

## Drugs: Contract Drugs List Part 1 – Prescription Drugs (N through R)

Page updated: October 2020

This section lists the codes and units for contract drugs. For additional help, refer to the *Drugs: Contract Drugs List Introduction* section of this manual.

### Nabumetone

«The following text is removed effective November 1, 2020:» Nabumetone is restricted to use for arthritis. **Note:** Subject to Step Therapy edits. See Drugs: Contract Drugs List Part 8 – Step Therapy for more information. «End of removed text.»

Dosage Form	Size and/or Strength	Billing Unit
Tablets	500 mg	each
Tablets	750 mg	each

### Nafcillin

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	500 mg/vial	each
Powder for injection	1 gm/vial	each
Powder for injection	2 gm/vial	each
Powder for injection	10 gm/vial	each
Powder for injection	1 gm, piggyback	each
Powder for injection	2 gm, piggyback	each

**Naftifine HCl \***

\* «Naftifine HCl is» restricted to claims with dates of service from October 1, 2005, through November 30, 2011, for all dosage forms.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Topical Cream	1 %, 15 gm	gram
Topical Cream	1 %, 30 gm	gram
Topical Cream	1 %, 60 gm	gram
Topical Cream	1 %, 90 gm	gram
Topical Gel	1 %, 20 gm	gram
Topical Gel	1 %, 40 gm	gram
Topical Gel	1 %, 60 gm	gram
Topical Gel	1 %, 90 gm	gram

**Nalidixic Acid**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	250 mg	each
Tablets	500 mg	each
Tablets	1 gm	each

**Note:** «Effective, March 1, 2017, these products are no longer manufactured or available.»

## Naloxegol Oxalate \*

\* Naloxegol Oxalate is restricted to use in the treatment of opioid-induced constipation in patients with chronic pain. Also restricted to NDC labeler code 00310 (AstraZeneca LP), and «(effective October 1, 2020,)» restricted to NDC labeler code 57841 only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	12.5 mg	each
Tablets	25 mg	each

## Naloxone HCl

Dosage Form	Size and/or Strength	Billing Unit
Injection	0.4 mg/ml	milliliter
Injection	1.0 mg/ml	milliliter
Intranasal Spray	4.0 mg/0.1 ml	each

## Naltrexone HCl

The following text is removed effective June 1, 2019: Naltrexone HCl is restricted to use in the treatment of alcohol dependence and for the prevention of relapse in opioid dependent patients, following opioid detoxification.

Naltrexone HCl is restricted to prescription only by prescribers trained in substance use disorder treatment.

Naltrexone HCl is restricted to a maximum dispensing quantity of 100 tablets and a maximum of three (3) dispensings in any 75 day period. End of removed text.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	50 mg	each

## Naphazoline HCl

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.1 %	milliliter

## Naphazoline HCl and Antazoline Phosphate

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.05 % – 0.5 %	milliliter

## Naproxen

«The following text is removed effective November 1, 2020:» **Note:** Subject to Step Therapy edits. See Drugs: Contract Drugs List Part 8 – Step Therapy for more information.  
«End of removed text.»

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules +	250 mg	each
Tablets or capsules +	375 mg	each
Tablets or capsules +	500 mg	each
Liquid	125 mg/5 ml	milliliter

## Natamycin

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	5 %, 15 ml	milliliter

**Nateglinide**

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	60 mg	each
Tablets +	120 mg	each

**Necitumumab ‡ \***

\* Necitumumab is restricted to use in the treatment of cancer only. Effective January 1, 2017, Necitumumab is also restricted to labeler code 00002 (Eli Lilly and Company) only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	800 mg/50 ml	milliliter

**Nelfinavir Mesylate ‡ \***

\* Nelfinavir Mesylate is restricted to use as combination therapy in the treatment of Human Immunodeficiency Virus (HIV) infection.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	250 mg	each
Tablets	625 mg	each
Oral powder	50 mg/gm	gram

**Nelarabine ‡ \***

\* «Effective October 1, 2020, Nelarabine is restricted to use in the treatment of cancer only. Also restricted to labeler code 00078 (Novartis Pharmaceuticals Corporation) only.»

Dosage Form	Size and/or Strength	Billing Unit
Injection	5 mg/ml	milliliter

**Neomycin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	0.5 gm	each
Liquid	125 mg/5 ml	milliliters

**Neomycin, Bacitracin and Polymyxin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Ophthalmic ointment	<<Blank>>	gram

**Neomycin and Polymyxin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Ampule – G.U. Irrigant	<<Blank>>	milliliters

**Neomycin, Polymyxin and Gramicidin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Ophthalmic solution	<<Blank>>	milliliters

## Neostigmine Bromide

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	15 mg	each

**Note:** This product is no longer manufactured or available.

## Nepafenac

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic suspension	0.1 % *	milliliter

\* «Effective March 1, 2020, Nepafenac is restricted to claims submitted with dates of service from August 1, 2005, through March 31, 2015, for the 0.1% only.»

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic suspension	0.3 % *	milliliter

\* «Effective March 1, 2020, Nepafenac is restricted to the 3 ml bottle size only. The 1.7 ml bottle size is restricted to claims submitted with dates of service from October 1, 2013, through February 29, 2020, for the 0.3% suspension only.»

## Netarsudil \*

\* «Netarsudil is» restricted to labeler code 70727 (Aerie Pharmaceuticals, Inc.)

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.02 %	milliliter

## Netarsudil/Latanoprost \*

\* «Netarsudil/Latanoprost is» restricted to labeler code 70727 (Aerie Pharmaceuticals, Inc.)

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.02 %/0.005%	milliliter

**Nevirapine ‡ \***

\* Nevirapine is restricted to use as combination therapy in the treatment of Human Immunodeficiency Virus (HIV) infection. «**Effective April 1, 2021: the liquid dose (50 mg/5 ml) is restricted to NDC labeler code 64370.**»

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	200 mg	each
Tablets, extended release	100 mg	each
Tablets, extended release	400 mg	each
Liquid	50 mg/5 ml	milliliter

**Niacin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets, extended release (includes film coated tablets)	500 mg	each
Tablets, extended release (includes film coated tablets)	750 mg	each
Tablets, extended release (includes film coated tablets)	1000 mg	each



## Niacin and Lovastatin

Dosage Form	Size and/or Strength	Billing Unit
Tablets (containing extended release niacin)	500 mg/20 mg	each
Tablets (containing extended release niacin)	750 mg/20 mg	each
Tablets (containing extended release niacin)	1000 mg/20 mg	each
Tablets (containing extended release niacin)	1000 mg/40 mg	each

**Note:** These products are no longer manufactured or available.

## Niacin and Simvastatin\*

\* «Niacin and Simvastatin is» restricted to NDC labeler code 00074 (Abbott Laboratories) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets (containing extended release niacin)	500 mg/20 mg	each
Tablets (containing extended release niacin)	500 mg/40 mg	each
Tablets (containing extended release niacin)	750 mg/20 mg	each
Tablets (containing extended release niacin)	1000 mg/20 mg	each
Tablets (containing extended release niacin)	1000 mg/40 mg	each

**Note:** These products are no longer manufactured or available.

## Nicardipine

Dosage Form	Size and/or Strength	Billing Unit
Capsules +	20 mg	each
Capsules +	30 mg	each
Tablets or capsules, long-acting +	30 mg	each
Tablets or capsules, long-acting +	45 mg	each
Tablets or capsules, long-acting +	60 mg	each

## Niclosamide

Dosage Form	Size and/or Strength	Billing Unit
Tablets	500 mg	each

**Note:** This product is no longer manufactured or available.

## Nifedipine

Dosage Form	Size and/or Strength	Billing Unit
Capsules +	10 mg	each
Capsules +	20 mg	each
Tablets or capsules, long-acting +	30 mg	each
Tablets or capsules, long-acting +	60 mg	each
Tablets or capsules, long-acting +	90 mg	each

**Nilotinib ‡ \***

\* «Nilotinib is» restricted to use in the treatment of cancer only for all strengths. Also restricted to NDC labeler code 00078 (Novartis Pharmaceuticals Corporation) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	150 mg	each
Capsules	200 mg	each

**Nilutamide ‡ \***

\* «Nilutamide is» restricted to use in the treatment of cancer only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	150 mg	each

**Niraparib ‡ \***

\* «Nilutamide is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 69656 (TESARO Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	100 mg	each

## Nisoldipine

Dosage Form	Size and/or Strength	Billing Unit
Tablets	8.5 mg	each
Tablets	10 mg *	each
Tablets	17 mg	each
Tablets	20 mg *	each
Tablets	25.5 mg	each
Tablets	30 mg *	each
Tablets	34 mg	each
Tablets	40 mg *	each

\* «Nisoldipine is» restricted to claims with dates of service from March 1, 1997, through March 31, 2010, for the 10 mg, 20 mg, 30 mg and 40 mg tablets only.

## Nitrofurantoin

Dosage Form	Size and/or Strength	Billing Unit
Capsules (macrocrystals only)	25 mg	each
Capsules (macrocrystals only)	50 mg	each
Capsules (macrocrystals only)	100 mg	each
Capsules (monohydrate/macrocrystals)	100 mg	each
Tablets	50 mg	each
Tablets	100 mg	each
Liquid	5 mg/ml	milliliter

**Nitroglycerin (Glyceryl Trinitrate)**

<b>Dosage Form</b>	<b>Percent, Size and/or Strength</b>	<b>Billing Unit</b>
Tablets (sublingual) (no long-acting forms) +	0.15 mg	each
Tablets (sublingual) (no long-acting forms) +	0.3 mg	each
Tablets (sublingual) (no long-acting forms) +	0.4 mg	each
Tablets (sublingual) (no long-acting forms) +	0.6 mg	each
Ointment	2 %, 20 gm	gram
Ointment	2 %, 30 gm	gram
Ointment	2 %, 60 gm	gram
Spray, lingual	2 %, 12 gm	gram

**Nivolumab ‡ \***

\* «Nivolumab is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00003 (E.R. Squibb & Sons, Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	40 mg/4 ml	milliliter
Injection	100 mg/10 ml	milliliter
Injection	240 mg/24 ml	milliliter

## Norelgestromin and Ethinyl Estradiol ‡ \*

\* «Norelgestromin and Ethinyl Estradiol are» restricted to a maximum dispensing quantity of up to 52 patches per client. The maximum quantity is intended for clients on a continuous cycle. A 12-month supply of the same product of contraceptive patches may be dispensed twice in one year. A *Treatment Authorization Request* (TAR) is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Transdermal patch	6 mg – 0.75 mg	each
Transdermal patch	4.86 mg – 0.53 mg	each

**Note:** Payment limited to a minimum dispensing quantity of three cycles except with the initial prescription or when authorization is obtained.

