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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

HYDROXYZINE HCL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Oral formulations only: Pruritus: Failure or clinically significant adverse effects to one of the following topical agents: betamethasone, hydrocortisone, triamcinolone, fluticasone, clobetasol, fluocinonide or fluocinolone. Anxiety: Failure or clinically significant adverse effects to one of the following: venlafaxine, buspirone, duloxetine or escitalopram. Oral and injectable formulations: All other FDA approved indications: Patient is continuing on this medication without adverse effects.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

HYDROXYZINE PAMOATE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Pruritus: Failure or clinically significant adverse effects to one of the following topical agents: betamethasone, hydrocortisone, triamcinolone, fluticasone, clobetasol, fluocinonide or fluocinolone. Anxiety: Failure or clinically significant adverse effects to one of the following: venlafaxine, buspirone, duloxetine or escitalopram. All other FDA approved indications: Patient is continuing on this medication without adverse effects.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ICLUSIG

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation of T315I mutation status. Acute Lymphoblastic Leukemia: Documentation of Philadelphia chromosome positive (Ph+) disease.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

T315I mutation-negative Chronic Myelogenous Leukemia: Failure of a trial of two tyrosine-kinase inhibitors (e.g., imatinib, nilotinib, dasatinib, bosutinib) used to treat CML, unless contraindicated or clinically significant adverse effects are experienced.





















## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

INLYTA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Thyroid carcinoma (i.e., Follicular carcinoma, Hurthle cell carcinoma, papillary carcinoma).

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Follicular carcinoma, Hurthle cell carcinoma, papillary carcinoma: Disease is unresectable, recurrent/persistent or metastatic. CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

INTERFERON BETA-1A (Avonex, Rebif)

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

INTERFERON BETA-1B (Betaseron, Extavia)

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

INTUNIV

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Attention Deficit Hyperactivity Disorder: Failure or clinically significant adverse effects to two of the following: dexamethylphenidate, methylphenidate or mixed amphetamine salts.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

JAKAFI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation of current platelet count and complete blood count (CBC). CONTINUATION OF THERAPY:  
Documentation of reduction in spleen volume or symptom improvement.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist or hematologist.

#### **Coverage Duration:**

Initial: 6 months. Reauthorization: Through the end of the Plan contract year.

#### **Other Criteria:**



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

JUXTAPID

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Repatha 420 mg (unless contraindicated).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

JYNARQUE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a nephrologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KADCYLA

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Kadcyla will be used as a single-agent therapy. Documentation that the patient has either received prior therapy for metastatic disease or developed disease recurrence during or within six months of completing adjuvant therapy.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Previously received trastuzumab and a taxane (e.g., paclitaxel, docetaxel), either separately or in combination.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KALYDECO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients with cystic fibrosis who are homozygous for the F508del mutation.

#### **Required Medical Information:**

Presence of one mutation in the CFTR gene that is responsive to ivacaftor as detected by an FDA-cleared cystic fibrosis mutation test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KAZANO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Tradjenta or Jentadueto AND Failure or clinically significant adverse effects to one of the following: Januvia, Janumet, Janumet XR.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KETOROLAC TROMETHAMINE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients with active peptic ulcer disease. Advanced renal impairment or at risk for renal failure due to volume depletion. Suspected or confirmed cerebrovascular bleeding, hemorrhagic diathesis, incomplete hemostasis and those at high risk for bleeding. Patient currently receiving aspirin or NSAIDs (Non-steroidal anti-inflammatory drugs). Patient currently receiving Probenecid or pentoxifylline.

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

5 days

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KEVZARA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a rheumatologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine, or auranofin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KINERET

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

RHEUMATOID ARTHRITIS: Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin AND Failure or clinically significant adverse effects to one of the following: Enbrel, Humira, Remicade, Cimzia, Simponi or Simponi Aria.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KISQALI (includes Kisqali Femara Co-Pack)

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Will be used in combination with an aromatase inhibitor (e.g., letrozole, anastrozole or exemestane) as initial endocrine-based therapy.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KOMBIGLYZE XR

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Tradjenta or Jentadueto AND Failure or clinically significant adverse effects to one of the following: Januvia, Janumet, Janumet XR.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KORLYM

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Pregnancy.

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KUVAN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response, demonstrated by a reduction of blood phenylalanine levels from baseline.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Initial: 2 months. Reauthorization: Through the end of the Plan contract year.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

KYNAMRO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Repatha 420 mg (unless contraindicated).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LATUDA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Schizophrenia: Failure or clinically significant adverse effects to two of the following generic atypical antipsychotics: risperidone, olanzapine, quetiapine, ziprasidone, aripiprazole.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LAZANDA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Age 18 or greater

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Through the end of the Plan contract year.

