (DMSO)

Dimethyl Sulfoxide (DMSO) is covered only when reasonable and necessary for the treatment of interstitial cystitis. See the NCD for Dimethyl Sulfoxide (DMSO) (230.12). (Accessed October 4, 2018)

m. Dermal Injections for the Treatment of Facial Lipodystrophy Syndrome (LDS)

Effective for claims with dates of service on and after March 23, 2010, dermal injections for LDS are only reasonable and necessary using dermal fillers approved by the Food and Drug Administration (FDA) for this purpose, and then only in HIV-infected beneficiaries when LDS caused by antiretroviral HIV treatment is a significant contributor to their depression. See the NCD for Dermal Injections for the Treatment of Facial Lipodystrophy Syndrome (LDS) (250.5). (Accessed October 4, 2018)

n. Drugs Treated as Hospital Outpatient Supplies

In certain circumstances, Medicare pays for drugs that may be considered usually self-administered by the patient when such drugs function as supplies. This is the case when the drugs provided are an integral component of a procedure or are directly related to it, i.e., when they facilitate the performance of or recovery from a particular procedure. Except for the applicable copayment, hospitals may not bill beneficiaries for these types of drugs because their costs, as supplies, are packaged into the payment for the procedure with which they are used. Listed below are examples of when drugs are treated as supplies and hospitals should bill Medicare for the drug as a supply and should not separately bill the member.

- Sedatives administered to a patient while he or she is in the preoperative area being prepared for a procedure
- Mydriatic drops instilled into the eye to dilate the pupils, anti-inflammatory drops, antibiotic drops/ointments, and ocular hypotensives that are administered to a patient immediately before, during, or immediately following an ophthalmic procedure. This does not refer to the patient’s eye drops that the patient uses pre-and postoperatively.
- Barium or low osmolar contrast media provided integral to a diagnostic imaging procedure
- Topical solution used with photodynamic therapy furnished at the hospital to treat nonhyperkeratotic actinic keratosis lesions of the face or scalp
• Antibiotic ointments such as bacitracin, placed on a wound or surgical incision at the completion of a procedure

The following are examples of when a drug is not directly related or integral to a procedure, and does not facilitate the performance of or recovery from a procedure. Therefore the drug is not considered a packaged supply. In many of these cases the drug itself is the treatment instead of being integral or directly related to the procedure, or facilitating the performance of or recovery from a particular procedure.

• Drugs given to a patient for his or her continued use at home after leaving the hospital
• Oral pain medication given to an outpatient who develops a headache while receiving chemotherapy administration treatment
• Daily routine insulin or hypertension medication given preoperatively to a patient
• A fentanyl patch or oral pain medication such as hydrocodone, given to an outpatient presenting with pain
• A laxative suppository for constipation while the patient waits to receive an unrelated X-ray

These two lists of examples may serve to guide hospitals in deciding which drugs are supplies packaged as a part of a procedure, and thus may be billed under Part B. Hospitals should follow CMS’ guidance for billing drugs that are packaged and paid as supplies, reporting coded and uncoded drugs with their charges under the revenue code associated with the cost center under which the hospital accumulates the costs for the drugs.

See the Medicare Benefit Policy Manual, Chapter 15, §50.2 - Determining Self-Administration of Drug or Biological, M – Drugs Treated as Hospital Outpatient Supplies. (Accessed October 4, 2018)

o. Hereditary Angioedema (HAE) Treatment (HCPCS codes J0596, J0597, J0598 and J1290)
• Medicare does not have a National Coverage Determination (NCD) for Hereditary Angioedema (HAE) treatment.
• Local Coverage Determinations (LCDs)/Local Coverage Articles (LCAs) do not exist at this time.
• For coverage guidelines, refer to the UnitedHealthcare Commercial Medical Benefit Drug Policy for Hereditary Angioedema (HAE), Treatment and Prophylaxis. (IMPORTANT NOTE: After searching the Medicare Coverage Database, if no state LCD/LCA is found, then use the above referenced policy.)
• Committee approval date: August 20, 2019
• Accessed August 12, 2019

4. Examples of medications/drugs that are not covered:
   a. Vitamin B12 Injections

Vitamin B12 injections to strengthen tendons, ligaments, etc., of the foot are not covered under Medicare because (1) there is no evidence that vitamin B12 injections are effective for the purpose of strengthening weakened tendons and ligaments, and (2) this is nonsurgical treatment under the subluxation exclusion. Accordingly, Vitamin B12 injections are not considered reasonable and necessary.

