### Physician Administered Drug Program Catalog

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| medicaid/medi       | icaid-ncci-ed | dit-files  |                            |                         |                              |   |   |                                   |             |             |                        |                 |                                 |   |                       |
|---------------------|---------------|--|----------------------------|-------------------------|------------------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category            | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                   | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Immune<br>Globulins | 90291         | Cytomegalovirus immune<br>globulin (CMV-IgIV), human,<br>for intravenous use   | 50 mL                      | 1/1/2000                | Cytogam®                     | cytomegalovirus immune<br>globulin intravenous, human   | Indicated for the prophylaxis of cytomegalovirus disease associated with transplantation of kidney, lung,<br>liver, pancreas, and heart. In transplants of these organs other than kidney from CMV seropositive donors<br>into seronegative recipients, prophylactic CMV-IGIV should be considered in combination with ganciclovir.   | 25.2                              | N/A         | N/A         | N/A                    | Y               | Y                               | 3/2024: Rebating Labeler<br>Required field updated to align<br>with policy that submitted<br>NDCs must come from<br>rebating labelers. Update not<br>due to a change in policy. | 3/28/2024             |
| Immune<br>Globulins | 90371         | Hepatitis B Immune Globulin<br>(HBig), human, for<br>intramuscular use   | 1 mL                       | 1/1/2000                | HyperHEP B* S/D,<br>Nabi-HB* | hepatitis b immune globulin,<br>(human)   | Indicated for treatment of acute exposure to blood containing HBsAg, perinatal exposure of infants born<br>to HBsAg-positive mothers, sexual exposure to HBsAg-positive persons and household exposure to persons<br>with acute HBV infection in the following settings:<br>• Acute Exposure to Blood Containing HBsAg. Following either parenteral exposure (needlestick, bite,<br>sharps), direct mucous membrane contact (acidential slpash), or onal ingestion (pipeting acident),<br>involving HBsAg-positive materials such as blood, plasma, or serum.<br>• Perinatal Exposure of Infants Born to HBsAg-positive Mothers: Infants born to mothers positive for<br>HBsAg with or without HBeAg.<br>• Scual Exposure to HBsAg-positive Persons: Secual partners of HBsAg-positive persons.<br>• Household Exposure to Persons with Acute HBV Infection: Infants less than 12 months old whose mother<br>or primary caregiver is positive for HBsAg. Other household contacts with an identifiable blood exposure   | 18                                | N/A         | N/A         | N/A                    | Y               | ¥                               | 3/2024: Rebating Labeler<br>Required field updated to align<br>with policy that submitted<br>NDCs must come from<br>rebating labelers. Update not<br>due to a change in policy. | 3/28/2024             |
| Immune<br>Globulins | 90375         | Rabies Immune Globulin (Rig),<br>human, for intramuscular<br>and/or subcutaneous use   | 150 IU                     | 1/1/2000                | HyperRAB* S/D,<br>HyperRAB*  | rables immune globulin,<br>(human) treated with<br>solvent/detergent, for<br>infiltration and intramuscular<br>administration<br>rables immune globulin,<br>(human) solution for<br>infiltration and intramuscular<br>injection | HyperRAB S(7): Bables vaccine and HyperRAB S(7) should be given to all persons suspected of exposure to<br>rables with one exception; persons who have been previously immunized with rables vaccine and have a<br>confirmed adequate rables antibody titer should receive only vaccine. HyperRAB S(7) should be<br>administered as promptly as possible after exposure, but can be administered up to the eighth day after<br>the first dose of vaccine is given.<br>HyperRAB: Indicated for post exposure prophylaxis, along with rables vaccine, for all persons suspected of<br>exposure to rables.<br>Limitations of use:<br>-Persons previously immunized with rables vaccine that have a confirmed adequate rables antibody titer<br>should receive only vaccine.<br>-Persons previously immunized with rables vaccine that have a confirmed adequate rables antibody titer<br>should receive only vaccine.<br>-Persons previously immunized with rables vaccine that have a confirmed adequate rables antibody titer<br>anonbite exposures regardless of the time interval between exposure and initiation of post-exposure<br>prophylaxis.<br>-Beyond 7 days (after the first vaccine dose), HyperRAB not indicated since an antibody response to<br>vaccine is previoused to have occurred. | 20                                | N/A         | N/A         | N/A                    | Y               | Y                               |   | 4/8/2020              |
| Immune<br>Globulins | 90376         | Rabies Immune Globulin, heat-<br>treated (Rig-HT), human, for<br>intramuscular and/or<br>subcutaneous use                              | 150 IU                     | 1/1/2000                | Imogam® Rabies -<br>HT       | rabies immune globulin<br>(human) USP, heat treated   | Indicated for individuals suspected of exposure to rables, particularly severe exposure, with one exception<br>persons who have been previously immunized with rabies vaccine prepared from human diploid cells<br>(HOCV) in a pre-exposure or post exposure treatment series should receive only vaccine. Persons who<br>have been previously immunized with rabies vaccines other than HOCV, RVA (Rabies Vaccine Adsorbed),<br>or PCEC (Purified Chick Embryo Cell Vaccine) vaccines should have confirmed adequate rabies antibody<br>titers if they are to receive only vaccine.  | 20                                | N/A         | N/A         | N/A                    | Y               | Ŷ                               |   | 9/21/2018             |