## Norethindrone ‡ \*

\* «Norethindrone is» restricted to a maximum quantity of up to 18 cycles (packs) per dispensing. The maximum supply is intended for clients on continuous cycle. A 12-month supply of the same product of oral contraceptive may be dispensed twice in one year. A *Treatment Authorization Request* (TAR) is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.35 mg Tablets from 28 tablet packet	each
Tablets	0.35 mg Tablets from 42 tablet packet	each

**Note:** Payment limited to a minimum dispensing quantity of three cycles. See *California Code of Regulations* (CCR), Title 22, Section 51513(b)(4) regarding exceptions.

## Norethindrone Acetate and Ethinyl Estradiol ‡ \*

\* Norethindrone Acetate and Ethinyl Estradiol are restricted to a maximum quantity of up to 18 cycles (packs) per dispensing. The maximum supply is intended for clients on continuous cycle. A 12-month supply of the same product of oral contraceptive may be dispensed twice in one year. A *Treatment Authorization Request* (TAR) is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	1 mg – 5 mcg	each §

The 1 mg to 5 mcg tablets are suspended until further notice.

Dosage Form	Size and/or Strength	Billing Unit
Tablets ‡	1 mg – 10 mcg/2 Fe tablets * Tablets from 28 tablet packet	each

\* The 1 mg to 10 mcg/2 Fe tablets are restricted to NDC Labeler Code 00430 (Warner Chilcott Laboratories) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	1 mg – 20 mcg Tablets from 21 tablet packet	each
Tablets	1 mg – 20 mcg/7 Fe tablets Tablets from 28 tablet packet	each
Tablets	1.5 mg – 30 mcg Tablets from 21 tablet packet	each
Tablets	1.5 mg – 30 mcg/7 Fe tablets Tablets from 28 tablet packet	each
Tablets from 5/7/9 combination packet (28 Tablets/packet) ‡	5 x 1 mg/20 mcg 7 x 1 mg/30 mcg 9 x 1 mg/35 mcg 7 inert	each

**Note:** Payment limited to a minimum dispensing quantity of three cycles. See *California Code of Regulations* (CCR), Title 22, Section 51513(b)(4) regarding exceptions.

## Norethindrone and Ethinyl Estradiol ‡ \*

\* «Norethindrone and Ethinyl Estradiol is» restricted to a maximum quantity of up to 18 cycles (packs) per dispensing. The maximum supply is intended for clients on continuous cycle. A 12-month supply of the same product of oral contraceptive may be dispensed twice in one year. A *Treatment Authorization Request* (TAR) is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.4 mg – 35 mcg Tablets from 21 tablet packet	each
Tablets	0.4 mg – 35 mcg Tablets from 28 tablet	each
Tablets	0.5 mg – 35 mcg Tablets from 21 tablet packet	each
Tablets	0.5 mg – 35 mcg Tablets from 28 tablet packet	each
Tablets	1 mg – 35 mcg Tablets from 21 tablet packet	each
Tablets	1 mg – 25 mcg Tablets from 28 tablet packet	each
Tablets	1 mg – 50 mcg Tablets from 21 tablet packet	each
Tablets	1 mg – 50 mcg Tablets from 28 tablet packet	each

**Note:** 1 mg – 50 mcg product is no longer manufactured or available.



**Norethindrone and Ethinyl Estradiol (continued) ‡ \***

Dosage Form	Size and/or Strength	Billing Unit
Tablets from 7/7/7 combination packet	7 x 0.5 mg/35 mcg	each
Tablets from 21 tablet packet	7 x 0.75 mg/35 mcg	
Tablets from 28 tablet packet	7 x 1.0 mg/35 mcg	
Tablets from 7/9/5 combination packet	7 x 0.5 mg/35 mcg	each
Tablets from 21 tablet packet	9 x 1.0 mg/35 mcg	
Tablets from 28 tablet packet	5 x 0.5 mg/35 mcg	
Tablets from 10/11 combination packet	10 x 0.5 mg/35 mcg	each
Tablets from 21 tablet packet	11 x 1 mg/35 mcg	
Tablets from 28 tablet packet		
Tablets from 7/14 combination packet (28 tablets packet)	7 x 0.5 mg/35 mcg 14 x 1 mg/35 mcg 7 inert	each

**Note:** 7/14 combination packet is no longer manufactured or available.

**Note:** Payment limited to a minimum dispensing quantity of three cycles. See *California Code of Regulations (CCR)*, Title 22, Section 51513(b)(4) regarding exceptions.

**Norethindrone and Mestranol ‡ \***

\* «Norethindrone and Mestranol is» restricted to a maximum quantity of up to 18 cycles (packs) per dispensing. The maximum supply is intended for clients on continuous cycle. A 12-month supply of the same product of oral contraceptive may be dispensed twice in one year. A *Treatment Authorization Request (TAR)* is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	1 mg – 50 mcg	each
	Tablets from 21 tablet packet	
Tablets	1 mg – 50 mcg	each
	Tablets from 28 tablet packet	

**Note:** Payment limited to a minimum dispensing quantity of three cycles. See *California Code of Regulations (CCR)*, Title 22, Section 51513(b)(4) regarding exceptions.

## Norfloxacin

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules	400 mg	each

**Note:** These products are no longer manufactured or available.

## Norgestimate and Ethinyl Estradiol ‡ \*

\* «Norgestimate and Ethinyl Estradiol is» restricted to a maximum quantity of up to 18 cycles (packs) per dispensing. The maximum supply is intended for clients on continuous cycle. A 12-month supply of the same product of oral contraceptive may be dispensed twice in one year. A *Treatment Authorization Request* (TAR) is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Tablets from 7/7/7 (tri-phasic) combination packet (21 tablets/packet)	7 x 0.180 mg/35 mcg 7 x 0.215 mg/35 mcg 7 x 0.250 mg/35 mcg	each
Tablets from 7/7/7 (tri-phasic) combination packet (28 tablets/packet)	7 x 0.180 mg/35 mcg 7 x 0.215 mg/35 mcg 7 x 0.250 mg/35 mcg 7 inert	each
Tablets from monophasic packet (28 tablets/packet)	21 x 0.25 mg/35 mcg 7 inert	each

**Note:** Payment limited to a minimum dispensing quantity of three cycles. See *California Code of Regulations* (CCR), Title 22, Section 51513(b)(4) regarding exceptions.

## Norgestrel and Ethinyl Estradiol ‡ \*

\* «Norgestrel and Ethinyl Estradiol is» restricted to a maximum quantity of up to 18 cycles (packs) per dispensing. The maximum supply is intended for clients on continuous cycle. A 12-month supply of the same product of oral contraceptive may be dispensed twice in one year. A *Treatment Authorization Request* (TAR) is required for the third supply of up to 12 months of the same product requested within a year.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.3 mg – 30 mcg Tablets from 21 tablet packet	each
Tablets	0.3 mg – 30 mcg Tablets from 28 tablet packet	each
Tablets	0.5 mg – 50 mcg Tablets from 21 tablet packet	each
Tablets	0.5 mg – 50 mcg Tablets from 28 tablet packet	each

**Note:** Payment limited to a minimum dispensing quantity of three cycles. See *California Code of Regulations* (CCR), Title 22, Section 51513(b)(4) regarding exceptions.

## Nortriptyline HCl \*

\* Use «of Nortriptyline HCl» in beneficiaries less than 18 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Capsules	10 mg	each
Capsules	25 mg	each
Capsules	50 mg	each
Capsules	75 mg	each
Liquid	10 mg/5 ml	milliliters

**Nystatin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets (oral) ‡	500,000 units	each
Suspension, oral ‡	100,000 units/ml, 48 ml	milliliter
Suspension, oral ‡	100,000 units/ml, 60 ml	milliliter
Suspension, oral ‡	100,000 units/ml, 480 ml	milliliter
Vaginal tablets ‡	15's	each
Vaginal tablets ‡	30's	each
Cream	100,000 units/gm, 15 gm	gram
Cream	100,000 units/gm, 30 gm	gram
Ointment	100,000 units/gm, 15 gm	gram
Ointment	100,000 units/gm, 30 gm	gram
Ointment	100,000 units/gm, 240 gm	gram
Topical powder	«Blank»	gram

**Obinutuzumab ‡ \***

\* «Obinutuzumab is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 50242 (Genentech, Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	25 mg/ml	milliliter

**Ofatumumab ‡ \***

\* «Ofatumumab is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00078 (Novartis Pharmaceuticals Corporation) or 00173 (GlaxoSmithKline) only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	100 mg/5 ml	milliliter
Injection	1000 mg/50 ml	milliliter

**Ofloxacin**

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.3 %	milliliter
Otic solution	0.3 %, 5 ml	milliliter
Otic solution	0.3 %, 10 ml	milliliter
Tablets *	200 mg	each
Tablets *	300 mg	each
Tablets *	400 mg	each

\* «Ofloxacin tablets» are restricted to use in the treatment of sexually transmitted diseases.

**Olanzapine \***

\* «Olanzapine is» restricted to: 1) The use of antipsychotics for Medi-Cal beneficiaries less than 18 years of age requires treatment authorization approval; 2) The use of antipsychotics for Medi-Cal beneficiaries residing in nursing facilities is restricted to FDA approved indications.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	2.5 mg	each
Tablets	5 mg	each
Tablets	7.5 mg	each
Tablets	10 mg	each
Tablets	15 mg	each
Tablets	20 mg	each
Tablets, orally disintegrating	5 mg	each
Tablets, orally disintegrating	10 mg	each
Tablets, orally disintegrating	15 mg	each
Tablets, orally disintegrating	20 mg	each

**Olaparib ‡ \***

\* «Olaparib is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00310 (AstraZeneca LP) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	50 mg	each
Tablets	100 mg	each
Tablets	150 mg	each

**Olaratumab ‡ \***

\* Olaratumab is restricted to use in the treatment of cancer only. Also restricted to labeler codes 00002 (Eli Lilly and Company) only «and effective November 1, 2021, it is restricted to dates of service from January 1, 2017 to November 1, 2021.»