See the NCD for Vitamin B12 Injections to Strengthen Tendons, etc., of the Foot (150.6). (Accessed October 4, 2018)
b. **Nesiritide for Heart Failure**

Nesiritide for heart failure is not covered. See the *NCD for Nesiritide for Treatment of Heart Failure Patients (200.1)*. (Accessed October 4, 2018)

c. **Laetrile**

Laetrile and the other drugs called by the various terms mentioned below, used primarily in the treatment or control of cancer, are not covered. Although the terms "Laetrile," "laetrile," "amygdalin," "Sarcarinase," "vitamin B-17," and "nitriloside" have been used interchangeably, the chemical identity of the substances to which these terms refer has varied.

*For more specific information, see the NCD for Laetrile and Related Substances (30.7).* (Accessed October 4, 2018)

d. **Outpatient L-Dopa**

Outpatient L-Dopa is not covered because it is a self–injectable medication, unless the member has Part D pharmacy benefit coverage. See the *NCD for L-Dopa (160.17)* for more specific information. (Accessed October 4, 2018)

*Also see Guideline 3.k.*

e. **Investigational or Experimental Drugs** are not covered. See the *Coverage Summary for Experimental Procedures and Items, Investigational Devices and Clinical Trials.*

f. **Placebos** are not covered.

g. **Outpatient Prescription Drugs**

Outpatient prescription drugs are not covered except those medications/drugs covered under the Member’s Part D Prescription Drug Plan benefit.

*Refer to the Member’s Pharmacy Program booklet or contact the Prescription Solutions Customer Services Department to determine coverage eligibility for Part D Prescription Drug benefit.*

h. **Medications for the Treatment of Sexual Dysfunction**

Medications for the treatment of sexual dysfunction including erectile dysfunction, impotence, anorgasm or hypoorgasm are not covered. See the *Coverage Summary for Impotence Treatment.*

i. **Medications for Elective Enhancement**

Medications for elective enhancement, such as those used for weight loss, hair growth, sexual performance, athletic performance, cosmetic purposes, anti-aging, and mental performance are not covered. See the *Coverage Summary for Cosmetic and Reconstructive Procedures.*

j. **Drugs Included in the CMS Self-Administered Drug Exclusion List**

Drugs included in the CMS Self-administered Drug Exclusion List are not covered.

*Notes:*

- **Self-Administered Drug (SAD) Exclusion List Report:** Local Contractors have self-administered drugs exclusion lists. Compliance with these lists is required where applicable. See the *Self-Administered Drug (SAD) Exclusion List Report.* (Accessed August 16, 2019)

- **PCSK9 Inhibitors:** PCSK9 Inhibitors, *i.e.*, Praluent™ (alirocumab) and Repatha™
(evolocumab) are considered self-administered drugs and are not covered under the Part B medical benefit. Refer to the Member’s Pharmacy Program booklet or contact the Prescription Solutions Customer Service Department to determine coverage eligibility for these drugs under the Part D Prescription Drug benefit.

k. Off-Label/Unlabeled Drug Use
Off-Label/unlabeled drug use is not covered unless criteria are met; see Guideline 2 above for coverage criteria and guidelines.

5. Other Examples of Specific Drugs/Medications – See Attachment A for the list of drugs/medications and coverage guidelines.

6. Review at Launch (RAL)
A pre-service organization determination is highly recommended for certain Part B medications (as defined above):

- that are new to the market;
- that have not yet undergone review by UnitedHealthcare; and
- for which a utilization management strategy has not been established.