| Immune<br>Globulins | 90377         | Rabies immune globulin, heat-<br>and solvent/detergent-treated<br>(RIg-HT S/D), human, for<br>intramuscular and/or<br>subcutaneous use | 150 IU                     | 1/1/2021                | Kedrab™                      | rabies immune globulin<br>(human) solution for<br>intramuscular injection   | Indicated for passive, transient post-exposure prophylaxis of rabies infection to persons of all ages when<br>given immediately after contact with a rabid or possibly rabid animal. Kedrab should be administered<br>concurrently with a full course of rabies vaccine.<br>• Do not exceed the recommended dose of Kedrab because this can partially suppress active production of<br>rabies.<br>• Do not administer additional doses of Kedrab, even if the antibody response to vaccination is delayed.  | 20                                | N/A         | N/A         | N/A                    | Y               | ¥                               |   | 9/21/2022             |
| Vaccines            | 90380         | Respiratory syncytial virus,<br>monoclonal antibody,<br>seasonal dose; 0.5 mL dosage,<br>for intramuscular use                         | 0.5 mL                     | 7/17/2023               | Beyfortus™                   | nirsevimab-alip injection, for<br>intramuscular use (0.5 mL<br>dosage)  | Indicated for the prevention of RSV lower respiratory tract disease in:<br>• Neonates and infants born during or entering their first RSV season.<br>• Children up to 24 months of age who remain vulnerable to severe RSV disease through their second RSV<br>season.  | 1                                 | N/A         | 24 months   | N/A                    | Y               | Ν                               |   | 9/28/2023             |
| Vaccines            | 90381         | Respiratory syncytial virus,<br>monoclonal antibody,<br>seasonal dose; 1 mL dosage,<br>for intramuscular use                           | 1 mL                       | 7/17/2023               | Beyfortus™                   | nirsevimab-alip injection, for<br>intramuscular use (1 mL<br>dosage)  | Indicated for the prevention of RSV lower respiratory tract disease in:<br>• Neonates and infants born during or entering their first RSV season.<br>• Children up to 24 months of age who remain vulnerable to severe RSV disease through their second RSV<br>season.  | 2                                 | N/A         | 24 months   | N/A                    | Ŷ               | Ν                               |   | 9/28/2023             |
| Immune<br>Globulins | 90389         | Tetanus Immune Globulin<br>(TIg), human, for<br>intramuscular use  | 250 units (1 mL)           | 1/1/2000                | HyperTET® S/D                | tetanus immune globulin<br>(human)  | Indicated for prophylaxis against tetanus following injury in patients whose immunization is incomplete or<br>uncertain. It is also indicated, although evidence of effectiveness is limited, in the regimen of treatment or<br>active cases of tetanus.  | 2                                 | N/A         | N/A         | N/A                    | Y               | Y                               |   | 6/4/2019              |
| Immune<br>Globulins | 90396         | Varicella-zoster Immune<br>Globulin (VZIG), human, for<br>intramuscular use (Code Price<br>is per 1 vial = 125 units)                  | 125 units (1 vial)         | 1/1/2000                | Varizig <sup>⊕</sup>         | varicella zoster immune<br>globulin (human) for<br>intramuscular administration<br>only   | Indicated for post exposure prophylaxis in high risk individuals. High risk groups include:<br>immunocompromised children and adults,<br>newborns of mothers with variacella shorthy before or after delivery,<br>premature infants,<br>infants less than one year of age,<br>adults without evidence of immunity,<br>pregnant women.<br>Administration is intended to reduce the severity of varicella.  | 10                                | N/A         | N/A         | N/A                    | Y               | Ŷ                               |   | 7/3/2018              |
| Vaccines            | 90585         | Bacillus Calmette-Guerin<br>Vaccine (BCG) for tuberculosis,<br>live, for percutaneous use.   | 50 mg                      | 1/1/2000                | BCG Vaccine                  | bacillus Calmette-Guérin<br>vaccine (BCG) for<br>tuberculosis, live, for<br>percutaneous use.   | Indicated for the prevention of tuberculosis (TB) in people not previously infected with Mycobacterium<br>tuberculosis, who are at high risk for exposure.  | 1                                 | N/A         | N/A         | N/A                    | Y               | N                               |   | 7/2/2018              |
| Vaccines            | 90589         | Chikungunya virus vaccine, live<br>attenuated, for intramuscular<br>use  | 0.5 mL (1 dose)            | 1/1/2024                | Ixchiq                       | Chikungunya vaccine, live<br>solution for intramuscular<br>injection  | Chikungunya Vaccine is indicated for the prevention of disease caused by chikungunya virus (CHIKV) in<br>individuals 18 years of age and older who are at increased risk of exposure to CHIKV.  | 1                                 | 18 years    | N/A         | N/A                    | Y               | Ν                               |   | 3/22/2024             |