Dosage Form	Size and/or Strength	Billing Unit
Injection	500 mg/50 ml	milliliter
Injection	190 mg/19 ml	milliliter

**Olmesartan Medoxomil**

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	20 mg *	each
Tablets +	40 mg *	each

\* The 20 mg and 40 mg tablets are restricted to claims with dates of service from April 1, 2003, through May 31, 2008, only.

**Olmesartan Medoxomil/Hydrochlorothiazide \***

\* Olmesartan Medoxomil/Hydrochlorothiazide is restricted to claims with dates of service from November 1, 2007, through May 31, 2008, only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	20 mg – 12.5 mg	each
Tablets +	40 mg – 12.5 mg	each
Tablets +	40 mg – 25 mg	each

**Olodaterol HCl \***

\* Olodaterol HCl is NDC labeler code 00597 (Boehringer Ingelheim Pharmaceuticals) only. Effective October 1, 2020

Dosage Form	Size and/or Strength	Billing Unit
Inhaler	2.5 mcg, 4 gm	gram

## Olopatadine HCl \*

\* «Olopatadine HCl is» restricted to NDC labeler code 00065 (Alcon Laboratories, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.1 % *	milliliter
Ophthalmic solution	0.2 % *	milliliter
Ophthalmic solution	0.7 %	milliliter
Nasal spray	0.6 %	grams

\* The ophthalmic solutions 0.1% and 0.2% are restricted to claims with dates of service through June 30, 2016.

## Omacetaxine Mepesuccinate ‡ \*

\* «Omacetaxine Mepesuccinate is» restricted to use in the treatment of cancer «and, effective July 1, 2019, to NDC labeler code 63459 (Cephalon, Inc.) only.»

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	3.5 mg	each

## Ombitasvir/Paritaprevir/Ritonavir and Dasabuvir \*

\* «Ombitasvir/Paritaprevir/Ritonavir and Dasabuvir» requires a *Treatment Authorization Request* (TAR). Restricted to use in the treatment of chronic Hepatitis C Virus (HCV) infection in adults (≥18 years of age). Also restricted to 1) a maximum quantity of 112 tablets per dispensing; and 2) duration of therapy lasting up to 12 or 24 weeks from the dispensing date of the first prescription.

Dosage Form	Size and/or Strength	Billing Unit
Tablets (dose-pack)	12.5 mg/75 mg/50 mg; 250 mg	each

**Note:** Providers must provide documentation of baseline HCV-RNA level and HCV genotype. In addition, when applicable, providers must document relevant clinical information (i.e., failure of prior treatment, presence of cirrhosis, etc.) in support of medical necessity for duration of therapy. Failure to submit supporting documentation may delay authorization of the TAR.

**Note:** “each” means tablet.



## Ombitasvir/Paritaprevir/Ritonavir/Dasabuvir \*

\* Ombitasvir/Paritaprevir/Ritonavir/Dasabuvir requires a *Treatment Authorization Request* (TAR). Restricted to use in the treatment of chronic Hepatitis C Virus (HCV) infection in adults ( $\geq 18$  years of age). Also restricted to 1) a maximum quantity of 84 tablets per dispensing; and 2) duration of therapy lasting up to 12 or 24 weeks from the dispensing date of the first prescription.

Dosage Form	Size and/or Strength	Billing Unit
Tablets, extended release (dose-pack)	8.33 mg/50 mg/33.33 mg/ 200 mg	each

**Note:** "each" means tablet.

**Note:** The following text is added effective February 1, 2021: Product is no longer available.

## Omeprazole

Dosage Form	Size and/or Strength	Billing Unit
Capsules, delayed release	10 mg	each
Capsules, delayed release	20 mg	each
Capsules, delayed release	40 mg	each

Effective May 1, 2019, Omeprazole is a contracted drug.

## «Omeprazole/Amoxicillin/Rifabutin \*

\* Omeprazole/Amoxicillin/Rifabutin is added effective October 1, 2021, and restricted to NDC labeler code 57841 only.

Dosage Form	Size and/or Strength	Billing Unit
Capsules, delayed release	10 mg/250 mg/12.5 mg	each»

## Omeprazole/Sodium Bicarbonate \*

\* Omeprazole/Sodium Bicarbonate is restricted to claims with dates of service from August 1, 2005, through September 30, 2009, only.

Dosage Form	Size and/or Strength	Billing Unit
Powder packet	20 mg	each
Powder packet	40 mg	each
Capsules	20 mg	each
Capsules	40 mg	each

## Ondansetron

«The 4 mg/5 mL liquid in the table below is effective November 1, 2020.»

Dosage Form	Size and/or Strength	Billing Unit
Injection + *	2 mg/ml, 2 ml	milliliter
Tablets +	4 mg	each
Tablets +	8 mg	each
Tablets, orally disintegrating +	4 mg	each
Tablets, orally disintegrating +	8 mg	each
«Liquid	4 mg/5mL	milliliter»

\* The 2 mg/ml, 2 ml injection is restricted to a maximum of 16 mg per dispensing.

## Oprelvekin

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	5 mg/vial	each

**Note:** This product is no longer manufactured or available.

## Oseltamivir Phosphate ‡

Dosage Form	Size and/or Strength	Billing Unit
Capsules	30 mg	each
Capsules	45 mg	each
Capsules	75 mg	each
Oral suspension	6 mg/ml, 60 ml	milliliter

## Osimertinib ‡ \*

\* Osimertinib is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00310 (AstraZeneca LP) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	40 mg	each
Tablets	80 mg	each

**Oxaliplatin ‡ \***

\* «Oxaliplatin is» restricted to use in the treatment of cancer only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	5 mg/ml	milliliter

**Oxandrolone**

Dosage Form	Size and/or Strength	Billing Unit
Tablets *	2.5 mg	each
Tablets *	10 mg	each

\* «The 2.5 mg tablets are» restricted to use in patients with AIDS wasting for claims submitted with dates of service from March 1, 2001, through May 31, 2003.

\* «The 10 mg tablets require authorization» for claims submitted with dates of service on or after September 1, 2002.

**Oxcarbazepine \***

\* Use «of Oxcarbazepine» in beneficiaries less than 2 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	150 mg	each
Tablets	300 mg	each
Tablets	600 mg	each

**Oxiconazole Nitrate \***

\* «Oxiconazole Nitrate is» restricted to NDC labeler code 00462 (PharmaDerm) only with dates of service through November 30, 2012, for all dosage forms.

Dosage Form	Size and/or Strength	Billing Unit
Cream	1%, 15 gm	gram
Cream	1%, 30 gm	gram
Cream	1%, 60 gm	gram
Lotion	1%, 30 ml	milliliter

## Oxybutynin \*

\* «Oxybutynin is» restricted to NDC labeler codes 52544 (Watson Laboratories, Inc.) and 00023 (Allergan, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Transdermal system	3.9 mg	each

## Oxybutynin Chloride

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	5 mg	each
Tablets, extended release + *	5 mg	each
Tablets, extended release + *	10 mg	each
Tablets, extended release + *	5 mg	each

\* «Extended release tablets are» restricted to NDC labeler code 17314 (Alza Corporation) and to claims submitted with dates of service from December 1, 1998, through December 31, 2008, only.

## Oxycodone and Acetaminophen \*

«The following text is removed effective August 1, 2020: Oxycodone and Acetaminophen» are restricted to a maximum quantity per dispensing of 20 tablets or capsules and a maximum of three (3) dispensings in any 75-day period.

**Note:** 5 mg to 500 mg is no longer manufactured or available.»

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules	5 mg to 500 mg	each

«End of removed text.

\* The following text is effective August 1, 2020: Oxycodone and Acetaminophen» are restricted to a maximum quantity per dispensing of «ninety (90) tablets» and a maximum of three (3) dispensings in any 75-day period.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg to 325 mg	each
Tablets	7.5 mg to 325 mg	each
Tablets	10 mg to 325 mg	each

## Oxycodone HCl

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules *	5 mg	each
Tablets or capsules *	15 mg	each
Tablets or capsules *	30 mg	each
Tablets, controlled release *	10 mg	each
Tablets, controlled release *	20 mg	each
Tablets, controlled release *	40 mg	each
Tablets, controlled release *	80 mg	each
Tablets, controlled release *	160 mg	each
Solution	«Blank»	milliliter
Concentrate	«Blank»	milliliter

\* «The 5 mg, 15 mg and 30 mg tablets or capsules are» restricted to a maximum of 90 tablets or capsules per dispensing and one dispensing every 25 days. Exceptions to this restriction require authorization.

\* «The controlled release tablets are» restricted to a maximum of 90 tablets per dispensing and a maximum of three (3) dispensings of any strength in a 75-day period and restricted to claims with dates of service from July 1, 1996, through August 31, 2008, only. Exceptions to this restriction require authorization.

## Oxycodone HCl and Aspirin

Dosage Form	Size and/or Strength	Billing Unit
Tablets *	4.8355 mg – 325 mg	each

\* «The tablets are» restricted to a maximum dispensing quantity of 120 tablets and a maximum of three (3) dispensings in any 75-day period.

## Oxycodone HCl with Oxycodone Terephthalate and Aspirin \*

\* «Oxycodone HCl with Oxycodone Terephthalate and Aspirin is» restricted to a maximum dispensing quantity of 120 tablets and a maximum of three (3) dispensings in any 75-day period.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	2.25 mg – 0.19 mg – 325 mg	each
Tablets	4.5 mg – 0.38 mg – 325 mg	each

**Note:** These products are no longer manufactured or available.

## Oxymorphone

Dosage Form	Size and/or Strength	Billing Unit
Ampule	1 mg/ml, 1 ml	milliliter
Ampule	1.5 mg/ml, 1 ml	each
Ampule	1.5 mg/ml, 10 ml	milliliter
Suppositories	5 mg	each

## Paclitaxel, Semi-Synthetic ‡

Dosage Form	Size and/or Strength	Billing Unit
Injection	«Blank»	milliliter

**Palbociclib ‡ \***

\* Palbociclib is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00069 (Pfizer, Inc.) only. The addition of tablets to the table is effective April 1, 2020.