These medications, referred to herein as RAL medications, are identified in Attachment A: Guideline 5 – Other Examples of Specific Drugs/Medications. Upon receipt of a pre-service organization determination, RAL medications will be reviewed against National Coverage Determinations (NCDs), Local Coverage Determinations (LCDs), and/or against available clinical evidence, which includes applicable Medical Benefit Drug Policies when there is no LCD uniformity or there is no clear Medicare guidance.

Providers are strongly encouraged to seek a pre-service organization determination for any RAL medication that has been identified in Attachment A: Guideline 5 – Other Examples of Specific Drugs/Medications. This will help to avoid gaps in coverage in the event that a prior authorization program becomes effective at a later date. If a provider believes an item or service may not be covered, or could only be covered under specific conditions, the appropriate process is to request a pre-service organization determination.

7. Step Therapy Program
Effective January 1, 2019, certain classes of medical benefit injectables covered under Medicare Part B will include non-preferred therapies that require prior authorization. Each class of medical injectables will include preferred therapies that do not required prior authorization. Prior authorization for a non-preferred therapy will generally require history of use of a preferred therapy within the same medical benefit injectable class, among other criteria.

This prior authorization requirement will apply to some, but not all, Medicare Advantage Plans.

A medical injectable is subject to step therapy when it is listed on Attachment A: Guideline 5 – Other Examples of Specific Drugs/Medications and a notation to refer to the Medicare Part B Step Therapy Program is provided in the Step Therapy column.

8. Other Specific Medications not listed above

b. If there are no applicable LCDs or LCAs found, refer to Guideline 1 and Guideline 2 of this
Coverage summary for Medicare guidelines for covered Part B medications and the use of compendia and evidence-based medical literature in determining coverage for specific medication.

9. **Shortage of Leucovorin (J0640)**

   There is currently a nationwide shortage of injectable racemic leucovorin, available only as a generic drug and only from two manufacturers in the US (Bedford Laboratories and Teva Pharmaceuticals). According to the FDA, the shortages are due to manufacturing delays; however, the American Society of Health-System Pharmacists (ASHP) reports that Bedford cannot provide a reason for the shortage. Information on current availability of specific vial sizes and expected release dates for others can be found on the [FDA website](https://www.fda.gov) and [American Society of Health-System Pharmacists (ASHP) Current Drug Shortage Bulletin](https://www.ashp.org). (Accessed October 4, 2018)

   For patients affected by this shortage, the UnitedHealthcare will consider levoleukovorin (J0641) as an alternative for leucovorin (J0640).

   Refer to *Coverage Summary for Chemotherapy, and Associated Drugs and Treatments* for information on Chemotherapy Drugs.

### II. DEFINITIONS

**FDA Approved Drug:** A drug that has received final marketing approval by the Food and Drug Administration (FDA) and as a part of its labeling contains its recommended uses and dosages as well as adverse reactions and recommended precautions in using it. *Medicare Benefit Policy Manual, Chapter 15, §50.4.2 - Unlabeled Use of Drug.* (Accessed October 4, 2018)

**Hemophilia:** For purposes of Medicare Part B coverage, hemophilia encompasses the following conditions:
- Factor VIII deficiency (classic hemophilia);
- Factor IX deficiency (also termed plasma thromboplastin component (PTC) or Christmas factor deficiency); and
- Von Willebrand’s disease


**Homebound:** The restricted ability of a member, due to an illness or injury, to leave home without the assistance of another or the aid of a supportive device (such as crutches, a cane, a wheelchair, or a walker), or if leaving the home is medically contraindicated. A person does not need to be bedridden to be confined to the home. However, the physical condition must be such that there exists a normal inability to leave home and leaving requires a considerable and taxing effort.

   If the patient does in fact leave the home, the patient may nevertheless be considered homebound if the absences from the home are infrequent or for periods of relatively short duration, or are attributable to the need to receive health care treatment.

   Any other absence of an individual from the home shall not so disqualify an individual if the absence is of infrequent or of relatively short duration. For purposes of the preceding sentence, any absence for the purpose of attending a religious service shall be deemed to be an absence of infrequent or short duration.