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|---------------|---------------|--|----------------------------|-------------------------|------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Vaccines      | 90611         | Smallpox and monkeypox<br>vaccine, attenuated vaccinia<br>virus, live, non-replicating,<br>preservative free, 0.5 m<br>dosage, suspension, for<br>subcutaneous use | 0.5 mL                     | 7/26/2022               | Jynneos™   | smallpox and monkeypox<br>vaccine, live, non-replicating<br>suspension for subcutaneous<br>and intradermal injection | FDA-Approved Indications:<br>Indicated for prevention of smallpox and monkeypox disease in adults 18 years of age and older<br>determined to be a high risk for smallpox or monkeypox infection.<br>Emergency Use Authorization:<br>The US Food and Drug Administration (FDA) has issued an Emergency Use Authorization (EUA) for the<br>emergency use of Jynneos for:<br>a - active immunization by subcutaneous injection for prevention of monkeypox disease in individuals less<br>than 18 years of age determined to be at high risk for monkeypox infection, and<br>- active immunization by subcutaneous injection for prevention of monkeypox disease in individuals less<br>than 18 years of age determined to be at high risk for monkeypox infection.<br>Justification for Emergency Use of Jynneos During the Monkeypox Public Health Emergency<br>There is currently an outbreak of monkeypox disease. Following a 3-17 day incubation period,<br>individuals infected with monkeypox virus davelop painful lesions that progress sequentially through<br>macular, papular, vesicular, and pustular stages, followed by scabbing over and desiguantion. Lesions<br>may occur anywhere on the body and may be limited to a single site or may be disseminated across many<br>sites. Individuals presentation of monkeypox mises is typically milder than smallpox disease but can be<br>fatal, particularly in severely immunocompromised individuals who do not receive antiviral therapy.<br>During the current monkeypox outbreak, monkeypox cases and exposures have occurred in individuals<br>across a wide range of ages, including infants and children.<br>Jynneos is not approved for these uses. | 2                                 | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | ¥               | Ν                               | Indication Specific Age<br>Restrictions:<br>FOA-Approved Indications:<br>18 years of age and older<br>Emergency Use Authorization:<br>N/A  | 5/31/2024             |
| Vaccines      | 90619         | Meningococcal conjugate<br>vaccine, serogroups A, C, W, Y,<br>quadrivalent, tetanus toxoid<br>carrier (MenACWY-TT), for<br>intramuscular use                       | 0.5 mL                     | 7/1/2019                | MenQuadfi™ | meningococcal [Groups A, C,<br>Y, W] conjugate vaccine,<br>solution for intramuscular<br>injection                   | Indicated for active immunization for the prevention of invasive meningococcal disease caused by<br>Neisseria meningitidis sergroups A, C, W, and Y. MenQuadfi vaccine is approved for use in individuals 2<br>varsar of age and older.<br>MenQuadfi does not prevent N. meningitidis serogroup B disease.  | 1                                 | 2 years   | N/A         | N/A                    | Y               | N                               |  | 8/5/2021              |
| Vaccines      | 90620         | Meningococcal recombinant<br>protein and outer membrane<br>vesicle vaccine, serogroup B<br>(MenB-4C), 2 dose schedule,<br>for intramuscular use                    | 0.5 mL                     | 2/1/2015                | Bexsero®   | meningococcal group b<br>vaccine suspension for<br>intramuscular injection   | Indicated for active immunization to prevent invasive disease caused by Neisseria meningitidis serogroup<br>B. Bexsero is approved for use in individuals 10 through 25 years of age.   | 2                                 | 10 years  | 25 years    | N/A                    | Ŷ               | N                               | 12/2023: Maximum age<br>restriction updated to align<br>with FDA-approved and ACIP-<br>recommended maximum age<br>effective 10/1/2023.   | 1/26/2024             |
| Vaccines      | 90621         | Meningococcal recombinant<br>lipoprotein vaccine, serogroup<br>B (MenB-FHbp), 2 or 3 dose<br>schedule, for intramuscular<br>use                                    | 0.5 mL                     | 2/1/2015                | Trumenba®  | meningococcal group b<br>vaccine suspension for<br>intramuscular injection   | Indicated for active immunization to prevent invasive disease caused by Neisseria meningitidis serogroup<br>B. Trumenba is approved for use in individuals 10 through 25 years of age.  | 2                                 | 10 years  | 25 years    | N/A                    | Y               | N                               | 12/2023: Maximum age<br>restriction updated to align<br>with FDA-approved and ACIP-<br>recommended maximum age<br>effective 10/1/2023.   | 1/26/2024             |
| Vaccines      | 90623         | Meningococcal pentavalent<br>vaccine, conjugated Men A, C,<br>W, Y- tetanus toxoid carrier,<br>and Men B-FHbp, for<br>intramuscular use                            | 0.5 mL                     | 1/1/2024                | Penbraya™  | meningococcal groups A, B,<br>C, W, and Y vaccine,<br>suspension for intramuscular<br>injection                      | Meningococcal groups A, B, C, W, and Y vaccine, suspension for intramuscular injection is indicated for<br>active immunization to prevent invasive disease caused by <i>Neisseria meningitidis</i> serogroups A, B, C, W,<br>and Y. Penbraya is approved for use in individuals 10 through 25 years of age.   | 1                                 | 10 years  | 25 years    | N/A                    | Y               | N                               | 7/2024: Addition to VFC<br>effective 7/18/2024 per DHB<br>request 7/23/2024.   | 7/29/2024             |