#### **Other Criteria:**

Patient is already taking and is tolerant to around-the-clock opioid therapy. Patients are considered opioid tolerant when taking another opioid daily for a week or longer (for example, at least 60 mg of oral morphine per day or an equianalgesic dose of another opioid).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LEMTRADA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to two of the following: Aubagio, Tecfidera, Gilenya, Avonex, Betaseron, Plegridy, Copaxone, Glatopa, Extavia or Rebif.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LENVIMA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Medullary thyroid carcinoma.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Renal Cell Carcinoma: Failure or clinically significant adverse effects to one of the following: Sutent, Nexavar, Votrient, Inlyta, Avastin in combination with Intron-A, Proleukin, Torisel AND Failure or clinically significant adverse effects to Opdivo or Cabometyx AND Must be used in combination with everolimus (Afinitor).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LEUKINE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Use Following Induction Chemotherapy in Acute Myelogenous Leukemia, Use in Mobilization and Following Transplantation of Autologous Peripheral Blood Progenitor Cells, Use in Myeloid Reconstitution After Autologous or Allogeneic Bone Marrow Transplantation: Failure or clinically significant adverse effects to Neupogen.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LIDODERM

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Diabetic peripheral neuropathy. Cancer-related neuropathic pain.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

---

### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LONSURF

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation that the patient does or does not have the KRAS wild type gene. CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following: 5-fluorouracil, capecitabine, oxaliplatin, irinotecan, Avastin, Cyramza, Zaltrap. If tumor expresses the KRAS wild type gene, failure or clinically significant adverse effects to Erbitux or Vectibix.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LOTROXEX

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Male patients.

#### **Required Medical Information:**

Female patient with irritable bowel symptoms persisting for at least 6 months.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LYNPARZA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Treatment of ovarian cancer: Mutations in the BRCA genes as detected by an FDA approved test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

LYNPARZA TABLET

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Treatment of ovarian cancer: Mutations in the BRCA genes as detected by an FDA approved test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

For maintenance therapy: Completed two or more platinum-based chemotherapy regimens and is in a complete or partial response.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MACRODANTIN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Urinary tract infectious disease, Acute treatment: Failure or clinically significant adverse effects to ONE of the following: sulfamethoxazole/trimethoprim or ciprofloxacin. Urinary tract infectious disease, Prophylaxis: Patient is continuing on this medication without adverse effects.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MAVYRET

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Treatment-experienced patients with both NS3/4A protease inhibitor and NS5A inhibitor.

#### **Required Medical Information:**

If cirrhosis is present, confirmation of Child-Pugh A status. Refer to the document, "Recommendations for Testing, Managing, and Treating Hepatitis C," by AASLD-IDSAs available at <http://www.hcvguidelines.org> for drug regimen and duration of treatment based on genotype, treatment status, previous drug regimens used, past medical history and comorbidities.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a gastroenterologist, hepatologist or infectious disease physician.

#### **Coverage Duration:**

8 to 16 weeks based on genotype, cirrhosis status, prior treatment regimen.

#### **Other Criteria:**

If patient has been previously treated with an HCV regimen containing NS5A inhibitor or an NS3/4A protease inhibitor, but not both, member has genotype 1.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MEGACE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Cachexia associated with cystic fibrosis.  
Cachexia associated with cancer.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Breast Cancer: Megestrol acetate is being used for palliative treatment.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Anorexia and cachexia associated with AIDS: Failure or clinically significant adverse effects to oxandrolone and dronabinol.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MEGACE ES

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Cachexia associated with cystic fibrosis.  
Cachexia associated with cancer.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Breast Cancer: Megestrol acetate is being used for palliative treatment.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Anorexia and cachexia associated with AIDS: Failure or clinically significant adverse effects to oxandrolone and dronabinol.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MEKINIST

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Uveal melanoma.

#### **Exclusion Criteria:**

MELANOMA: Monotherapy for patients who have disease progression on prior BRAF inhibitor therapy.

#### **Required Medical Information:**

MELANOMA: Positive for the BRAF V600E or V600K mutation detected by an FDA-approved test. NON-SMALL CELL LUNG CANCER, ANAPLASTIC THYROID CANCER: Positive for BRAF V600E mutation.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

NON-SMALL CELL LUNG CANCER, ANAPLASTIC THYROID CANCER: Used in combination with Tafinlar.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MEKTOVI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Positive for BRAF V600E or V600K mutation as detected by an FDA-approved test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Prescribed in combination with Braftovi.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

METHAMPHETAMINE

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Treatment of obesity.

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

METHOCARBAMOL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient is continuing on this medication without adverse effects.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

METHOTREXATE INJ (Otrexup, Rasuvo)

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to generic methotrexate injection.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MIRVASO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Diagnosis of persistent facial erythema of rosacea with papules and pustules of rosacea: Failure or clinically significant adverse effects to topical metronidazole, Finacea or oral doxycycline.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

MOZOBIL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation of patient's current weight and absolute neutrophil count (ANC dated within 30 days prior to the request).

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist or hematologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Documented failure to reach and/or maintain a target absolute neutrophil count (ANC) with an adequate trial of Neupogen alone. Must be administered in combination with a granulocyte-colony stimulating factor (G-CSF) (i.e., filgrastim, filgrastim-sndz, or tbo-filgrastim).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NAMENDA (includes Namenda XR)

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Vascular dementia.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 59 years and younger. Prior authorization is not required for patients 60 years and older.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NATPARA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NERLYNX

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months total duration of therapy.