Dosage Form	Size and/or Strength	Billing Unit
Capsules and tablets	75 mg	each
Capsules and tablets	100 mg	each
Capsules and tablets	125 mg	each

**Palonosetron HCl**

Dosage Form	Size and/or Strength	Billing Unit
Injection + *	0.25 mg/5 ml	milliliter

\* The 0.25 mg/5 ml injection is restricted to a maximum of 5 ml per dispensing and to NDC labeler code 62856 (Eisai, Inc.) only. «**The following text is added effective August 1, 2021: Also restricted to claims submitted with dates of service through July 31, 2021.**»

**Pamidronate Disodium ‡**

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	Blank	each

**«Pancrelipase \***

\* Added effective January 1, 2021. The following strengths are restricted to NDC labeler code 00032 only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules, delayed release	3,000 USP units of lipase 9,500 USP units of protease 15,000 USP units of amylase	each
Capsules, delayed release	6,000 USP units of lipase 19,000 USP units of protease 30,000 USP units of amylase	each
Capsules, delayed release	12,000 USP units of lipase 38,000 USP units of protease 60,000 USP units of amylase	each
Capsules, delayed release	24,000 USP units of lipase 76,000 USP units of protease 120,000 USP units of amylase	each
Capsules, delayed release	36,000 USP units of lipase 114,000 USP units of protease 180,000 USP units of amylase	each»



**«Pancrelipase (continued) \***

\* Effective January 1, 2021: the following strengths are restricted to NDC labeler code 00023 only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules, delayed release	3,000 USP units of lipase; 10,000 USP units of protease; 14,000 USP units of amylase	each
Capsules, delayed release	5,000 USP units of lipase; 17,000 USP units of protease; 24,000 USP units of amylase	each
Capsules, delayed release	10,000 USP units of lipase; 32,000 USP units of protease; 42,000 USP units of amylase	each
Capsules, delayed release	15,000 USP units of lipase; 47,000 USP units of protease; 63,000 USP units of amylase	each
Capsules, delayed release	20,000 USP units of lipase; 63,000 USP units of protease; 84,000 USP units of amylase	each
Capsules, delayed release	25,000 USP units of lipase; 79,000 USP units of protease; 105,000 USP units of amylase	each
Capsules, delayed release	40,000 USP units of lipase; 126,000 USP units of protease; 168,000 USP units of amylase	each»

## Pancrelipase (Amylase/Lipase/Protease)

Dosage Form	Size and/or Strength	Billing Unit
Tablets	Blank	each
Capsules	Blank	each
Capsules with enteric coated granules	Blank	each
Powder	Blank	gram

## Panitumumab \*

\* The following text is effective October 1, 2020: Panitumumab is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 55513 (Amgen USA, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	20 mg/ml, 5 ml	milliliter
Injection	20 mg/ml, 10 ml	milliliter
Injection	20 mg/ml, 20 ml	milliliter

The following text is removed «effective October 1, 2020:» Panitumumab is restricted to use in the treatment of cancer and to and to claims submitted with dates of service from October 13, 2006, through February 28, 2010, only. Continuing care with a date of service on or after March 1, 2010, is available when the following conditions are met:

- 1) The beneficiary had a paid fee-for-service claim for this drug on or before February 28, 2010; 2) A claim has been submitted and paid within the past 100 days; and
  - 3) The claim being submitted is within 100 days of the date of service of the last paid claim.
- End of removed text.

## Panobinostat ‡ \*

\* Panobinostat is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00078 (Novartis Pharmaceuticals Corporation) or 00173 (GlaxoSmithKline) only.

Dosage Form	Size and/or Strength	Billing Unit
Capsules	10 mg	each
Capsules	15 mg	each
Capsules	20 mg	each

## Pantoprazole Sodium

Dosage Form	Size and/or Strength	Billing Unit
Tablets, delayed release +	20 mg	each
Tablets, delayed release +	40 mg	each

## Papain and Urea \*

\* Papain and Urea is restricted to NDC labeler codes 50484 (Smith & Nephew, Inc.) and 58980 (Stratus Pharmaceuticals) only and with dates of service on or before February 28, 2009.

Dosage Form	Size and/or Strength	Billing Unit
Ointment	Blank	gram

## Papain-Urea-Chlorophyllin Copper Complex Sodium

Dosage Form	Size and/or Strength	Billing Unit
Ointment *	30 gm	gram
Spray *	33 ml	milliliters

\* The 30 gm ointment is restricted to NDC labeler codes 00064 (Healthpoint, Ltd.), 50484 (Smith & Nephew, Inc.) and 58980 (Stratus Pharmaceuticals) only until April 30, 2006.

\* Effective May 1, 2006, the 30 gm ointment is restricted to NDC labeler codes 50484 (Smith & Nephew, Inc.) and 58980 (Stratus Pharmaceuticals) only and with dates of service on or before February 28, 2009.

\* The 33 ml spray is restricted to claims submitted with dates of service from January 1, 2004, through April 30, 2006.

## Paregoric

Dosage Form	Size and/or Strength	Billing Unit
Liquid	Blank	Blank

## Paregoric and Protective

Dosage Form	Size and/or Strength	Billing Unit
Liquid	Blank	milliliters

**Paromomycin Sulfate ‡**

Dosage Form	Size and/or Strength	Billing Unit
Capsules	Blank	each

**Paroxetine HCl \***

\* Use of Paroxetine HCl is in beneficiaries less than 18 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Suspension, oral	10 mg/5 ml	milliliters
Tablets	10 mg	each
Tablets	20 mg	each
Tablets	30 mg	each
Tablets	40 mg	each
Tablets, controlled release *	12.5 mg	each
Tablets, controlled release *	25 mg	each
Tablets, controlled release *	37.5 mg	each

\* The controlled release tablets are restricted to dates of service from October 1, 2002, through December 31, 2011, only. Continuing care with a date of service on or after December 31, 2011, is available when the following conditions are met: 1) The beneficiary had a paid fee-for-service claim for this drug on or before December 31, 2011; 2) A claim has been submitted and paid within the past 100 days; and 3) The claim being submitted is within 100 days of the date of service of the last paid claim.

**Paroxetine Mesylate \***

\* Paroxetine Mesylate is restricted to claims with dates of service from September 1, 2004, through May 31, 2009 only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	10 mg	each
Tablets	20 mg	each
Tablets	30 mg	each
Tablets	40 mg	each

**Pazopanib Hydrochloride ‡ \***

\* Pazopanib Hydrochloride is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00078 (Novartis Pharmaceuticals Corporation).

Dosage Form	Size and/or Strength	Billing Unit
Tablets	200 mg	each

**Pegaspargase ‡**

Dosage Form	Size and/or Strength	Billing Unit
Injection	750 units/ml	milliliter

**Peginterferon Alfa-2A \***

\* Peginterferon Alfa-2A is restricted to use in the treatment of chronic viral Hepatitis B or C infection. Also restricted to therapy lasting up to 48 weeks from the dispensing date of the first prescription for all dosage forms and strengths and to labeler code 00004 (Roche Laboratories Inc.).

Dosage Form	Size and/or Strength	Billing Unit
Injection kit with alcohol pads *	180 mcg/0.5 ml	each kit
Syringes, package of four, without alcohol pads *	180 mcg/0.5 ml	milliliter
Pen injector, package of four *	180 mcg/0.5 ml	milliliter
Pen injector, package of four *	135 mcg/0.5 ml	milliliter
Injection *	180 mcg/ml	milliliter

\* The 180 mcg/0.5 ml injection kit with alcohol pads is restricted to a maximum of one injection kit per dispensing for the 180 mcg/0.5 ml injection kit. Injection kit with alcohol pads 180 mcg/0.5 ml is currently no longer manufactured or available and will not be payable after June 30, 2012.

\* The syringes are restricted to a maximum of 2 ml per dispensing for the 180 mcg/0.5 ml syringes, package of four, without alcohol pads.

\* The pen injectors are restricted to a maximum of 2 ml per dispensing, package of four, for both strengths of the auto injector pens only.

## Peginterferon Alfa-2A \* (continued)

\* The injection is restricted to a maximum of four vials per dispensing for the 180 mcg/ml injection only.

**Note:** Previously, the injection kit (NDC 00004-0352-39) was billed in units of each kit (four 0.5 ml syringes as one kit). As recently as October 2011, new packaging (NDC 00004-0357-30) was released without alcohol pads and therefore is no longer designated as a kit by NCPDP standards. Bill NDC 00004-0357-30 in units of ml (four 0.5 ml syringes, totaling two ml per package).

## Peginterferon Alfa-2B \*

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection kit *	50 mcg/0.5 ml	each
Powder for injection kit *	80 mcg/0.5 ml	each
Powder for injection kit *	120 mcg/0.5 ml	each
Powder for injection kit *	150 mcg/0.5 ml	each
Powder for injection, single dose delivery system *	50 mcg/0.5 ml	each
Powder for injection, single dose delivery system *	80 mcg/0.5 ml	each
Powder for injection, single dose delivery system *	120 mcg/0.5 ml	each
Powder for injection, single dose delivery system *	150 mcg/0.5 ml	each
Lyophilized powder for injection ‡ *	296 mcg (200 mcg deliverable)	each
Lyophilized powder for injection ‡ *	444 mcg (300 mcg deliverable)	each
Lyophilized powder for injection ‡ *	888 mcg (600 mcg deliverable)	each

\* The powder for injection kits and powder for injection, single dose delivery system are restricted to use in the treatment of Hepatitis C. Also restricted to a maximum of four injection kits or single dose delivery systems per dispensing and therapy lasting up to 48 weeks from the dispensing date of the first prescription for the powder for injection, single dose delivery system only. Also restricted to labeler code 00085 (Schering Corporation) only.

\* The lyophilized powder for injection is restricted to use in the treatment of cancer only.

**Pembrolizumab ‡ \***

\* Pembrolizumab is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00006 (Merck & Company, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	50 mg/vial	each
Solution for injection	100 mg/4 ml	milliliter

**Pemetrexed ‡ \***

\* Pemetrexed is restricted to use in the treatment of cancer only. Also restricted to NDC code 00002 (Eli Lilly and Company) only.

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	100 mg/vial	each
Powder for injection	500 mg/vial	each

**Pemirolast Potassium \***

\* Pemirolast Potassium is restricted to claims with dates of service from December 1, 2000, through September 30, 2010, only.