| Vaccines      | 90625         | Cholera vaccine, live, adult<br>dosage, 1 dose schedule, for<br>oral use   | 1 adult dosage (100<br>mL) | 1/1/2016                | Vaxchora*  | cholera vaccine, live, oral<br>suspension for oral<br>administration   | Indicated for active immunization against disease caused by Vibrio cholerae serogroup O1. Vaxchora is<br>approved for use in persons 2 through 64 years of age traveling to cholera-affected areas.<br>Limitations of Use:<br>• The effectiveness of Vaxchora has not been established in persons living in cholera-affected areas.<br>• The effectiveness of Vaxchora has not been established in persons who have pre-existing immunity due<br>to previous exposure to V. cholerae or receipt of a cholera vaccine.<br>• Vaxchora has not been shown to protect against disease caused by V. cholerae serogroup O139 or other<br>non-O1 serogroups.   | 1                                 | 2 years   | 64 years    | N/A                    | Y               | N                               |  | 10/27/2023            |
| Vaccines      | 90626         | Tick-borne encephalitis virus<br>vaccine, inactivated; 0.25 mL<br>dosage, for intramuscular use  | 0.25 mL                    | 7/1/2021                | TicoVac™   | tick-borne encephalitis<br>vacine, suspension for<br>intramuscular injection (0.25<br>mL dose)                       | Tick-borne encephalitis vaccine is indicated for active immunization to prevent tick-borne encephalitis<br>(TBE). It is approved for use in individuals 1 year of age and older.  | 1                                 | 1 year  | 15 years    | N/A                    | Y               | N                               | 1/2024: Coverage effective<br>11/10/2023<br>6/2024: Rebating Labeler<br>Required field updated to align<br>with policy that submitted<br>warcine NDCs do not need to<br>come from rebating labelers.<br>Update not due to a change in<br>policy. | 6/7/2024              |

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|--------------|---------------|---|----------------------------|-------------------------|-----------------|---|--|---|-------------|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
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| Vaccines     | 90627         | Tick-borne encephalitis virus<br>vaccine, inactivated; 0.5 mL<br>dosage, for intramuscular use  | 0.5 mL                     | 7/1/2021                | TicoVac™        | tick-borne encephalitis<br>vaccine, suspension for<br>intramuscular injection (0.5<br>mL dose)                | Tick-borne encephalitis vaccine is indicated for active immunization to prevent tick-borne encephalitis<br>(TBE). It is approved for use in individuals 1 year of age and older.   | 2   | 16 years    | N/A         | N/A                    | Y               | N                               | 1/2024: Coverage effective<br>11/10/2023<br>6/2024: Rebating Labeler<br>Required field updated to align<br>with policy that submitted<br>vaccine NDCs do not need to<br>come from rebating labelers.<br>Update not due to a change in<br>policy. | 6/7/2024              |
| Vaccines     | 90632         | Hepatitis A vaccine (Hep A),<br>adult dosage, for<br>intramuscular use  | 1 mL                       | 1/1/2000                | Havrix®, Vaqta® | hepatitis a vaccine, adult<br>dosage, suspension for<br>intramuscular injection                               | Indicated for active immunization against disease caused by hepatitis A virus (HAV). Approved for use in<br>persons 12 months of age and older. Primary immunization should be administered at least 2 weeks prior<br>to expected exposure to HAV.   | 1   | 19 years    | N/A         | N/A                    | Y               | N                               |  | 7/3/2018              |
| Vaccines     | 90633         | Hepatitis A vaccine (Hep A),<br>pediatric/adolescent dosage -<br>2-dose schedule, for<br>intramuscular use  | 0.5 mL                     | 1/1/2000                | Havrix®, Vaqta® | hepatitis a vaccine,<br>pediatric/adolescent dosage-<br>2 dose schedule, for<br>intramuscular injection       | Indicated for active immunization against disease caused by hepatitis A virus (HAV). Approved for use in<br>persons 12 months of age and older. Primary immunization should be administered at least 2 weeks prior<br>to expected exposure to HAV.   | 1   | 12 months   | 18 years    | N/A                    | Y               | N                               |  | 7/3/2018              |
| Vaccines     | 90636         | Hepatitis A and Hepatitis B<br>Vaccine (HepA-HepB), adult<br>dosage, for intramuscular use  | 1 mL                       | 1/1/2000                | Twinrix®        | hepatitis a & hepatitis b<br>(recombinant) vaccine<br>suspension for intramuscular<br>injection               | Indicated for active immunization against disease caused by hepatitis A virus and infection by all known<br>subtypes of hepatitis B virus. Twinrix is approved for use in persons 18 years of age or older.  | з   | 18 years    | N/A         | N/A                    | Y               | N                               |  | 9/12/2018             |
| Vaccines     | 90647         | Haemophilus influenzae type b<br>vaccine (Hib), PRP-OMP<br>conjugate, 3-dose schedule,<br>for intramuscular use                                   | 0.5 mL                     | 1/1/2000                | PedvaxHib*      | haemophilus b conjugate<br>vaccine (meningococcal<br>protein conjugate)                                       | For routine vaccination against invasive disease caused by haemophilus influenzae type B in infants and<br>children 2 – 71 months of age.  | 1   | 2 months    | 71 months   | N/A                    | Y               | N                               |  | 7/2/2018              |
| Vaccines     | 90648         | Haemophilus influenzae b<br>vaccine (Hib), PRP-T<br>conjugate, 4-dose schedule,<br>for intramuscular use  | 0.5 mL                     | 1/1/2000                | ActHIB®         | haemophilus b conjugate<br>vaccine (tetanus toxoid<br>conjugate) solution for<br>intramuscular injection      | Indicated for the prevention of invasive disease caused by Haemophilus influenzae type b. ActHB vaccine<br>is approved for use as a four dose series in infants and children 2 months through 5 years of age.  | 1   | 2 months    | 5 years     | N/A                    | Y               | N                               |  | 7/3/2018              |