#### **Other Criteria:**

Documentation of previous treatment with Herceptin (trastuzumab) as adjuvant therapy.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NESINA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Tradjenta or Jentadueto AND Failure or clinically significant adverse effects to one of the following: Januvia, Janumet, Janumet XR.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NEULASTA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Mobilization of peripheral-blood progenitor cells prior to autologous transplantation.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NEUPOGEN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Myelodysplastic syndrome. Neutropenia in patients with HIV/AIDS.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NINLARO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one prior therapy [e.g., Velcade (bortezomib), cyclophosphamide (Cytoxan), doxorubicin, Revlimid (lenalidomide), Thalomid (thalidomide), Alkeran (melphalan)]. Ninlaro must be used in combination with dexamethasone.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NORPACE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient is continuing on this medication without adverse effects.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NORTHERA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NUCALA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient has a blood eosinophil count of greater than or equal to 150 cells/mcL within the past 3 months.  
CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

Patient is 12 years of age or older.

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an allergist, pulmonologist, or immunologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Must be used in combination with ONE inhaled corticosteroid (e.g., beclomethasone, budesonide, flunisolide, fluticasone, mometasone, ciclesonide) AND must be used in combination with ONE long-acting beta-agonist (e.g., salmeterol, formoterol, vilanterol), unless contraindicated.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NUEDEXTA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

NUPLAZID

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OCALIVA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Must be used in combination with ursodeoxycholic acid unless patient is intolerant to ursodeoxycholic acid.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OCREVUS

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Member will not use other disease modifying therapies for MS concurrently. CONTINUATION OF THERAPY: Member is maintained on therapy with positive response (e.g., improved or maintained disease control evidenced by decreased or stabilized Expanded Disability Status Scale (EDSS) score or reduction in relapses or MRI lesions).

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a Neurologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Relapsing Forms Of Multiple Sclerosis: Failure or clinically significant adverse effects to one of the following: Aubagio, Tecfidera, Gilenya, Avonex, Betaseron, Plegridy, Copaxone, Glatopa, Extavia or Rebif.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ODOMZO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Basal cell carcinoma has recurred following surgery or radiation therapy, or member is not a candidate for surgery or radiation therapy.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OFEV

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ONGLYZA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Tradjenta or Jentadueto AND Failure or clinically significant adverse effects to one of the following: Januvia, Janumet, Janumet XR.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OPSUMIT

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ORENCIA CLICKJECT

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

RHEUMATOID ARTHRITIS: Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin. PSORIATIC ARTHRITIS: Failure or clinically significant adverse effects to methotrexate unless contraindicated.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ORENCIA IV

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

RHEUMATOID ARTHRITIS: Failure or clinically significant adverse effects to Remicade AND one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ORENCIA SC

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

RHEUMATOID ARTHRITIS: Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin. PSORIATIC ARTHRITIS: Failure or clinically significant adverse effects to methotrexate unless contraindicated.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ORKAMBI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Presence of homozygous F508del mutation in an FDA-cleared cystic fibrosis mutation test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OSENI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to Tradjenta or Jentadueto AND Failure or clinically significant adverse effects to one of the following: Januvia, Janumet, Janumet XR.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OSMOLEX ER

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Medical justification supports inability to use immediate-release amantadine.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of immediate-release amantadine unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

OTEZLA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

PSORIATIC ARTHRITIS, PLAQUE PSORIASIS: Prescribed by or in consultation with a rheumatologist or dermatologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

PLAQUE PSORIASIS: Failure or clinically significant adverse effects to ONE of the following: methotrexate, cyclosporine or acitretin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PARAFON FORTE DSC

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient is continuing on this medication without adverse effects.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PERSERIS

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of TWO of the following atypical antipsychotics, unless contraindicated or clinically significant adverse effects are experienced: aripiprazole, ziprasidone, quetiapine, olanzapine, risperidone, asenapine, iloperidone, paliperidone.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PHENOBARBITAL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Partial seizures: Failure or clinically significant adverse effects to one of the following: carbamazepine, phenytoin, topiramate, tiagabine, levetiracetam, gabapentin, lamotrigine, oxcarbazepine, primidone or divalproex. Generalized seizures: Failure or clinically significant adverse effects to one of the following: carbamazepine, phenytoin, topiramate, levetiracetam, primidone or lamotrigine. Sedation: patient is continuing on this medication without adverse effects.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PLEGRIDY

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PRALUENT

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Heterozygous Familial Hypercholesterolemia : Documentation (e.g., medical records, chart notes, laboratory values) of LDL level suggestive of a diagnosis of heterozygous familial hypercholesterolemia (e.g., Adults: LDL greater than 190 mg/dL). Hypercholesterolemia: Documentation of an LDL of 100 mg/dL or greater AND documented history of clinical atherosclerotic cardiovascular disease defined as one of the following: Acute coronary syndromes, Myocardial Infarction, Stable or unstable angina, Coronary or other arterial revascularization (e.g., percutaneous coronary intervention or coronary artery bypass graft surgery), Stroke, Peripheral artery disease presumed to be of atherosclerotic origin, Transient ischemic attack (TIA), Clinically significant coronary heart disease (CHD) diagnosed by invasive or noninvasive testing (such as coronary angiography, stress test using treadmill, stress echocardiography, or nuclear imaging), Carotid artery occlusion greater than 50% without symptoms, Renal artery stenosis or renal artery stent procedure. Reauthorization requests require documentation of LDL reduction while on Praluent therapy AND, if tolerated, confirmation of continued statin therapy at the maximally tolerated dose.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a cardiologist, endocrinologist, or lipid specialist.