   Generally speaking, a patient will be considered to be homebound if they have a condition due to an illness or injury that restricts their ability to leave their place of residence except with the aid of: supportive devices such as crutches, canes, wheelchairs, and walkers; the use of special transportation; or the assistance of another person; or if leaving home is medically contraindicated.

Not Usually Self-Administered by the Patient (as defined by Medicare):

a. **Administered**: The term “administered” refers only to the physical process by which the drug enters the patient’s body. It does not refer to whether the process is supervised by a medical professional (for example, to observe proper technique or side-effects of the drug). Injectable drugs (including intravenous drugs) are typically eligible for inclusion under the “incident to” benefit. With limited exclusions, other routes of administration including, but not limited to, oral drugs, suppositories, topical medications are all considered to be usually self-administered by the patient.

b. **Usually**: For the purposes of applying this exclusion, the term "usually" means more than 50 percent of the time for all Medicare beneficiaries who use the drug. Therefore, if a drug is self-administered by more than 50 percent of Medicare beneficiaries, the drug is excluded from coverage and you may not make any Medicare payment for it.

c. **By the patient**: The term "by the patient" means Medicare beneficiaries as a collective whole. Include only the patients themselves and not other individuals (which do not include spouses, friends, or other caregivers).

*Medicare Benefit Policy Manual, Chapter 15, §50.2 Determining Self-Administration of Drug or Biological.* (Accessed October 4, 2018)

**Unlabeled Use of Drug**: A use that is not included as an indication of the drug’s label as approved by FDA. *Medicare Benefit Policy Manual, Chapter 15, §50.4.2 - Unlabeled Use of Drug.* (Accessed October 4, 2018)

### III. REFERENCES

See above

### IV. REVISION HISTORY

08/20/2019 **Guideline 3.o [Hereditary Angioedema (HAE) Treatment (HCPCS codes J0596, J0597, J0598 and J1290)]** (new to policy; )

- Added coverage guidelines (previously outlined in the UnitedHealthcare Medicare Advantage Coverage Summary titled *Genetic Testing*) to indicate:
  - Medicare does not have a National Coverage Determination (NCD) for Hereditary Angioedema (HAE) treatment
  - Local Coverage Determinations (LCDs)/Local Coverage Articles (LCAs) do not exist
  - Refer to the UnitedHealthcare Commercial Medical Benefit Drug Policy titled *Hereditary Angioedema (HAE), Treatment and Prophylaxis* for applicable coverage guidelines

**Guideline 5 (Other Examples of Specific Drugs/Medications)**

- Revised Attachment A - Guideline 5: Other Examples of Specific Drugs/Medications:
  - Updated reference links to reflect the most current LCDs for:
    - Erythropoietin
    - Rituxan® (rituximab) for Non-Chemotherapeutic Indications
    - Sodium Hyaluronate Injections for Osteoarthritis of Knee
  - Updated list of applicable CPT/HCPCS codes for Testopel® (testosterone pellet):
    - Added J3490
    - Removed S0189

**Attachments**

- Updated LCD Availability Grid to reflect the most current reference links
V. ATTACHMENT(S)

Attachment A

**Guideline 5 - Other Examples of Specific Drugs/Medications**

*IMPORTANT NOTE: After searching the *Medicare Coverage Database*, if no state LCD/LCA is found, then use the applicable referenced default policy below for coverage guidelines.*