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.1 %, 10 ml	milliliter

**Pemoline \***

\* Pemoline is restricted to use in Attention Deficit Disorder in individuals from 4 years through 16 years of age with a Medi-Cal fee-for-service paid claim for this drug prior to December 1, 2005, and with the claim being submitted within 100 days of the date of service of the last paid claim submitted.

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules	18.75 mg	each
Tablets or capsules	37.5 mg	each
Tablets or capsules	75 mg	each
Tablets (chewable)	37.5 mg	each

**Penbutolol Sulfate**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets +	20 mg	each

**Penicillin G**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Powder for injection	1,000,000 units/vial	each
Powder for injection	5,000,000 units/vial	each
Powder for injection	10,000,000 units/vial	each
Powder for injection	20,000,000 units/vial	each

**Penicillin G Benzathine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	300,000 U/ml, 10 ml	milliliter
Injection	600,000 U/ml, 1 ml	milliliter
Injection	600,000 U/ml, 2 ml	milliliter
Injection	600,000 U/ml, 4 ml	milliliter

**Penicillin G Procaine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	Blank	milliliter



**Penicillin V (K)**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	125 mg	each
Tablets	250 mg	each
Tablets	500 mg	each
Liquid	125 mg/5 ml, 100 ml	milliliter
Liquid	125 mg/5 ml, 150 ml	milliliter
Liquid	125 mg/5 ml, 200 ml	milliliter
Liquid	250 mg/5 ml, 100 ml	milliliter
Liquid	250 mg/5 ml 150 ml	milliliter
Liquid	250 mg/5 ml, 200 ml	milliliter

**Pentamidine ‡ \***

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Powder for injection	300 mg/vial	each
Powder for aerosolized administration *	300 mg/vial	each

\* The powder for aerosolized administration is restricted to the prevention of pneumocystis carinii pneumonia (PCP) and must meet all of the following criteria: 1) Patient is HIV-infected, with a history of PCP or with a CD4 (T4) lymphocyte count less than or equal to 200 cells/mm<sup>3</sup>. 2) Nebulizer system must comply with the specifications in the package insert for the drug product.

**Pentobarbital**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Suppositories	30 mg	each
Suppositories	60 mg	each
Suppositories	120 mg	each
Suppositories	200 mg	each

**Pentosan Polysulfate Sodium**

Dosage Form	Size and/or Strength	Billing Unit
Capsules	100 mg	each

**Pentostatin ‡**

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	10 mg/vial	each

**Pentoxifylline \***

\* Pentoxifylline is restricted to use for patients 65 years of age or older diagnosed with intermittent claudication, or for diabetic patients of any age diagnosed with intermittent claudication.

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules +	400 mg	each

**«Perampanel \***

\* The following text is effective July 1, 2021. Perampanel is restricted to treatment of seizures in patients with epilepsy 4 years of age and older. Also restricted to NDC labeler code 62856 only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	2 mg	each
Tablets	4 mg	each
Tablets	6 mg	each
Tablets	8 mg	each
Tablets	10 mg	each
Tablets	12 mg	each
Oral suspension	0.5 mg/ml	milliliter»

## Pergolide Mesylate

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.05 mg *	each
Tablets	0.25 mg *	each
Tablets	1.0 mg	each

\* The 0.05 and 0.25 mg tablets are restricted to NDC labeler codes 59075 (Elan Pharmaceuticals, Inc./Athena Neuroscience) and 65234 (Valeant Pharmaceuticals North America).

**Note:** These products are no longer manufactured or available.

## Permethrin

Dosage Form	Size and/or Strength	Billing Unit
Cream	5 %, 60 gm	gram

## Perphenazine \*

\* Perphenazine is restricted to: 1) The use of antipsychotics for Medi-Cal beneficiaries less than 18 years of age requires treatment authorization approval; 2) The use of antipsychotics for Medi-Cal beneficiaries residing in nursing facilities is restricted to FDA approved indications.

Dosage Form	Size and/or Strength	Billing Unit
Injection	5 mg/ml, 1 ml	milliliter
Tablets +	2 mg	each
Tablets +	4 mg	each
Tablets +	8 mg	each
Tablets +	16 mg	each
Liquid	16 mg/5 ml	milliliter

**Pertuzumab ‡ \***

\* Pertuzumab is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 50242 (Genentech USA, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	420 mg/14 ml	milliliter

**Pertuzumab, Trastuzumab and Hyaluronidase-zzxf ‡ \***

« \* Effective, April 1, 2019, **Pertuzumab, Trastuzumab and Hyaluronidase-zzxf is** restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 50242 (Genentech, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Subcutaneous Injection	600 mg/600 mg/ 20,000 units/ 10 ml	milliliter
Subcutaneous Injection	1200 mg/600 mg/ 30,000 units/ 15 ml	milliliter

**Pexidartinib ‡ \***

\* Effective, October 1, 2019, Pexidartinib is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 65597 (Daiichi Sankyo, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Capsules	200 mg	each

**Phenazopyridine HCl**

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.1 gm	each
Tablets	0.2 gm	each

## Phenobarbital

Dosage Form	Size and/or Strength	Billing Unit
Injection	120 to 130 mg/ml, 1 ml	milliliter
Tablets +	15 mg	each
Tablets +	16.2 mg	each
Tablets +	30 mg	each
Tablets +	32.4 mg	each
Tablets +	60 mg	each
Tablets +	65 mg	each
Tablets +	97.2 mg	each
Tablets +	100 mg	each
Liquid	20 mg/5 ml	milliliter

## Phenylephrine

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic solution	0.12 %, 15 ml	milliliter
Ophthalmic solution	2.5 % ,15 ml	milliliter
Ophthalmic solution	10 %, 5 ml	milliliter
Ophthalmic solution	10 %, 15 ml	milliliter

## Phenytoin

Dosage Form	Size and/or Strength	Billing Unit
Tablets, chewable +	50 mg	each
Capsules, extended release +	30 mg	each
Capsules, extended release +	100 mg, 1,000's √	each
Capsules, extended release +	200 mg, 1,000's √	each

**Phenytoin (continued)**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules, extended release +	300 mg, 1000's y	each
Capsules, prompt +	100 mg, 1,000's y	each
Suspension	125 mg/5 ml	milliliter
Suspension	30 mg/5 ml	milliliter

**Phenytoin with Phenobarbital**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets, or capsules +	100 mg/15 mg	each
Tablets, or capsules +	100 mg/30 mg	each

**Note:** These products are no longer manufactured or available.

**Phytonadione**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	10 mg/ml, 1 ml	milliliter
Injection	10 mg/ml, 2.5 ml	milliliter
Injection	10 mg/ml, 5 ml	milliliter
Tablets	5 mg	each

**Pilocarpine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Ophthalmic gel	4 %, 3.5 gm	gram
Ophthalmic gel	4 %, 5 gm	gram
Ophthalmic solution	¼ %, 15 ml	milliliter
Ophthalmic solution	½ %, 15 ml	milliliter
Ophthalmic solution	½ %, 30 ml	milliliter
Ophthalmic solution	1 %, 15 ml	milliliter
Ophthalmic solution	1 %, 30 ml	milliliter

**Pilocarpine (continued)**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Ophthalmic solution	2 %, 15 ml	milliliter
Ophthalmic solution	2 %, 30 ml	milliliter
Ophthalmic solution	3 %, 15 ml	milliliter
Ophthalmic solution	3 %, 30 ml	milliliter
Ophthalmic solution	4 %, 15 ml	milliliter
Ophthalmic solution	4 %, 30 ml	milliliter
Ophthalmic solution	5 %, 15 ml	milliliter
Ophthalmic solution	6 %, 15 ml	milliliter
Ophthalmic solution	6 %, 30 ml	milliliter
Ophthalmic solution	8 % or 10 %, 30 ml	milliliter
Tablets *	5 mg	each
Tablets *	7.5 mg	each

\* The 5 mg and 7.5 mg tablets are restricted to NDC labeler code 58063 (MGI Pharma) and to claims with dates of service from November 1, 2000, through April 30, 2010, for tablets only.

**Pilocarpine with Epinephrine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Ophthalmic solution	1 %, 10 ml	milliliter
Ophthalmic solution	1 %, 15 ml	milliliter
Ophthalmic solution	2 %, 10 ml	milliliter
Ophthalmic solution	2 %, 15 ml	milliliter
Ophthalmic solution	3 %, 10 ml	milliliter
Ophthalmic solution	3 %, 15 ml	milliliter
Ophthalmic solution	4 %, 10 ml	milliliter
Ophthalmic solution	4 %, 15 ml	milliliter
Ophthalmic solution	6 %, 10 ml	milliliter
Ophthalmic solution	6 %, 15 ml	milliliter

**Note:** These products are no longer manufactured or available.

**Pindolol**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets +	5 mg	each
Tablets +	10 mg	each

**Pioglitazone HCl**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	15 mg	each
Tablets	30 mg	each
Tablets	45 mg	each

**Pioglitazone HCl/Glimepiride**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	30 mg/2 mg	each
Tablets	30 mg/4 mg	each

**Pioglitazone HCl/Metformin HCl**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets +	15 mg/500 mg	each
Tablets +	15 mg/850 mg	each

**Piperacillin Sodium**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Powder for injection	Blank	each



**Pipobroman ‡**

Dosage Form	Size and/or Strength	Billing Unit
Tablets	10 mg	each
Tablets	25 mg	each

**Note:** These products are no longer manufactured or available.

**Pirbuterol Acetate**

Dosage Form	Size and/or Strength	Billing Unit
Aerosol inhaler with adapter *	14 gm	gram
Aerosol inhaler with adapter *	25.6 gm	gram

\* The aerosol inhaler with adapter is restricted to dates of service from March 1, 1994, to January 31, 2007.

**Piroxicam**

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules +	10 mg	each
Tablets or capsules +	20 mg	each

**Pitavastatin Calcium \***

\* Pitavastatin Calcium is restricted to claims submitted with dates of service from October 1, 2011, through October 31, 2014, only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	1 mg	each
Tablets	2 mg	each
Tablets	4 mg	each

**Plicamycin ‡**

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	2.5 mg/vial	each

**Note:** This product is no longer manufactured or available.

## Pneumococcal Vaccine, 13-Valent, Conjugated \*

\* «Pneumococcal Vaccine, 13-Valent, Conjugated is» restricted to: 1) Medi-Cal beneficiaries 19 years of age and older. 2) One dose of vaccine per lifetime. 3) Use of this vaccine must be based on the guidelines published by the Centers for Disease Control and Prevention (CDC).