#### **Coverage Duration:**

6 months.

#### **Other Criteria:**

Clinically significant adverse effect(s), contraindication(s), intolerance or failure to two of the following at maximally tolerated doses: atorvastatin, rosuvastatin, simvastatin, Vytorin, pitavastatin, pravastatin, fluvastatin, or lovastatin.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PREVYMIS

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Member is receiving pimozide or ergot alkaloids. Member is receiving cyclosporine co-administered with pitavastatin or simvastatin.

#### **Required Medical Information:**

Intravenous (IV) Prevymis: Medical justification why the member cannot use oral therapy.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncology, hematology, infectious disease, or transplant specialist.

#### **Coverage Duration:**

Through day 100 post-transplantation.

#### **Other Criteria:**

Failure of generic valacyclovir or generic ganciclovir, unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PROLIA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Hypocalcemia (unless corrected prior to initiating therapy).

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

For men with non-metastatic prostate cancer: Receiving or has received androgen deprivation therapy [i.e. leuprolide (Lupron), bicalutamide (Casodex) or Nilandron]. For women with breast cancer: Receiving or has received adjuvant aromatase inhibitor therapy [i.e. anastrozole (Arimidex), exemestane (Aromasin) or letrozole (Femara)].

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PROMACTA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Thrombocytopenia in Chronic Hepatitis C: Documentation of current or planned interferon-based treatment of chronic hepatitis C.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Chronic Immune (Idiopathic) Thrombocytopenia: Failure of a corticosteroid (e.g., oral prednisone, intravenous methylprednisolone or oral dexamethasone), unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PROTOPIC

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Tacrolimus 0.1%: 16 years and older.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of two medium to high potency topical corticosteroids (e.g., amcinonide, fluticasone propionate, triamcinolone acetonide, betamethasone valerate, fluocinolone acetonide, hydrocortisone butyrate, mometasone furoate, desoximetasone, fluocinonide or betamethasone dipropionate), unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PROVIGIL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Multiple sclerosis-related fatigue.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

PURIXAN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Lymphoblastic lymphoma.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Member has a documented swallowing disorder or an inability to swallow tablets or capsules. CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with oncologist or hematologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to mercaptopurine tablets.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

QUALAQUIN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Babesiosis. Plasmodium vivax malaria.

#### **Exclusion Criteria:**

For the treatment or prevention of nocturnal leg cramps.

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Malaria: 7 days. Babesiosis: 7-10 days

#### **Other Criteria:**

Plasmodium vivax malaria: Infection is chloroquine-resistant.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

RADICAVA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Forced vital capacity greater than or equal to 80%, disease duration of less than or equal to 2 years, functionally retains most activities of daily living (defined as a baseline revised ALS Functional Rating Scale (ALSFRS-R) score with greater than or equal to 2 points in each of the 12 items, meets diagnostic criteria of definite or probable amyotrophic lateral sclerosis (ALS) based on El Escorial revised criteria. CONTINUATION OF THERAPY: Member continues to retain most activities of daily living, forced vital capacity greater than or equal to 80%, and ALSFRS-R score with greater than or equal to 2 points in each of the 12 items.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist.

#### **Coverage Duration:**

6 months.

#### **Other Criteria:**

Prescribed in combination with riluzole unless contraindicated or clinically significant adverse effects are experienced.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

RANEXA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients on strong CYP3A inhibitors (e.g., ketoconazole, HIV protease inhibitors, clarithromycin) or CYP3A inducers (e.g., rifampin, phenobarbital).

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

RAYALDEE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient has stage 3 or 4 chronic kidney disease (CKD) and serum total 25-hydroxyvitamin D level less than 30 ng/mL.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

REMICADE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Wegener's Granulomatosis.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: RHEUMATOID ARTHRITIS and PLAQUE PSORIASIS: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Psoriatic Arthritis/Plaque Psoriasis: Prescribed by or in consultation with a rheumatologist or dermatologist. Crohn's Disease/Ulcerative Colitis: Prescribed by or in consultation with a gastroenterologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Rheumatoid Arthritis: Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin. Plaque Psoriasis: Failure or clinically significant adverse effects to one of the following: methotrexate, cyclosporine or acitretin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

REPATHA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Heterozygous or Homozygous Familial Hypercholesterolemia : Documentation (e.g., medical records, chart notes, laboratory values) of LDL level suggestive of a diagnosis of familial hypercholesterolemia (e.g., Adults: LDL greater than 190 mg/dL). Hypercholesterolemia: Documentation of an LDL of 100 mg/dL or greater AND documented history of clinical atherosclerotic cardiovascular disease defined as one of the following: Acute coronary syndromes, Myocardial Infarction, Stable or unstable angina, Coronary or other arterial revascularization (e.g., percutaneous coronary intervention or coronary artery bypass graft surgery), Stroke, Peripheral artery disease presumed to be of atherosclerotic origin, Transient ischemic attack (TIA), Clinically significant coronary heart disease (CHD) diagnosed by invasive or noninvasive testing (such as coronary angiography, stress test using treadmill, stress echocardiography, or nuclear imaging), Carotid artery occlusion greater than 50% without symptoms, Renal artery stenosis or renal artery stent procedure. Reauthorization requests require documentation of LDL reduction while on Repatha therapy AND, if tolerated, confirmation of continued statin therapy at the maximally tolerated dose.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a cardiologist, endocrinologist, or lipid specialist.