| Vaccines     | 90651         | Human Papillomavirus vaccine<br>types 6, 11, 16, 18, 31, 33, 45,<br>52, 58, nonavalent (9vHPV), 2<br>or 3 dose schedule, for<br>intramuscular use | 0.5 mL                     | 7/1/2017                | Gardasil* 9     | human papillomavirus 9-<br>valent vaccine, recombinant<br>suspension for intramuscular<br>injection           | Indicated in girs and women 9 through 45 years of age for the prevention of the following diseases:<br>• Cervical, vulvar, vaginal, and anal cancer caused by HPV types 16, 18, 31, 33, 45, 52, and 58<br>• Gential warts (condytoma acuminata) caused by HPV types 6, 11, 16, 18, 31, 33, 45, 52, and 58<br>• Gential warts (condytoma acuminata) caused by HPV types 6, 11, 16, 18, 31, 33, 45, 52, and 58:<br>• Cervical intracpithelian neoplasis (CNI) grade 2/3 and cervical adenocarcinoma in situ (AIS).<br>• Cervical intracpithelian neoplasis (CNI) grade 2 and grade 3.<br>• Vulvar intracpithelian neoplasis (CNI) grade 2 and grade 3.<br>• Vaginal intracpithelian neoplasis (CNI) grade 2 and grade 3.<br>• Anal intracpithelian neoplasis (CNI) grades 1, 2, and 3.<br>indicated in boys and men 9 through 45 years of age for the prevention of the following diseases:<br>• Anal cancer caused by HPV types 16, 18, 31, 33, 45, 52, and 58.<br>• Centual warts (condytoma acuminata) caused by HPV types 6, 11, 16, 18, 31, 33, 45, 52, and 58.<br>• Anal intracepithelian neoplasis (LNI) grades 1, 2, and 3.<br>• Anal intracepithelian neoplasis (LNI) grades 1, 2, and 3.<br>• Anal intracepithelian neoplasis (LNI) grades 1, 2, and 3.<br>• Anal intracepithelian neoplasis (LNI) grades 1, 2, and 3.<br>• Anal intracepithelian neoplasis (LNI) grades 1, 2, and 3.<br>• Indicated in girs and women 9 through 45 years of age for the prevention of oropharyngeal and other head and neck cancers caused by HPV types 16, 18, 31, 33, 45, 52, and 58. | 2   | 9 years     | 45 years    | N/A                    | ¥               | N                               |  | 6/25/2024             |
| Vaccines     | 90653         | Influenza vaccine, inactivated<br>(IIV), subunit, adjuvanted, for<br>intramuscular use  | 0.5 mL                     | 1/1/2013                | Fluad®          | influenza vaccine, inactivated<br>(IIV), subunit, adjuvanted, for<br>Intramuscular use, 2024-<br>2025 Formula | Indicated for active immunization for the prevention of influenza disease caused by influenza A subtype<br>viruses and type B virus contained in the vaccine for use in persons 65 years of age and older.   | 1   | 65 years    | N/A         | N/A                    | ¥               | N                               |  | 7/29/2024             |

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| nedicaid/medi | caid-ncci-ed  | lit-files   |                            | 1                       | 1   | 1   |  |                                   |              | 1           |                        |                 | Rebating            |   |                       |
|---------------|---------------|---|----------------------------|-------------------------|---|---|--|-----------------------------------|--------------|-------------|------------------------|-----------------|---------------------|---|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name  | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age  | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Labeler<br>Required | Comments                                      | Last Modified<br>Date |
| Vaccines      | 90656         | Influenza virus vaccine,<br>trivalent (IIV3), split virus,<br>preservative free, 0.5 mL<br>dosage, for intramuscular use  | 0.5 mL                     | 1/1/2005                | Afluria®, Fluarix,<br>FluLaval, Fluzone®  | influenza virus vaccine,<br>trivalent (IIV3), split virus,<br>preservative free, 0.5 mL<br>dosage, for intramuscular<br>use, 2024-2025 Formula                            | Indicated for active immunization for the prevention of disease caused by influenza A subtype viruses and<br>type B virus contained in the vaccine.<br>Afluria (0.5 mL): Approved for use in persons 36 months of age and older.<br>Fluarix: Approved for use in persons 6 months of age and older.<br>Fluare: Approved for use in persons 6 months of age and older.  | 2                                 | 6 months     | N/A         | N/A                    | Y               | N                   |   | 7/29/2024             |
| Vaccines      | 90657         | Influenza virus vaccine,<br>trivalent (IIV3), split virus,<br>0.25 mL dosage, for<br>intramuscular use  | 0.25 mL                    | 1/1/1999                | Afluria®, Fluzone®  | Influenza Virus Vaccine, split<br>virus, for IM or jet injection<br>use, contains preservatives<br>(0.25 mL dosage), 2024-2025<br>Formula                                 | Vaccination against influenza types A and B in children 6-35 months of age.  | 2                                 | 6 months     | 35 months   | N/A                    | Y               | N                   |   | 7/29/2024             |
| Vaccines      | 90658         | Influenza virus vaccine,<br>trivalent (IIV3), split virus, 0.5<br>mL dosage, for intramuscular<br>use   | 0.5 mL                     | 1/1/1999                | Afluria®, Fluzone®  | influenza virus vaccine,<br>trivalent (IIV3), split virus, 0.5<br>mL dosage, for intramuscular<br>use, 2024-2025 Formula  | Indicated for active immunization for the prevention of disease caused by influenza A subtype viruses and<br>type 8 virus contained in the vaccine.<br>Afluria (0.5 mL dose): Approved for use in persons 36 months of age and older.<br>Flucture: Approved for use in persons 6 months of age and older.  | 2                                 | 6 months     | N/A         | N/A                    | Y               | Ν                   |   | 7/29/2024             |