Dosage Form	Size and/or Strength	Billing Unit
Injection	1-dose syringe	milliliter

## Pneumococcal Vaccine, 23-Valent, Non-Conjugated \*

\* «Pneumococcal Vaccine, 13-Valent, Non-Conjugated is» restricted to: 1) Medi-Cal beneficiaries 19 years of age and older. 2) Two doses of vaccine per lifetime. 3) Use of this vaccine must be based on the guidelines published by the Centers for Disease Control and Prevention (CDC).

Dosage Form	Size and/or Strength	Billing Unit
Injection	1-dose vial	milliliter
Injection	1-dose syringe	milliliter

## Podofilox

Dosage Form	Size and/or Strength	Billing Unit
Topical solution	0.5 %	milliliter
Topical gel	0.5 %	gram

**Polatuzumab Vedotin-piiq ‡ \***

\* «Effective June 11, 2019, Polatuzumab Vedotin-piiq is» restricted to use in the treatment of cancer only. Also restricted to labeler codes 50242 (Genentech, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	140 mg	each

**Polyestradiol Phosphate ‡**

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	40 mg/vial	each

**Note:** This product is no longer manufactured or available.

**Poethylene Glycol 3350**

«The following text is removed effective June 1, 2020.»

Dosage Form	Size and/or Strength	Billing Unit
Powder	17 gm	gram

«End of removed text. The following text is effective June 1, 2020.»

Dosage Form	Size and/or Strength	Billing Unit
Powder	238 gm	gram
Powder	510 gm	gram

**Poethylene Glycol 3350 and Electrolytes**

Dosage Form	Size and/or Strength	Billing Unit
Solution	4000 ml	milliliter

**Polymyxin, Bacitracin**

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic ointment	Blank	gram

**Ponatinib ‡ \***

\* Ponatinib is restricted to use in the treatment of cancer only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	15 mg	each
Tablets	45 mg	each

**Porfimer Sodium ‡**

Dosage Form	Size and/or Strength	Billing Unit
Powder for injection	Blank	each

**Potassium Bicarbonate/Citric Acid**

Dosage Form	Size and/or Strength	Billing Unit
Tablets, effervescent +	10 meq	each
Tablets, effervescent +	20 meq	each
Tablets, effervescent +	25 meq	each

«The following text is removed effective June 1, 2021: The effervescent tablets exclude NDC labeler codes 00245 (Upsher-Smith Laboratories, Inc.) and 66758 (Parenta Pharmaceuticals, Inc.). End of removed text.»

## Potassium Chloride

Dosage Form	Size and/or Strength	Billing Unit
Tablets, long acting +	8 meq	each
Injection	Blank	milliliters
Liquid	10 %	milliliters
Liquid	20 %	milliliters

The following text is effective May 1, 2020:

Dosage Form	Size and/or Strength	Billing Unit
Capsules, long acting +	8 meq	each
Capsules, long acting +	10 meq	each
Tablets, long acting +	10 meq	each
Tablets, long acting +	20 meq	each

«The following text is removed effective June 1, 2021: Effective May 1, 2020, the 8 meq, 10 meq, and 20 meq long acting tablets exclude NDC labeler codes 00074 (Abbvie, Inc.), 00245 (Upsher-Smith Laboratories, Inc.) and 66758 (Parenta Pharmaceuticals, Inc.). End of removed text.»

**Note:** Payment for oral liquid limited to a minimum dispensing quantity of 480 ml. See *California Code of Regulations (CCR)*, Title 22, Section 51513(b)(5) regarding exceptions.

## Potassium Citrate

The following text is effective May 1, 2020:

Dosage Form	Size and/or Strength	Billing Unit
Tablets, extended release +	5 meq	each
Tablets, extended release +	10 meq	each

## Potassium Iodide Saturated Solution

Dosage Form	Size and/or Strength	Billing Unit
Liquid	Blank	milliliters

## Pralatrexate ‡ \*

\* Pralatrexate is restricted to use in the treatment of cancer and restricted to claims submitted with dates of service from October 28, 2009, through September 30, 2014, only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	20 mg/1 ml	milliliters
Injection	40 mg/2 ml	milliliters

## Pralsetinib ‡ \*

\* Effective September 14, 2020, Pralsetinib is restricted to use in the treatment of cancer. Also restricted to NDC labeler code «50242, effective July 1, 2021, and» 72064. «Effective July 1, 2021, the following text is removed: (Blueprint Medicines Corporation). End of removed text.»

Dosage Form	Size and/or Strength	Billing Unit
Capsules	100 mg	each

## Pramipexole Dihydrochloride \*

\* Use of Pramipexole Dihydrochloride in beneficiaries less than 18 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.125 mg	each
Tablets	0.25 mg	each
Tablets	0.5 mg	each
Tablets	1.0 mg	each
Tablets	1.5 mg	each
Tablets, extended-release	0.375 mg	each
Tablets, extended-release	0.75 mg	each
Tablets, extended-release	1.5 mg	each
Tablets, extended-release	3.0 mg	each
Tablets, extended-release	4.5 mg	each

«Effective April 1, 2021, the following text is removed: The extended-release tablets are restricted to NDC labeler code 00597 (Boehringer Ingelheim Pharmaceuticals) only. End of removed text.»

## Pramlintide Acetate \*

\* Pramlintide Acetate is restricted to use in the treatment of Type 2 diabetes and labeler code 00310 (AstraZeneca LP) only.

Dosage Form	Size and/or Strength	Billing Unit
60 Pen injector	1.5 ml	milliliter
120 Pen injector	2.7 ml	milliliter

## Prasugrel \*

\* Prasugrel is restricted to NDC labeler code 00002 (Eli Lilly and Company) and to claims with dates of service from April 1, 2011, through May 31, 2017, only. Continuing care with a date of service on or after June 1, 2017, is available when the following conditions are met: 1) The beneficiary had a paid fee-for-service claim for this drug on or before May 31, 2017; 2) A claim has been submitted and paid within the past 100 days; and 3) The claim being submitted is within 100 days of the date of service of the last paid claim.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg	each
Tablets	10 mg	each

## Pravastatin

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	10 mg	each
Tablets +	20 mg	each
Tablets +	40 mg	each
Tablets +	80 mg	each

## Prazosin HCl

Dosage Form	Size and/or Strength	Billing Unit
Capsules +	1 mg	each
Capsules +	2 mg	each
Capsules +	5 mg	each

## Prednicarbate

Dosage Form	Size and/or Strength	Billing Unit
Cream	0.1 %, 15 gm	gram
Cream	0.1 %, 60 gm	gram



## Prednisolone

Dosage Form	Size and/or Strength	Billing Unit
Injection	20 mg/ml, 2 ml	milliliter
Injection	20 mg/ml, 5 ml	milliliter
Injection	20 mg/ml, 10 ml	milliliter
Injection	25 mg/ml, 10 ml	milliliter
Injection	25 mg/ml, 30 ml	milliliter
Tablets	5 mg	each
Ophthalmic solution	0.12 % - 0.125 %, 5 ml	milliliter
Ophthalmic solution	0.12 % - 0.125 %, 10 ml	milliliter
Ophthalmic solution	1.0 %, 5 ml	milliliter
Ophthalmic solution	1.0 %, 10 ml	milliliter
Ophthalmic solution	1.0 %, 15 ml	milliliter
Liquid	5 mg/5 ml	milliliter
Liquid	15 mg/5 ml	milliliter

## Prednisolone, Neomycin, Polymyxin B

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic suspension	5 mg/5 mg/10,000 U/ml, 5 ml	milliliter
Ophthalmic suspension	5 mg/5 mg/10,000 U/ml, 10 ml	milliliter

**Note:** These products are no longer manufactured or available.

## Prednisolone Sodium Phosphate

Dosage Form	Size and/or Strength	Billing Unit
Oral solution *	20.2 mg/5 ml	milliliter
Orally disintegrating tablets	10 mg	each
Orally disintegrating tablets	15 mg	each
Orally disintegrating tablets	30 mg	each

\* The oral solution is restricted to labeler code 68135 (Biomarin Pharmaceuticals, Inc.) only and to claims with dates of service from April 1, 2005, through September 30, 2008, only.

## Prednisolone with Sulfacetamide

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic ointment	0.25 %	gram
Ophthalmic ointment	0.5 %	gram
Ophthalmic solution	0.2 % - 0.25 %, 5 ml	milliliter
Ophthalmic solution	0.2 % - 0.25 %, 10 ml	milliliter
Ophthalmic solution	0.2 % - 0.25 %, 15 ml	milliliter
Ophthalmic solution	0.5 %, 5 ml	milliliter
Ophthalmic solution	0.5 %, 15 ml	milliliter

## Prednisone

Dosage Form	Size and/or Strength	Billing Unit
Tablets	1 mg	each
Tablets	2.5 mg	each
Tablets	5 mg	each
Tablets	10 mg	each
Tablets	20 mg	each
Tablets	50 mg	each

## «Pregabalin

The following text is effective September 1, 2020:

Dosage Form	Size and/or Strength	Billing Unit
Capsules	25 mg	each
Capsules	50 mg	each
Capsules	75 mg	each
Capsules	100 mg	each
Capsules	150 mg	each»

**«Pregabalin (continued)**

The following text is effective September 1, 2020:

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	200 mg	each
Capsules	225 mg	each
Capsules	300 mg	each»

**Primaquine ‡**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	26.3 mg	each

**Primidone**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets +	50 mg	each
Tablets +	250 mg	each
Liquid	0.25 gm/5 ml	milliliter

**Probenecid**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets +	500 mg	each

**Probenecid with Colchicine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets +	«Blank»	each

**Procainamide**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	100 mg/ml, 10 ml	milliliter
Capsules or tablets +	250 mg	each
Capsules or tablets +	375 mg	each
Capsules or tablets +	500 mg	each
Capsules or tablets, long-acting +	250 mg	each
Capsules or tablets, long-acting +	500 mg	each
Capsules or tablets, long-acting +	750 mg	each
Capsules or tablets, long-acting +	1000 mg	each

**Procarbazine ‡**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	50 mg	each

**Prochlorperazine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	5 mg/ml	milliliter
Injection, prefilled syringe	«Blank»	milliliter
Tablets +	5 mg	each
Tablets +	10 mg	each
Tablets +	25 mg	each
Liquid	5 mg/5 ml	
Capsules, sustained release +	10 mg	each
Capsules, sustained release +	15 mg	each

**Prochlorperazine (continued)**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules, sustained release +	30 mg	each
Suppositories	2.5 mg	each
Suppositories	5 mg	each
Suppositories	25 mg	each

**Procyclidine**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	5 mg	each

**Note:** This product is no longer manufactured or available.

**Progesterone**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	50 mg/ml, 10 ml	milliliter

**Promethazine \***

\* «Promethazine is» restricted to individuals 2 years of age or older.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	25 mg/ml, 1 ml	milliliter
Injection	25 mg/ml, 10 ml	milliliter
Injection	50 mg/ml	milliliter
Tablets +	12.5 mg	each
Tablets +	25 mg	each
Tablets +	50 mg	each
Liquid	6.25 mg/5 ml	milliliter
Liquid Fortis	25 mg/5 ml	milliliter

**Promethazine\* (continued)**

\* «Promethazine is» restricted to individuals 2 years of age or older.