<table>
<thead>
<tr>
<th>Drug/Medication</th>
<th>NCD, Medicare Manual, or LCDs/LCAs MAC and States</th>
<th>Default Policy for state with no LCDs</th>
<th>Individual Consideration (IC)</th>
<th>Review at Launch (RAL)</th>
<th>Step Therapy</th>
<th>UMBIC Approval Date</th>
<th>Accessed Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>a. Botulinum Toxin for the Treatment of Migraine Headaches</td>
<td>First Coast L33274 NGS L33646 CGS L33949 Wisconsin L34635 Noridian L35170 Noridian L35172</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Botulinum Toxins A and B</td>
<td>None</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>b. Crysvita® (burosumab-twza)</td>
<td>None</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Crysvita® (Burosumab-Twza)</td>
<td>Not Applicable (N/A)</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>c. Denosumab (Xgeva®, Prolia®)</td>
<td>First Coast L33270 CGS A56534 NGS L33394 Palmetto A55297</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Denosumab (Prolia® &amp; Xgeva®)</td>
<td>None</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>d. Erythropoietin</td>
<td>N/A</td>
<td>N/A</td>
<td>No</td>
<td>Yes</td>
<td>See the Medicare Part B Step Therapy Programs policy</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>Drug/Medication</td>
<td>NCD, Medicare Manual, or LCDs/LCAs MAC and States</td>
<td>Default Policy for state with no LCDs</td>
<td>Individual Consideration (IC)</td>
<td>Review at Launch (RAL)</td>
<td>Step Therapy</td>
<td>UMBIC Approval Date</td>
<td>Accessed Date</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>--------------------------------------------------</td>
<td>--------------------------------------</td>
<td>--------------------------------</td>
<td>------------------------</td>
<td>--------------</td>
<td>--------------------</td>
<td>---------------</td>
</tr>
<tr>
<td></td>
<td>applicable:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>First Coast</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>CGS L3456</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Wisconsin L34633</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>e. Infliximab (Remicade®, Inflectra™, Renflexis™)</td>
<td>Palmetto L35677 First Coast L33704 NGS L33394</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Infliximab (Remicade®, Inflectra™, Renflexis™)</td>
<td>None</td>
<td>No</td>
<td>Yes</td>
<td>See the Medicare Part B Step Therapy Programs policy</td>
<td>11/20/2018 08/12/2019</td>
</tr>
<tr>
<td>f. Luxturna™ (voretigene neparvovec)</td>
<td>Palmetto L37863</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Luxturna™ (Voretigene Neparvovec-Rzyl)</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>04/16/2019 08/12/2019</td>
<td></td>
</tr>
<tr>
<td>g. ONPATTRO™ (patisiran)</td>
<td>None</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Onpattro™ (Patisiran)</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>01/15/2019 08/12/2019</td>
<td></td>
</tr>
</tbody>
</table>
### Attachment A

**Guideline 5 - Other Examples of Specific Drugs/Medications**

*(IMPORTANT NOTE: After searching the Medicare Coverage Database, if no state LCD/LCA is found, then use the applicable referenced default policy below for coverage guidelines.)*

<table>
<thead>
<tr>
<th>Drug/Medication</th>
<th>NCD, Medicare Manual, or LCDs/LCAs MAC and States</th>
<th>Default Policy for state with no LCDs</th>
<th>Individual Consideration (IC)</th>
<th>Review at Launch (RAL)</th>
<th>Step Therapy</th>
<th>UMBIC Approval Date</th>
<th>Accessed Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>h. Primacor® (milrinone) - Use in Home Setting</td>
<td>DME MAC LCD for External Infusion Pumps L33794</td>
<td>All states have LCDs</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>i. Radicava™ (edaravone)</td>
<td>None</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Radicava™ (Edaravone)</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>j. Rituxan® (rituximab) for Non-Chemotherapeutic Indications</td>
<td>First Coast L33746 NGS L33394 Palmetto L35026</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Rituxan® (Rituximab)</td>
<td>Waldenstrom’s Macroglobulinemia</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>k. Sodium Hyaluronate Injections for Osteoarthritis of Knee</td>
<td>Palmetto L33432 Novitas L35427 Wisconsin A56157 NGS L33394 First Coast L33767</td>
<td>UnitedHealthcare Commercial Medical Policy for Sodium Hyaluronate</td>
<td>None</td>
<td>No</td>
<td>Yes</td>
<td>See the Medicare Part B Step Therapy Programs policy</td>
<td>11/20/2018</td>
</tr>
<tr>
<td>l. Spinraza™ (nusinersen)</td>
<td>Novitas L37682</td>
<td>UnitedHealthcare</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>11/20/2018</td>
<td>08/12/2019</td>
</tr>
</tbody>
</table>
### Guideline 5 - Other Examples of Specific Drugs/Medications

*IMPORTANT NOTE:* After searching the Medicare Coverage Database, if no state LCD/LCA is found, then use the applicable referenced default policy below for coverage guidelines.