#### **Coverage Duration:**

6 months.

#### **Other Criteria:**

Clinically significant adverse effect(s), contraindication(s), intolerance or failure to two of the following at maximally tolerated doses: atorvastatin, rosuvastatin, simvastatin, Vytorin, pitavastatin, pravastatin, fluvastatin, or lovastatin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

REVATIO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients taking nitrates (e.g., Nitrodur, Nitrobid, Nitrostat, Isordil, Ismo). Patients taking a guanylate cyclase stimulator, such as riociguat (Adempas).

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

REVLIMID

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients who are pregnant.

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Mantle Cell Lymphoma: Failure of maximally tolerated doses of two prior chemo therapies (e.g., CHOP [cyclophosphamide, doxorubicin, vincristine, and prednisone], hyperCVAD [cyclophosphamide, vincristine, doxorubicin, and dexamethasone]) including Velcade unless contraindicated or clinically significant adverse effects are experienced. Multiple Myeloma: Must be used in combination with dexamethasone unless being used as maintenance therapy following autologous hematopoietic stem cell transplantation or as maintenance therapy for active (symptomatic) myeloma responding to primary myeloma therapy.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

REXULTI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Schizophrenia: Failure or clinically significant adverse effects to two of the following generic atypical antipsychotics: risperidone, olanzapine, quetiapine, ziprasidone, aripiprazole.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

RUBRACA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Mutations in the BRCA genes as detected by an FDA-approved test or member has a complete or partial response to two or more platinum-based chemotherapy regimens.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

RYDAPT

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Acute Myeloid Leukemia: Positive for the FLT3 mutation as detected by an FDA-approved test (e.g., LeukoStrat CDx FLT3 Mutation Assay). CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Acute Myeloid Leukemia: Prescribed by or in consultation with an oncologist. Advanced Systemic Mastocytosis: Prescribed by or in consultation with an oncologist, allergist, or immunologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Acute Myeloid Leukemia: Prescribed in combination with daunorubicin for induction therapy AND in combination with cytarabine for induction and consolidation therapy.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SEROSTIM

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

REAUTHORIZATION: Continued treatment will be approved with documentation of response to therapy.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

6 months.

#### **Other Criteria:**

Patient is being treated with concomitant antiretroviral therapy

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SILIQ

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a rheumatologist or dermatologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following as recommended by the American Academy of Dermatology: methotrexate, cyclosporine or acitretin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SIMPONI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

PSORIATIC ARTHRITIS: Prescribed by or in consultation with a rheumatologist or dermatologist. ULCERATIVE COLITIS: Prescribed by or in consultation with a gastroenterologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

RHEUMATOID ARTHRITIS: Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SIMPONI ARIA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SKELAXIN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient is continuing on this medication without adverse effects.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SOMA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient is continuing on this medication without adverse effects.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SOMAVERT

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Inadequate response to surgery or radiation therapy, unless surgery or radiation therapy is not appropriate for the patient.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SONATA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to two of the following: Rozerem, Silenor 6 mg/day or less, trazodone or temazepam.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SOVALDI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. For the treatment of hepatitis C virus genotypes 5 and 6.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Diagnosis of chronic hepatitis C (CHC) and genotype 1, 2, 3, 4, 5 or 6 confirmed by detectable serum hepatitis C virus RNA by quantitative assay OR For treatment of CHC in patients with hepatocellular carcinoma (HCC) meeting Milan criteria (awaiting liver transplantation). Milan criteria is defined as the presence of a tumor 5 cm or less in diameter in patients with single hepatocellular carcinomas and no more than three tumor nodules, each 3 cm or less in diameter in patients with multiple tumors and no extrahepatic manifestations of the cancer or evidence of vascular invasion of tumor. Refer to the document, "Recommendations for Testing, Managing, and Treating Hepatitis C," by AASLD-IDSA available at <http://www.hcvguidelines.org> for drug regimen and duration of treatment based on genotype, treatment status, previous drug regimens used, past medical history and comorbidities.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a gastroenterologist, hepatologist or infectious disease physician.

#### **Coverage Duration:**

GT 1 to 6: 12 to 24 weeks or HCC with CHC: up to 48 weeks or until liver transplantation

#### **Other Criteria:**

For Sovaldi in combination with Daklinza for genotype 1: Failure or clinically significant adverse effects to Harvoni (sofosbuvir/ledipasvir). For Sovaldi in combination with Daklinza for genotype 2: Failure or clinically significant adverse effects to sofosbuvir/ribavirin. For patients with hepatocellular carcinoma (HCC) meeting Milan criteria (awaiting liver transplantation): must be used in combination with ribavirin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SPRITAM

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Medical justification must be provided why patient cannot take generic levetiracetam tablets or liquid.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SPRYCEL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Accelerated, or myeloid or lymphoid blast phase Philadelphia chromosome-positive chronic myeloid leukemia (CML) or Philadelphia chromosome-positive acute lymphoblastic leukemia (ALL): Failure of imatinib, unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

STELARA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

PSORIATIC ARTHRITIS, PLAQUE PSORIASIS: Prescribed by or in consultation with a rheumatologist or dermatologist. CROHN'S DISEASE: Prescribed by or in consultation with a gastroenterologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

PLAQUE PSORIASIS: Failure or clinically significant adverse effects to ONE of the following: methotrexate, cyclosporine, or acitretin. PSORIATIC ARTHRITIS: Failure or clinically significant adverse effects to methotrexate unless contraindicated.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

STELARA IV

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a gastroenterologist.