| Vaccines      | 90661         | Influenza virus vaccine<br>(ccIIV3), derived from cell<br>cultures, subunit, antibiotic<br>free, for intramuscular use  | 0.5 mL                     | 1/1/2008                | Flucelvax®  | Influenza virus vaccine,<br>trivalent (ccIIV3), derived<br>from cell cultures, subunit,<br>antibiotic free, 0.5 mL<br>dosage, for intramuscular<br>use, 2024-2025 Formula | Influenza Vaccine Injectable Suspension is indicated for active immunization for the prevention of<br>influenza disease caused by influenza virus subtypes A and type B contained in the vaccine. Flucelvax is<br>approved for use in persons 6 months of age and older.   | 2                                 | 6 months     | N/A         | N/A                    | Y               | N                   |   | 7/29/2024             |
| Vaccines      | 90662         | Influenza virus vaccine (IIV),<br>split virus, preservative free,<br>enhanced immunogenicity via<br>increased antigen content, for<br>intramuscular use                             | 0.5 mL                     | 1/1/2008                | Fluzone® High-<br>Dose  |   | Indicated for active immunization for the prevention of influenza disease caused by influenza A subtype<br>viruses and type B virus contained in the vaccine for use in persons 65 years of age and older.   | 1                                 | 65 years     | N/A         | N/A                    | Y               | N                   |   | 7/29/2024             |
| Vaccines      | 90670         | Pneumococcal conjugate<br>vaccine, 13 valent (PCV13), for<br>intramuscular use  | 0.5 mL                     | 7/1/2009                | Prevnar 13*   | pneumococcal 13-valent<br>conjugate vaccine (diphtheria<br>CRM197 protein) suspension<br>for intramuscular injection  | In children 6 weeks through 5 years of age (prior to the 6th birthday), Prevnar 13 is indicated for:<br>A citive immunization for the prevention of invasive disease caused by Streptococcus pneumoniae<br>serotypes 1, 3, 4, 5, 6A, 6B, 7F, 9V, 14, 18C, 19A, 19F and 23F.<br>-active immunization for the prevention of dits media caused by 5, pneumoniae serotypes 4, 6B, 9V, 14,<br>18C, 19F, and 23F. No otitis media efficacy data are available for serotypes 1, 3, 5, 6A, 7F, and 19A.<br>In children 6 years through 17 years of age (prior to the 18th birthday), Prevnar 13 is indicated for:<br>* Active immunization for the prevention of invasive disease caused by S. pneumoniae serotypes 1, 3, 4, 5,<br>6A, 6B, 7F, 9V, 14, 18C, 19A, 19F and 23F.<br>In adults 18 years of age and older, Prevnar 13 is indicated for:<br>* Active immunization for the prevention of pneumonia and invasive disease caused by S. pneumoniae<br>serotypes 1, 3, 4, 5, 6A, 6B, 7F, 9V, 14, 18C, 19A, 19F and 23F. | 1                                 | 6 weeks      | N/A         | N/A                    | ¥               | N                   |   | 7/3/2018              |
| Vaccines      | 90671         | Pneumococcal conjugate<br>vaccine, 15 valent (PCV15), for<br>intramuscular use  | 0.5 mL (1 dose)            | 7/1/2021                | Vaxneuvance™  | pneumococcal 15-valent<br>conjugate vaccine suspension<br>for intramuscular injection   | Indicated for active immunization for the prevention of invasive disease caused by Streptococcus<br>pneumonize serotypes 1, 3, 4, 5, 6A, 6B, 7F, 9V, 14, 18C, 19A, 19F, 22F, 23F, and 33F in individuals 6<br>weeks of age and older.<br>ACIP recommends use of PCV15 as an option for pneumococcal conjugate vaccination of persons aged <19<br>years, according to currently recommended PCV13 dosing and schedules.   | 1                                 | 6 weeks      | N/A         | N/A                    | Y               | N                   | ACIP recommends for 6 weeks of age and older  | 10/20/2022            |
| Vaccines      | 90673         | Influenza virus vaccine,<br>trivalent (RIV3), derived from<br>recombinant DNA,<br>hemagglutinin (HA) protein<br>only, preservative and<br>antibiotic free, for<br>intramuscular use | 0.5 mL                     | 1/1/2014                | Flublok®  | influenza virus vaccine,<br>trivalent (RIV3), derived from<br>recombinant DNA,<br>preservative and antibiotic<br>free, for intramuscular use,<br>2024-2025 Formula        | Indicated for active immunization against disease caused by influenza A virus subtypes and influenza type<br>B virus contained in the vaccine. Flublok is approved for use in persons 18 years of age and older.   | 1                                 | 18 years     | N/A         | N/A                    | Y               | N                   |   | 7/29/2024             |
| Vaccines      | 90675         | Rabies vaccine, for<br>intramuscular use  | 1 mL                       | 1/1/2000                | Imovax® Rabies<br>(Human Diploid-<br>Cell Vaccine) and<br>RabAvert®<br>(Purified Chick<br>Embryo Cell<br>Culture) | rabies vaccine, for<br>intramuscular use  | Indicated for pre-exposure and post-exposure prophylaxis against rables in all age groups.   | 5                                 | N/A          | N/A         | N/A                    | Y               | N                   |   | 7/3/2018              |
| Vaccines      | 90677         | Pneumococcal conjugate<br>vaccine, 20 valent (PCV20), for<br>intramuscular use  | 0.5 mL                     | 7/1/2021                | Prevnar 20™   | pneumococcal 20-valent<br>conjugate vaccine,<br>suspension for intramuscular<br>injection   | Prevnar 20 is a vaccine indicated for active immunization for the prevention of:<br>• pneumonia caused by S. <i>pneumonice</i> serotypes 1, 3, 4, 5, 6A, 6B, 7F, 8, 9V, 10A, 11A, 12F, 14, 15B, 18C,<br>19A, 19F, 22F, 23F, and 33F in individuals 18 years of age and older.<br>• invasive disease caused by Streptococcus pneumonine serotypes 1, 3, 4, 5, 6A, 6B, 7F, 8, 9V, 10A, 11A,<br>12F, 14, 15B, 18C, 13A, 19F, 22F, 23F, and 33F in individuals 6 weeks of age and older.<br>• otitis media caused by S. pneumoniae serotypes 4, 6B, 9V, 14, 18C, 19F, and 23F in individuals 6 weeks<br>through 5 years of age.  | 1                                 | See Comments | N/A         | N/A                    | Y               | N                   | ACIP recommends for 2 months of age and older | 9/28/2023             |