<table>
<thead>
<tr>
<th>Drug/Medication</th>
<th>NCD, Medicare Manual, or LCDs/LCAs MAC and States</th>
<th>Default Policy for state with no LCDs</th>
<th>Individual Consideration (IC)</th>
<th>Review at Launch (RAL)</th>
<th>Step Therapy</th>
<th>UMBIC Approval Date</th>
<th>Accessed Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>m. Soliris® (eculizumab)</td>
<td>NGS L33394</td>
<td>Commercial Medical Benefit Drug Policy for Spinraza™ (Nusinersen)</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>03/19/2019</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>n. Testopel® (testosterone pellet)</td>
<td>First Coast L33412, Noridian L36569, L36538</td>
<td>First Coast L33412</td>
<td>None</td>
<td>No</td>
<td>No</td>
<td>08/20/2019</td>
<td>08/14/2019</td>
</tr>
<tr>
<td>o. Ultomiris® (ravulizumab)</td>
<td>None</td>
<td>UnitedHealthcare Commercial Medical Benefit Drug Policy for Complement Inhibitors (Soliris &amp; Ultomiris)</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>07/23/2019</td>
<td>08/12/2019</td>
</tr>
<tr>
<td>p. Zolgensma® (Onasemnogene abeparvovec-Xioi)</td>
<td>None</td>
<td>UnitedHealthcare Drug Policy for Zolgensma® (Onasemnogene abeparvovec-Xioi)</td>
<td>N/A</td>
<td>Yes</td>
<td>No</td>
<td>07/23/2019</td>
<td>08/12/2019</td>
</tr>
</tbody>
</table>

---

End of Attachment A
## Intravenous Immune Globulin (IVIG)

**Attachment B - LCD Availability Grid**

**CMS website accessed 08/12/2019**

<table>
<thead>
<tr>
<th>LCD ID</th>
<th>LCD Title</th>
<th>Contractor Type</th>
<th>Contractor</th>
<th>States</th>
</tr>
</thead>
<tbody>
<tr>
<td>L33610</td>
<td>Intravenous Immune Globulin</td>
<td>DME MAC</td>
<td>Noridian Healthcare Solutions, LLC (16013)</td>
<td>CT, DC, DE, MA, MD, ME, NH, NJ, NY, PA, RI, VT</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CGS Administrators, LLC (18003)</td>
<td>AL, AR, CO, FL, GA, LA, MS, NC, NM, OK, PR, SC, TN, TX, VA, VI, WV</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Noridian Healthcare Solutions, LLC (19003)</td>
<td>AK, AS, AZ, CA, GU, HI, IA, ID, KS, MO, MT, ND, NE, MP, NV, OR, SD, UT, WA, WY</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CGS Administrators (17013)</td>
<td>IL, IN, KY, MI, MN, OH, WI</td>
</tr>
<tr>
<td>L34007</td>
<td>Intravenous Immune Globulin</td>
<td>A and B MAC</td>
<td>First Coast Service Options, Inc.</td>
<td>FL, PR, VI</td>
</tr>
<tr>
<td>L34771</td>
<td>Immune Globulins</td>
<td>MAC Part A and B</td>
<td>Wisconsin Physicians Service Insurance Corporation</td>
<td>IA, IN, KS, MI, MO, NE</td>
</tr>
<tr>
<td>L34771</td>
<td>Immune Globulins</td>
<td>MAC Part A</td>
<td>Wisconsin Physicians Service Insurance Corporation</td>
<td>AK, AL, AR, AZ, CA, CO, CT, DE, FL, GA, HI, IA, ID, IL, IN, KS, KY, LA, MA, MD, ME, MI, MO, MS, MT, NC, ND, NE, NH, NJ, NM, NV, OH, OK, OR, PA, RI, SC, SD, TN, TX, UT, VA, WI, WV, WY</td>
</tr>
<tr>
<td>L33394</td>
<td>Drugs and Biologicals, Coverage of, for Label and Off-Label Uses</td>
<td>MAC Part A and B A and B MAC</td>
<td>National Government Services, Inc.</td>
<td>CT, IL, MA, ME MN, NH, NY, RI, VT, WI</td>
</tr>
<tr>
<td>L35093</td>
<td>Intravenous Immune Globulin (IVIG)</td>
<td>A and B MAC</td>
<td>Novitas Solutions, Inc.</td>
<td>CO, NM, OK, TX, AR, LA, MS DE, DC, MD, NJ, PA</td>
</tr>
<tr>
<td>L34074</td>
<td>Immune Globulin Intravenous (IVIg)</td>
<td>A and B MAC</td>
<td>Noridian Healthcare Solutions, LLC</td>
<td>AK, ID, OR, WA, AZ, MT, ND, SD, UT, WY</td>
</tr>
<tr>
<td>L34314</td>
<td>Immune Globulin Intravenous (IVIg)</td>
<td>A and B MAC</td>
<td>Noridian Healthcare Solutions, LLC</td>
<td>CA, AS, GU, HI, MP, NV</td>
</tr>
<tr>
<td>L34580</td>
<td>Intravenous Immunoglobulin (IVIG)</td>
<td>A and B MAC</td>
<td>Palmetto GBA</td>
<td>AL, GA, NC, SC, TN, VA, WV</td>
</tr>
<tr>
<td>L35891</td>
<td>Intravenous Immune Globulin</td>
<td>A and B MAC</td>
<td>CGS Administrators, LLC</td>
<td>KY, OH</td>
</tr>
</tbody>
</table>