#### **Coverage Duration:**

4 weeks.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

STIVARGA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

METASTATIC COLORECTAL CANCER: Documentation that the patient does or does not have the RAS wild type gene. Documentation that the patient does or does not have the BRAF V600E mutation. CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

METASTATIC COLORECTAL CANCER: If tumor does not have the RAS wild type gene, failure or clinically significant adverse effects to one of the following: 5-fluorouracil, capecitabine, oxaliplatin, irinotecan, Cyramza, Avastin, Zaltrap OR If tumor expresses the RAS wild type gene without the BRAF V600E mutation, failure or clinically significant adverse effects to Erbitux or Vectibix. GASTROINTESTINAL STROMAL TUMOR: Failure or clinically significant adverse effects to one of the following: Gleevec or Sutent.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

STRENSIQ

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SUBSYS

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Age 18 or greater

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Through the end of the Plan contract year.

#### **Other Criteria:**

Patient is already taking and is tolerant to around-the-clock opioid therapy. Patients are considered opioid tolerant when taking another opioid daily for a week or longer (for example, at least 60 mg of oral morphine per day or an equianalgesic dose of another opioid).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SUBUTEX

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Patient is pregnant OR Written documentation of intolerance to naloxone.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

Non pregnant: 3 months initial. Pregnant patients: 9 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SURMONTIL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Irritable bowel syndrome.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Depression: Failure or clinically significant adverse effects to one of the following: bupropion, bupropion SR, bupropion XL, citalopram, desvenlafaxine succinate, duloxetine, escitalopram, fluoxetine, mirtazapine, sertraline, venlafaxine, or venlafaxine XR.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SYMDEKO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Presence of homozygous F508del mutation or at least one mutation in the CFTR gene that is responsive to tezacaftor/ivacaftor.

#### **Age Restrictions:**

Age greater than or equal to 12 years.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

SYMLINPEN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation that the current HbA1c level is greater than 7%.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Diabetes Type 2: Failure of a metformin-containing regimen, unless contraindicated or clinically significant adverse effects are experienced. Diabetes Type 1: Failure of an insulin-containing regimen, unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TAGRISSO

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Disease is positive for any of the following, as detected by an FDA-approved test: exon 19 deletions, exon 21 L858R mutations, or T790M mutation with progression on or after an EGFR TKI therapy (e.g., Tarceva, Iressa, or Gilotrif).

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TALTZ

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a rheumatologist or dermatologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following: methotrexate, cyclosporine or acitretin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TARCEVA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Non-small cell lung cancer: Documentation that the patient has EGFR exon 19 deletions or exon 21 (L858R) substitution mutations as detected by an FDA-approved test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Pancreatic cancer: Tarceva is being prescribed in combination with gemcitabine.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TASIGNA

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients with hypokalemia, hypomagnesemia, or long QT syndrome.

#### **Required Medical Information:**

Chronic Myelogenous Leukemia (CML), Acute Lymphoblastic Leukemia (ALL): Documentation that the patient has Philadelphia chromosome positive disease.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Soft Tissue Sarcoma Gastrointestinal Stromal Tumor: Failure to imatinib, sunitinib, or regorafenib, unless contraindicated or clinically significant adverse effects are experienced

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TAVALISSE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of a corticosteroid (e.g., prednisone, methylprednisolone, or dexamethasone), unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TECENTRIQ

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of platinum-containing chemotherapy (e.g., cisplatin or carboplatin), OR the patient is not eligible for cisplatin-containing chemotherapy. Non-small cell lung cancer: If a known epidermal growth factor receptor (EGFR) or anaplastic lymphoma kinase (ALK) genomic tumor aberration exists, then for ALK+ disease: prior trial of Xalkori or Alecensa OR for EGFR+ disease: prior trial of Tarceva, Gilotrif or Iressa.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TECFIDERA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a neurologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TENEX

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to two of the following: amlodipine/benazepril, benazepril, benazepril/hydrochlorothiazide, captopril, captopril/hydrochlorothiazide, fosinopril, fosinopril/hydrochlorothiazide, lisinopril, lisinopril/hydrochlorothiazide, quinapril, quinapril/hydrochlorothiazide, losartan, losartan/hydrochlorothiazide, valsartan, valsartan/hydrochlorothiazide, irbesartan, irbesartan/hydrochlorothiazide, candesartan, candesartan/hydrochlorothiazide, carvedilol, labetalol, acebutolol, atenolol, bisoprolol, bisoprolol/hydrochlorothiazide, timolol, nadolol, propranolol, metoprolol, metoprolol/hydrochlorothiazide, pindolol, nifedipine SR, amlodipine, nicardipine.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TETRABENAZINE

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TIBSOVO

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Presence of an isocitrate dehydrogenase-1 (IDH1) mutation as detected by an FDA-approved test (e.g., Abbott RealTime IDH1 Assay).