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|--------------|---------------|--|----------------------------|-------------------------|-------------------------------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|----------------------|
| Vaccines     | 90678         | Respiratory syncytial virus<br>vaccine, pref, subunit,<br>bivalent, for intramuscular use  | 0.5 mL                     | 1/1/2023                | Abrysvo**                           | respiratory syncytial virus<br>vaccine solution for<br>intramuscular injection                  | Indicated for:<br>- active immunization for the prevention of lower respiratory tract disease (LRTD) caused by respiratory<br>syncytial virus (RSV) in individuals 60 years of age and older.<br>- active immunization of pregnant individuals at 32 through 36 weeks gestational age for the prevention of<br>lower respiratory tract disease (LRTD) and severe LRTD caused by respiratory syncytial virus (RSV) in<br>infamts from birth through 6 months of age.  | , 1                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | N                               | Indication specific age<br>restrictions:<br>• Active immunization for the<br>prevention of RID caused by<br>RSV: 60 years of age and older<br>• Active immunization of<br>pregnant individuals at 32<br>through 36 weeks gestational<br>age for the prevention of RITD<br>and severe LRTD caused by<br>RSV in infants from birth<br>through 6 months of age: use<br>after menarche<br>1/2024: Addition to VFC<br>Effective 1/2/2024 per<br>DHB Request 12/21/2023 | 1/26/2024            |
| Vaccines     | 90679         | Respiratory syncytial virus<br>vaccine, preF, recombinant,<br>subunit, adjuvanted, for<br>intramuscular use  | 0.5 mL                     | 5/3/2023                | Arexvy                              | respiratory syncytial virus<br>vaccine, adjuvanted<br>suspension for intramuscular<br>injection | Indicated for active immunization for the prevention of lower respiratory tract disease (LRTD) caused by<br>respiratory syncytial virus in:<br>- individual 50 queras of age and older;<br>- individual 50 through 59 years of age who are at increased risk for LRTD caused by RSV.   | 1                                 | 60 years  | N/A         | N/A                    | Y               | N                               |   | 7/29/2024            |
| Vaccines     | 90680         | Rotavirus vaccine, pentavalent<br>(RV5), 3 dose schedule, live,<br>for oral use  | t 2 mL                     | 7/1/2005                | RotaTeq®                            |   | Indicated for the prevention of rotavirus gastroenteritis in infants and children caused by types G1, G2,<br>G3, G4, and G9 when administered as a 3-dose series to infants between the ages of 6 to 32 weeks.   | 2                                 | 6 weeks   | 8 months    | N/A                    | Y               | N                               | ACIP recommends for 6 weeks<br>of age to 8 months of age  | 3/30/2023            |
| Vaccines     | 90681         | Rotavirus vaccine, human,<br>attenuated (RV1), 2 dose<br>schedule, live, for oral use  | 1 mL                       | 1/1/2008                | Rotarix                             | rotavirus vaccine, live, oral   | Indicated for the prevention of rotavirus gastroenteritis caused by G1 and non-G1 types (G3, G4, and G9).<br>Rotarix is approved for use in infants 6 weeks to 24 weeks of age.  | 2                                 | 6 weeks   | 8 months    | N/A                    | Y               | N                               | ACIP recommends for 6 weeks<br>of age to 8 months of age  | 3/30/2023            |
| Vaccines     | 90683         | Respiratory syncytial virus<br>vaccine, mRNA lipid<br>nanoparticles, for<br>intramuscular use  | 0.5 mL                     | 1/1/2024                | mRESVIA™                            | respiratory syncytial virus<br>vaccine injectable<br>suspension, for intramuscular<br>use       | Respiratory Syncytial Virus Vaccine is a vaccine indicated for active immunization for the prevention of<br>lower respiratory tract disease (LRTD) caused by respiratory syncytial virus (RSV) in individuals 60 years of<br>age and older.  | 1                                 | 60 years  | N/A         | N/A                    | Y               | Ν                               |   | 7/29/2024            |
| Vaccines     | 90690         | Typhoid vaccine, live, oral  | 4 capsules                 | 1/1/2000                | Vivotif®                            | typhoid vaccine live oral<br>Ty21a  | Indicated for immunization of adults and children greater than 6 years of age against disease caused by<br>Salmonella typhi. Routine typhoid vaccination is not recommended in the United States of America.<br>Selective immunization against typhoid fever is recommended for the following groups:<br>1) travelers to areas in which there is a recognized risk of exposure to 5. typhi,<br>2) persons with intimate exposure (e.g. household contact) to a 5. typhi carrier, and<br>3) microbiology laboratorians who work frequently with 5. typhi.<br>There is no evidence to support the use of typhoid vaccine to control common source outbreaks, disease<br>following natural disasters or in persons attending rural summer camps.  | 1                                 | 6 years   | N/A         | N/A                    | ¥               | N                               |   | 10/27/2023           |