Dosage Form	Size and/or Strength	Billing Unit
Suppositories	12.5 mg	each
Suppositories	25 mg	each
Suppositories	50 mg	each

**Promethazine with Codeine \***

\* «Promethazine with Codeine is» restricted to individuals 2 years of age and older. Also restricted to a maximum dispensing quantity of 360 ml and a maximum of three (3) dispensings in any 75-day period.

Dosage Form	Size and/or Strength	Billing Unit
Liquid	«Blank»	milliliter

**Promethazine with Dextromethorphan \***

\* «Promethazine with Dextromethorphan is» restricted to individuals 2 years of age and older.

Dosage Form	Size and/or Strength	Billing Unit
Liquid	«Blank»	milliliter

**Promethazine with Phenylephrine \***

\* «Promethazine with Phenylephrine is» restricted to individuals 2 years of age and older.

Dosage Form	Size and/or Strength	Billing Unit
Liquid	«Blank»	milliliter

**Promethazine with Phenylephrine and Codeine \***

\* «Promethazine with Phenylephrine and Codeine is» restricted to individuals 2 years of age and older. Also restricted to a maximum dispensing quantity of 360 ml and a maximum of three (3) dispensings in any 75-day period.

Dosage Form	Size and/or Strength	Billing Unit
Liquid	«Blank»	milliliter

## Proprantheline Bromide

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	7.5 mg	each
Tablets +	15 mg	each

## Proparacaine HCl

Dosage For	Size and/or Strength	Billing Unit
Ophthalmic solution	0.5 %, 2 ml	milliliter
Ophthalmic solution	0.5 %, 15 ml	milliliter

## Propranolol

Dosage Form	Size and/or Strength	Billing Unit
Injection	1 mg/ml, 1 ml	milliliter
Tablets +	10 mg	each
Tablets +	20 mg	each
Tablets +	40 mg	each
Tablets +	60 mg	each
Tablets +	80 mg	each
Tablets +	90 mg	each
Liquid	4 mg/ml	milliliter
Liquid	8 mg/ml	milliliter

## «Propranolol Hydrochloride \*

\* Added effective April 1, 2021: Propranolol Hydrochloride is restricted to use in the treatment of proliferating infantile hemangioma. Also restricted to NDC labeler code 64370.

Dosage Form	Size and/or Strength	Billing Unit
Oral Solution	4.28mg/ml	milliliter»»

## Propylthiouracil

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	50 mg	each

## Protriptyline HCl

\* Use «of Protriptyline HCl» in beneficiaries less than 12 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg	each
Tablets	10 mg	each

## Pyrantel Pamoate

Dosage Form	Size and/or Strength	Billing Unit
Liquid	«Blank»	each

## Pyrazinamide

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules	500 mg	each

## Pyridostigmine

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	60 mg	each
Tablets, long acting +	180 mg	each
Liquid	«Blank»	milliliter



## Pyridoxine

Dosage Form	Size and/or Strength	Billing Unit
Injection	100 mg/ml, 10 ml	milliliter
Injection	100 mg/ml, 30 ml	milliliter

## Pyrimethamine ‡

Dosage Form	Size and/or Strength	Billing Unit
Tablets	25 mg	each

## Quetiapine Fumarate \*

\* «Quetiapine Fumarate is» restricted to: 1) The use of antipsychotics for Medi-Cal beneficiaries less than 18 years of age requires treatment authorization approval; 2) The use of antipsychotics for Medi-Cal beneficiaries residing in nursing facilities is restricted to FDA approved indications.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	25 mg	each
Tablets	50 mg	each
Tablets	100 mg	each
Tablets	200 mg	each
Tablets	300 mg	each
Tablets	400 mg	each
Extended-release tablets	50 mg	each
Extended-release tablets	150 mg	each
Extended-release tablets	200 mg	each
Extended-release tablets	300 mg	each
Extended-release tablets	400 mg	each

**Quinapril HCl \* ♦**

\* Authorization is always required for «Quinapril HCl.» ♦

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg	each ♦
Tablets	10 mg	each ♦
Tablets	20 mg	each ♦
Tablets	40 mg	each ♦

«Quinapril HCl is» suspended until further notice.

**Quinidine Gluconate**

Dosage Form	Size and/or Strength	Billing Unit
Injection	80 mg/ml, 10 ml	milliliter
Tablets, long acting +	324 mg	each

**Quinidine Sulfate**

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	100 mg	each
Tablets +	200 mg, 1,000's †	each
Tablets +	300-325 mg	each
Tablets or capsules	«Blank»	each

**Quinine Sulfate \***

\* «Quinine Sulfate is» restricted to claims submitted with dates of service prior to May 1, 2007.

Dosage Form	Size and/or Strength	Billing Unit
Tablets or capsules	«Blank»	each

**Rabies Vaccine \***

\* «Rabies Vaccine is» restricted to: 1) Medi-Cal beneficiaries 19 years of age and older.  
2) Use of this vaccine must be based on the guidelines published by the Centers for Disease Control and Prevention (CDC).

Dosage Form	Size and/or Strength	Billing Unit
Injection	1-dose vial	each

**Raloxifene HCl \***

\* «Raloxifene HCl is» restricted to use for the treatment and prevention of osteoporosis and reduction in the risk of invasive breast cancer in postmenopausal women with osteoporosis or postmenopausal women at high risk of invasive breast cancer and to claims submitted with dates of service from October 1, 1998, through June 30, 2009, only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	60 mg	each

**Raltegravir ‡ \***

\* «Raltegravir is» restricted to use as combination therapy in the treatment of Human Immunodeficiency Virus (HIV) infection. Also restricted to NDC labeler code 00006 (Merck & Company, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	400 mg	each
Tablets	600 mg	each
Chewable tablets	25 mg	each
Chewable tablets	100 mg	each
Oral Suspension packets	100 mg	each

**Ramelteon \***

\* Ramelteon is restricted to use in the treatment of insomnia.

\* The following text is effective December 1, 2020: Ramelteon is restricted to a maximum quantity per dispensing of 60 tablets in 30 days. Use in beneficiaries less than 18 years of age requires treatment authorization approval.

The following text is removed «effective December 1, 2020:» and to a maximum dispensing quantity of thirty (30) tablets. and a maximum of three (3) dispensings in any seventy-five (75) day period and to claims with dates of service from November 1, 2006, through March 31, 2010, only. End of removed text.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	8 mg	each

**Ramipril**

Dosage Form	Size and/or Strength	Billing Unit
Capsules +	1.25 mg	each
Capsules +	2.5 mg	each
Capsules +	5 mg	each
Capsules +	10 mg	each

**Ramucirumab ‡ \***

\* Ramucirumab is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00002 (Eli Lilly and Company) only.

Dosage Form	Size and/or Strength	Billing Unit
Injection	100 mg/10 ml	milliliter
Injection	500 mg/50 ml	milliliter

**Ranitidine HCl**

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	150 mg	each
Tablets +	300 mg	each

The following text is effective April 1, 2020:

Dosage Form	Size and/or Strength	Billing Unit
Syrup	15 mg/ml	milliliters

**Regorafenib ‡ \***

\* Regorafenib is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 50419 (Bayer HealthCare Pharmaceuticals) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	40 mg	each

**«Relugolix ‡ \***

\* Added effective January 21, 2021: Relugolix is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 72974 only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	120 mg	each»

**Repaglinide\***

\* Repaglinide is restricted to claims submitted with dates of service from April 1, 1999, through July 31, 2005.

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	0.5 mg	each
Tablets +	1.0 mg	each
Tablets +	2.0 mg	each

## Ribavirin \*

\* «Ribavirin is» restricted to use as combination therapy in the treatment of Hepatitis C. Also restricted to therapy lasting up to 48 weeks from the dispensing date of the first prescription.

Dosage Form	Size and/or Strength	Billing Unit
Capsules	200 mg	each
Tablets	200 mg	each
Dose Pack Tablets (56 tablets per pack) *	600 mg Dose Pack (200 mg x 28, 400 mg x 28)	each
Dose Pack Tablets (56 tablets per pack) *	800 mg Dose Pack (400 mg x 56)	each
Dose Pack Tablets (56 tablets per pack) *	1000 mg Dose Pack (400 mg x 28, 600 mg x 28)	each
Dose Pack Tablets (56 tablets per pack) *	1200 mg Dose Pack (500 mg x 56)	each

\* «The Dose Pack Tablets is» restricted to brand name Ribasphere RibaPak labeler code 66435 (Kadmon Pharmaceuticals, LLC) for Dose Pack (tablets) only. Also restricted to claims with dates of service from July 1, 2012, through June 30, 2015. Continuing care with a date of service on or after July 1, 2015, is available when the following conditions are met: 1) The beneficiary had a paid fee-for-service claim for this drug on or before June 30, 2015; 2) A claim has been submitted and paid within the past 100 days; and 3) The claim being submitted is within 100 days of the date of service of the last paid claim.

**Note:** “each” means tablet. Bill using outer package NDCs ending in -99 for proper reimbursement.

## Ribavirin and Interferon Alfa-2B \*

\* «Ribavirin and Interferon Alfa-2B are» restricted to use in the treatment of Hepatitis C. Also restricted to a maximum quantity of two kits per dispensing, to therapy lasting up to 48 weeks from the dispensing date of the first prescription and to dates of service from July 1, 1999, through June 30, 2005.