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist or hematologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

For age less than 60 years, disease has relapsed or is refractory following treatment with a first-line chemotherapy regimen (e.g., cytarabine, idarubicin, daunorubicin, cladribine, midostaurin).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TREMFYA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a rheumatologist or dermatologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to ONE of the following as recommended by the American Academy of Dermatology: methotrexate, cyclosporine, or acitretin.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TRIHEXYPHENIDYL

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Parkinsons disease/Parkinsonism: Failure or clinically significant adverse effects to two of the following: amantadine, levodopa/carbidopa, entacapone, pramipexole, ropinirole, selegiline. All other FDA-approved indications: Patient is continuing on this medication without adverse effects.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TYMLOS

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response. Tymlos has not been used for more than two years.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of a bisphosphonate (e.g., alendronate) unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

TYSABRI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients who have or have had progressive multifocal leukoencephalopathy.

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

MULTIPLE SCLEROSIS: Prescribed by or in consultation with a neurologist. CROHN'S DISEASE: Prescribed by or in consultation with a GI specialist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

RELAPSING FORMS OF MULTIPLE SCLEROSIS: Failure or clinically significant adverse effects to one of the following: Aubagio, Tecfidera, Gilenya, Avonex, Betaseron, Plegridy, Copaxone, Glatopa, Extavia or Rebif.

CROHN'S DISEASE: Failure or clinically significant adverse effects to Humira or Remicade.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

UPTRAVI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VALCHLOR

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following skin-directed therapies: topical corticosteroids (e.g., clobetasol, triamcinolone), Targetin gel, Tazorac, or imiquimod. FOR CONTINUATION OF THERAPY: Maintained on therapy with positive response.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VANCOGIN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a gastroenterologist, infectious disease specialist or hospitalist.

#### **Coverage Duration:**

C. Diff diarrhea: 14 days. Staph enterocolitis: 10 days. Recurrent C. Diff: 10 weeks

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VENCLEXTA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of clinically significant adverse effects to one previous therapy (e.g., Imbruvica, Campath, high-dose methylprednisolone with Rituxan).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VERSACLOZ

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Psychotic disorder associated with Parkinson's disease.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure of or clinically significant adverse effects to clozapine (Clozaril) or FazaClo.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VERZENIO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Hormone receptor (HR)-positive, human epidermal growth factor receptor 2 (HER2)-negative advanced or metastatic breast cancer.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VIBERZI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to loperamide and either diphenoxylate-atropine or dicyclomine, unless patient is 65 years or older.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VINBLASTINE

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation that vinblastine is being used as palliative therapy.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VINCRIStINE

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

Patients with the demyelinating form of Charcot-Marie-Tooth syndrome.

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Hodgkin's disease, non-Hodgkin's malignant lymphomas, rhabdomyosarcoma, neuroblastoma, Wilms' tumor: use in combination with other oncolytic agents.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VOSEVI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

For members with cirrhosis, documentation of Child-Pugh A status. Refer to the document, "Recommendations for Testing, Managing, and Treating Hepatitis C," by AASLD-IDSAs available at <http://www.hcvguidelines.org> for drug regimen and duration of treatment based on genotype, treatment status, previous drug regimens used, past medical history and comorbidities.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a gastroenterologist, hepatologist or infectious disease physician.

#### **Coverage Duration:**

12 weeks.

#### **Other Criteria:**

If HCV genotype 1, 2, 3, 4, 5 or 6, member has previously been treated with an HCV regimen containing one of the following NS5A inhibitors: daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir. Alternatively, if HCV genotype is 1a or 3, member has previously been treated with an HCV regimen containing sofosbuvir.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VOTRIENT

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

SOFT TISSUE SARCOMA: Member has received prior chemotherapy (e.g., regimens containing doxorubicin or epirubicin).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

VRAYLAR

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to TWO of the following atypical antipsychotics: aripiprazole, ziprasidone, quetiapine, olanzapine, risperidone.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XALKORI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation that the patient is ALK-positive as detected by an FDA-approved test or that the patient is ROS-1 positive as confirmed by a laboratory-developed break-apart FISH or RT-PCR clinical trial assay.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XATMEP

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

Less than 18 years of age.

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist (for acute lymphoblastic leukemia) or rheumatologist (for polyarticular juvenile idiopathic arthritis).

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Medical justification as to why member cannot use methotrexate tablets.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XELJANZ (includes Xeljanz XR)

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to one of the following: methotrexate, sulfasalazine, hydroxychloroquine, d-penicillamine, azathioprine or auranofin.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XEOMIN

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

---

### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XERMELO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Prescribed in combination with a somatostatin analog (e.g., octreotide, lanreotide) unless contraindicated or clinically significant adverse effects are experienced. CONTINUATION OF THERAPY: Maintained on therapy with positive response (e.g., reduction in bowel movement frequency, reduction in urinary 5-HIAA levels).