### End of Attachment B

#### Addendum – Attachment A

**Part A and B MACs and States**

<table>
<thead>
<tr>
<th>Provider</th>
<th>States</th>
</tr>
</thead>
<tbody>
<tr>
<td>Palmetto</td>
<td>NC, SC, VA, WV and AL, GA, TN</td>
</tr>
<tr>
<td>Noridian</td>
<td>AK, AZ, ID, MT, ND, OR, SD, UT, WA, WY</td>
</tr>
<tr>
<td>Noridian</td>
<td>AS, CA, HI, NV, No. Mariana Islands</td>
</tr>
<tr>
<td>CGS</td>
<td>KY, OH</td>
</tr>
<tr>
<td>Wisconsin</td>
<td>AK, AL, AR, AZ, CA, CO, CT, DE, FL, GA, HI, IA, ID, IL, IN, KS, KY, LA, MA, MD, ME, MI, MO, MS, MT, NC, ND, NE, NH, NJ, NM, NV, OH, OK, OR, PA, RI, SC, SD, TN, TX, UT, VA, VT, WA, WI, WV, WY</td>
</tr>
</tbody>
</table>

<Back to Attachment A>
<table>
<thead>
<tr>
<th>DME MACs and States</th>
</tr>
</thead>
<tbody>
<tr>
<td>Novitas AR, CO, DE, LA, MD, MS, NJ, NM, OK, PA, TX</td>
</tr>
<tr>
<td>First Coast FL, PR, VI</td>
</tr>
<tr>
<td>NGS CT, IL, ME, MA, MN, NH, NY, RI, VT, WI</td>
</tr>
</tbody>
</table>
| Noridian (16013) CT, DC, DE, MA, MD, ME, NH, NJ, NY, PA, RI, VT | <Back to Attachment A>  
| CGS (18003) AL, AR, CO, FL, GA, LA, MS, NC, NM, OK, PR, SC, TN, TX, VA, VI, WV |  
| Noridian (19003) AK, AS, AZ, CA, GU, HI, IA, ID, KS, MO, MT, ND, NE, MP, NV, OR, SD, UT, WA, WY |  
| CGS (17013) IL, IN, KY, MI, MN, OH, WI |