Dosage Form	Size and/or Strength	Billing Unit
Capsules and injection, multi-dose pen +	«Blank»	each kit

**Note:** Product is no longer manufactured or available.

**Ribociclib ‡ \***

\* «Ribociclib is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00078 (Novartis Pharmaceuticals Corporation) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	600 mg daily dose (3 x 21 tablet blister packs)	each
Tablets	400 mg daily dose (3 x 14 tablet blister packs)	each
Tablets	200 mg daily dose (1 x 21 tablet blister packs)	each

**Note:** “each” means number of tablets per box of either 63, 42 or 21.

**Ribociclib and Letrozole ‡ \***

\* «Ribociclib and Letrozole is» restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 00078 (Novartis Pharmaceuticals Corporation) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	600 mg daily dose (3 x 21 tablet blister packs) and 2.5 mg (bottle of 28 tablets)	each
Tablets	400 mg daily dose (3 x 14 tablet blister packs) and 2.5 mg (bottle of 28 tablets)	each
Tablets	200 mg daily dose (1 x 21 tablet blister packs) and 2.5 mg (bottle of 28 tablets)	each

**Note:** “each” means total number of tablets carton of either 91, 70 or 49.

**Rifabutin ‡ \***

\* «Rifabutin is» restricted to use in the prevention of disseminated Mycobacterium Avium Complex (MAC) disease in patients with advanced HIV infection.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	150 mg	each

**Rifampin**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	150 mg	each
Capsules	300 mg	each

**Rifampin and Isoniazid**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	300 mg/150 mg	each

**Rifampin and Isoniazid and Pyrazinamide**

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	120 mg/50 mg/300 mg	each



**Rifapentine**

Dosage Form	Size and/or Strength	Billing Unit
Tablets	150 mg	each

**Rilpivirine ‡ \***

\* «Rilpivirine is» restricted to use as combination therapy in the treatment of Human Immunodeficiency Virus (HIV) infection only. Also restricted to NDC labeler code 59676 (Janssen Products, LP.) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	25 mg	each

**Riluzole \***

\* «Riluzole is» restricted to use in the treatment of amyotrophic lateral sclerosis.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	50 mg	each

**Rimegepant \***

«The following text is effective August 1, 2020:»

\* «Rimegepant» requires a *Treatment Authorization Request (TAR)*. Rimegepant is restricted to 1) Use in patients who have failed or are unable to tolerate a drug in the triptan class of medication; 2) Acute treatment of migraine headache; 3) Maximum fill quantity of 8 tablets per dispensing and one dispensing in 30 days. Also restricted to NDC labeler code 72618 (Biohaven Pharmaceuticals, Inc.) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets, orally disintegrating	75 mg	each»

**Rimexolone \***

\* Rimexolone is restricted to NDC labeler code 00065 (Alcon Laboratories, Inc.) only and to claims submitted with dates of service through September 29, 2018, only.

Dosage Form	Size and/or Strength	Billing Unit
Ophthalmic suspension	1 %	milliliter

**Ripretinib ‡ \***

\* <<Ripretinib is added effective October 1, 2020:>> it is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 73207 (Deciphera Pharmaceuticals, LLC) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	50 mg	each

**Risedronate Sodium**

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg	each
Tablets	30 mg	each
Tablets	35 mg	each
Tablets *	150 mg	each

\* Restricted to claims with dates of service from January 1, 2009, through April 30, 2012, for the 150 mg tablets only.

**Risperidone \***

\* Risperidone is restricted to: 1) The use of antipsychotics for Medi-Cal beneficiaries less than 18 years of age requires treatment authorization approval; 2) The use of antipsychotics for Medi-Cal beneficiaries residing in nursing facilities is restricted to FDA approved indications.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	0.25 mg	each
Tablets	0.5 mg	each
Tablets	1 mg	each
Tablets	2 mg	each
Tablets	3 mg	each
Tablets	4 mg	each
Solution	1 mg/ml	milliliter

**Ritonavir ‡ \***

\* Ritonavir is restricted to use as combination therapy in the treatment of Human Immunodeficiency Virus (HIV) infection. Also restricted to NDC labeler code 00074 (AbbVie Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Capsules	100 mg	each
Tablets	100 mg	each
Solution	80 mg/ml	milliliter
Oral powder packets	100 mg	each

**Rituximab ‡ \***

\* Rituximab is restricted to use in the treatment of cancer. Also restricted to NDC labeler code 50242 (Genentech, Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	10 mg/ml	milliliter

**Rituximab-abbs ‡ \***

\* Rituximab is restricted to use in the treatment of cancer. Also restricted to NDC labeler 63459 (Teva Pharmaceuticals USA, Inc.) only. Drug addition and dosages effective July 1, 2020.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	100 mg/10 ml	milliliter
Injection	500 mg/50 ml	milliliter

**Rituximab and Hyaluronidase Human ‡ \***

\* Rituximab and Hyaluronidase Human is restricted to use in the treatment of cancer only. Also restricted to NDC labeler code 50242 (Genentech, Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Injection	1400 mg/23400 units/11.7 ml	milliliter
Injection	1600 mg/26800 units/13.4 ml	milliliter

**Rivaroxaban \***

\* Rivaroxaban is restricted to NDC labeler code 50458 (Janssen Pharmaceuticals, Inc.) only.

<b>Dosage Form</b>	<b>Size and/or Strength</b>	<b>Billing Unit</b>
Tablets	10 mg	each
Tablets	15 mg	each
Tablets	20 mg	each
Starter Pack Tablets	15 mg to 20 mg Tablets from 51-tablet pack	each

## Rivastigmine \*

\* Rivastigmine is restricted to treatment of dementia of the Alzheimer's type and to treatment of mild to moderate dementia associated with Parkinson's disease. Use in beneficiaries less than 18 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Transdermal system	4.6 mg/24 hr	each
Transdermal system	9.5 mg/24 hr	each
Transdermal system	13.3 mg/24 hr	each

## Rivastigmine Tartrate \*

\* Rivastigmine Tartrate is restricted to treatment of mild to moderate dementia of the Alzheimer's type and to treatment of mild to moderate dementia associated with Parkinson's disease. Use in beneficiaries less than 18 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Capsules	1.5 mg	each
Capsules	3.0 mg	each
Capsules	4.5 mg	each
Capsules	6.0 mg	each
Solution, oral *	2 mg/ml	milliliter

\* Restricted to claims with dates of service from November 1, 2002, through June 30, 2014, for the oral solution only.

## Rizatriptan \*

\* The following text is effective August 1, 2020: Rizatriptan is restricted to a maximum quantity per dispensing of nine (9) tablets.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg	each
Tablets	10 mg	each
Tablets, orally disintegrating	5 mg	each
Tablets, orally disintegrating	10 mg	each

## Ropinirole HCl \*

\* Use «of Ropinirole HCl» in beneficiaries less than 18 years of age requires treatment authorization approval.

Dosage Form	Size and/or Strength	Billing Unit
Tablets	0.25 mg	each
Tablets	0.5 mg	each
Tablets	1 mg	each
Tablets	2 mg	each
Tablets	3 mg	each
Tablets	4 mg	each
Tablets	5 mg	each
Tablets, extended-release *	2 mg	each
Tablets, extended-release *	4 mg	each
Tablets, extended-release *	6 mg	each
Tablets, extended-release *	8 mg	each
Tablets, extended-release *	12 mg	each

\* «Extended-release tablets are restricted to» NDC labeler code 00007 (GlaxoSmithKline) and to claims with dates of service from January 1, 2009, through June 30, 2012, only. Continuing care with a date of service on or after June 30, 2012, is available when the following conditions are met: 1) The beneficiary had a paid fee-for-service claim for this drug on or before June 30, 2012; 2) A claim had been submitted and paid within the past 100 days; 3) The claim being submitted is within 100 days of the date of service of the last paid claim.

## Rosiglitazone Maleate \*

\* «Rosiglitazone Maleate is» restricted to NDC labeler code 00029 (GalxoSmithKline) and to claims with dates of service through November 18, 2011, only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	2 mg	each
Tablets +	4 mg	each
Tablets +	8 mg	each

## Rosiglitazone Maleate/Glimepiride \*

\* «Rosiglitazone Maleate/Glimepiride is» restricted to NDC labeler code 00007 (GlaxoSmithKline) only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	4 mg/1 mg	each
Tablets +	4 mg/2 mg	each
Tablets +	4 mg/4 mg	each
Tablets +	8 mg/2 mg	each
Tablets +	8 mg/4 mg	each

**Note:** These products are no longer manufactured or available.

## Rosiglitazone Maleate/Metformin HCl \*

\* «Rosiglitazone Maleate/Metformin HCl is» restricted to NDC labeler code 00007 (GlaxoSmithKline) and to claims with dates of service through November 18, 2011, only.

Dosage Form	Size and/or Strength	Billing Unit
Tablets +	1 mg/500 mg	each
Tablets +	2 mg/500 mg	each
Tablets +	4 mg/500 mg	each
Tablets +	2 mg/1000 mg	each
Tablets +	4 mg/1000 mg	each

## Rosuvastatin Calcium

Dosage Form	Size and/or Strength	Billing Unit
Tablets	5 mg	each
Tablets	10 mg	each
Tablets	20 mg	each
Tablets	40 mg	each



## Legend

Symbols used in the document above are explained in the following table.

Symbol	Description
«	This is a change mark symbol. It is used to indicate where on the page the most recent change begins.
»	This is a change mark symbol. It is used to indicate where on the page the most recent change ends.
*	Code I. See paragraph (2) of “General Provisions” in the Drugs: Contract Drugs List Introduction section of this manual regarding authorization and prescription documentation requirements.
+	Frequency of billing requirement. See paragraph (3) of “General Provisions” in the Drugs: Contract Drugs List Introduction section regarding information and exceptions.
∩	Cost is based on this package size. See paragraph (4) of “General Provisions” in the Drugs: Contract Drugs List Introduction section for more information.
§	Authorization not needed for continuing care. See paragraph (6) of “General Provisions” in the Drugs: Contract Drugs List Introduction section for more information.
‡	Drug is exempt from the monthly drug claim line limit. See paragraph (7) of “General Provisions” in the Drugs: Contract Drugs List Introduction section for more information.
«◆»	Suspended until further notice»