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure to a trial of a somatostatin analog (e.g., octreotide, lanreotide) unless contraindicated or clinically significant adverse effects are experienced.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XOLAIR

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Moderate to severe persistent asthma: Patient has a positive skin test or in vitro reactivity to a perennial aeroallergen AND Patient has a confirmed total serum IgE level greater than 30 IU/ml. CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

Asthma: Patient is 6 years of age or older. Chronic Idiopathic Urticaria: Patient is 12 years of age or older.

#### **Prescriber Restrictions:**

Asthma: Prescribed by or in consultation with a pulmonologist, immunologist, or allergist. Urticaria: Prescribed by or in consultation with an allergist, dermatologist, or immunologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Moderate to severe persistent asthma: Failure or clinically significant adverse effects to one inhaled corticosteroid (e.g., beclomethasone, budesonide, flunisolide, fluticasone, mometasone, ciclesonide). Chronic Idiopathic Urticaria: Failure or clinically significant adverse effects to one H1 Antihistamine (e.g., levocetirizine or desloratadine).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

XTANDI

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

For patients without visceral metastases: failure or clinically significant adverse effects to Zytiga.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

YERVOY

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Small cell lung cancer.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Small cell lung cancer: Disease relapse within 6 months following complete or partial response or stable disease.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Small cell lung cancer: Disease relapse with initial treatment (e.g., cisplatin, carboplatin containing regimen).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

YONSA

#### **Covered Uses:**

All medically accepted indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist or urologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Prescribed in combination with methylprednisolone. Member has previously had bilateral orchiectomy, failed androgen deprivation therapy (ADT) or will use ADT concurrently with Yonsa.



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZALTRAP

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Colorectal cancer is resistant or has progressed following an oxaliplatin-containing regimen AND Zaltrap will be used in combination with 5-fluorouracil, leucovorin, and irinotecan (FOLFIRI).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZARXIO

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Myelodysplastic syndrome. Neutropenia in patients with HIV/AIDS. Hematopoietic syndrome of acute radiation syndrome.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

---

### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZEJULA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Completed two or more platinum-based chemotherapy regimens and are in a complete or partial response.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZELBORAF

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Non-small cell lung cancer. Hairy Cell Leukemia.

#### **Exclusion Criteria:**

MELANOMA: Patients with wild-type BRAF melanoma.

#### **Required Medical Information:**

MELANOMA, NON-SMALL CELL LUNG CANCER: Positive for the BRAF V600E mutation detected by an FDA-approved test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

HAIRY CELL LEUKEMIA: Condition is non-responsive to purine analog therapy (e.g., pentostatin, cladribine).

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZEPATIER

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Refer to the document, "Recommendations for Testing, Managing, and Treating Hepatitis C," by AASLD-IDSA available at <http://www.hcvguidelines.org> for drug regimen and duration of treatment based on genotype, treatment status, previous drug regimens used, past medical history and comorbidities.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a gastroenterologist, hepatologist or infectious disease physician.

#### **Coverage Duration:**

12 to 16 wks based on genotype,presence of NS5A resistance-associated polymorphisms,prior treatment.

#### **Other Criteria:**

## Prior Authorization Protocol

---

### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZINPLAVA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation of positive Clostridium difficile test.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

4 weeks.

#### **Other Criteria:**

Will receive or is currently receiving antibacterial drug treatment for Clostridium difficile infection (e.g., metronidazole, vancomycin, fidaxomicin) concomitantly with Zinplava. Has received appropriate treatment for past CDI recurrences, including a pulsed vancomycin regimen.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZOLPIDEM

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

Prior authorization is required for patients 65 years and older. Prior authorization is not required for patients 64 years and younger.

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Failure or clinically significant adverse effects to two of the following: Rozerem, Silenor 6 mg/day or less, trazodone or temazepam.

## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZYDELIG

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. MALT lymphoma (gastric and nongastric). Splenic Marginal Zone Lymphoma. Primary Cutaneous Marginal Zone B-Cell Lymphoma. Nodal Marginal Zone Lymphoma.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with a hematologist or oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

For relapsed follicular B-cell non-Hodgkin lymphoma (FL) or relapsed small lymphocytic lymphoma (SLL): failure or clinically significant adverse effects to two prior systemic therapies (e.g., For FL: Leukeran, Rituxan, Treanda, R-CHOP, R-CVP, FCMR or for SL: Leukeran, Gazyva, FCR, FR, BR or PCR).



## Prior Authorization Protocol

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### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZYKADIA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D. Soft tissue sarcoma - inflammatory myofibroblastic tumor.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

Documentation that the patient does or does not have anaplastic lymphoma kinase (ALK)-positive disease.  
CONTINUATION OF THERAPY: Maintained on therapy with positive response.

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

Prescribed by or in consultation with an oncologist.

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

## Prior Authorization Protocol

---

### Medicare Part D – 2018

#### **Prior Authorization Group Description**

ZYTIGA

#### **Covered Uses:**

All FDA-approved indications not otherwise excluded from Part D.

#### **Exclusion Criteria:**

#### **Required Medical Information:**

#### **Age Restrictions:**

#### **Prescriber Restrictions:**

#### **Coverage Duration:**

12 months.

#### **Other Criteria:**

Must be used in combination with prednisone.

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