| Vaccines     | 90691         | Typhoid vaccine, Vi capsular<br>polysaccharide (ViCPs), for<br>intramuscular use   | 0.5 mL                     | 1/1/2000                | Typhim Vi®                          | typhoid vi polysaccharide<br>vaccine  | Indicated for active immunization for the prevention of typhoid fever caused by S typhi and is approved<br>for use in persons two years of age or older, immunization with Typhim VI vaccine should occur at least<br>two weeks piror to expected exposure to S typhi.<br>Typhim VI vaccine is not indicated for routine immunization of individuals in the United States (US).<br>Selective immunization against typhoid fever is recommended under the following circumstances:<br>1) travelers to areas where a recognized risk of exposure to typhoid exists, particularly ones who will have<br>prolonged exposure to potentially contaminated food and water,<br>2) persons with intimate exposure (ec, continued household contact) to a documented typhoid carrier, and<br>3) workers in microbiology laboratories who frequently work with S typhi.<br>An optimal reimmunization schedule has not been established. Reimmunization every two years under<br>conditions of repeated or continued exposure to the S typhi organism is recommended at this time. |                                   | 2 years   | N/A         | N/A                    | Y               | N                               |   | 10/27/2023           |
| Vaccines     | 90696         | Diphtheria, tetanus toxoids,<br>acellular pertussis vaccine and<br>inactivated poliovirus vaccine,<br>(DTa-PI-V), when<br>administered to children 4<br>years through 6 years of age,<br>for intramuscular use |                            | 1/1/2008                | Kinrix <sup>®</sup> ,<br>Quadracel™ |   | <ul> <li>Kinric: A single dose of Kinrix is indicated for active immunization against diphtheria, tetanus, pertussis, and poliomyelitis as the fifth dose in the diphtheria, tetanus, and acellular pertussis (DTaP) vaccine oseries and the fourth dose in the inactivated poliovirus vaccine (IPV) series in children 4 through 6 years of age whose previous DTaP vaccine doses have been with INFANRIX and/or PEDIARIX for the first three doses and INFANRIX and/or PEDIARIX for the first three doses and INFANRIX for the fourth dose.</li> <li>Quadracel: indicated for active immunization against diphtheria, tetanus, pertussis and poliomyelitis. A single dose of Quadracel is approved for use in children four through ix years of age as a fifth dose in the diphtheria, tetanus, pertussis vaccination (IPTaP) series, in children who have received four doses of Pentacel and/or Daptacel vaccine.</li> </ul>   | 1                                 | 4 years   | б years     | N/A                    | Y               | N                               |   | 7/2/2018             |

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| edicaid/medi | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                                     | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age  | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modifie<br>Date |
|--------------|---------------|--|----------------------------|-------------------------|--|---|---|-----------------------------------|---|--|------------------------|-----------------|---------------------------------|--|----------------------|
| Vaccines     | 90697         | Diphtheria, tetanus toxoids,<br>aellular pertussis vaccine,<br>inactivated poliovirus<br>vaccine, Haemophilus<br>influenzae type b PR-OMP<br>conjugate vaccine, and<br>hepatitis B vaccine (DTAP-IPV-<br>Hib-HepB), for intramuscular<br>use | 0.5 mL                     | 1/1/2015                | Vaxelis™                                       | diphtheria and tetanus<br>toxoids and acellular<br>pertussis, inactivated<br>poliovirus, haemophilus b<br>conjugate and hepatitis B<br>vaccine suspension for<br>intramuscular injection                            | Indicated for active immunization to prevent diphtheria, tetanus, pertussis, poliomyelitis, hepatitis B, and<br>invasive disease due to Haemophilus influenzae type b. Vaxelis is approved for use as a 3-dose series in<br>children from 6 weeks through 4 years of age (prior to the 5th birthday).   | 1                                 | 6 weeks   | 4 years  | N/A                    | Y               | N                               |  | 12/20/2022           |
| Vaccines     | 90698         | Diphtheria, tetanus toxoids,<br>acellular pertussis vaccine,<br>Haemophilus influenzae type<br>b, and inactivated poliovirus<br>vaccine, (DTaP-IPV / Hib), for<br>intramuscular use  | 0.5 mL                     | 1/1/2004                | Pentacel®                                      | diphtheria and tetanus<br>toxoids and acellular<br>pertussis adsorbed,<br>inactivated poliovirus and<br>haemophilus b conjugate<br>(tetanus toxoid conjugate)<br>vaccine, suspension for<br>intramuscular injection | Indicated for active immunization against diphtheria, tetanus, pertussis, poliomyelitis, and invasive<br>disease due to Haemophilus influenzae type b. Pentacel vaccine is approved for use as a four dose series in<br>children 6 weeks through 4 years of age (prior to fifth birthday).  | 1                                 | 6 weeks   | 4 years  | N/A                    | Y               | N                               |  | 7/2/2018             |
| Vaccines     | 90700         | Diphtheria, tetanus toxoids,<br>and acellular pertussis vaccine<br>(DTaP), when administered to<br>individuals younger than seven<br>years, for intramuscular use  | 0.5 mL                     | 1/1/2004                | Daptacel®,<br>Infanrix®                        | diphtheria, tetanus toxoids,<br>and acellular pertussis<br>vaccine adsorbed suspension<br>for intramuscular injection   | Indicated for active immunization against diphtheria, tetanus and pertussis as a five dose series in infants<br>and children 6 weeks through 6 years of age (prior to 7th birthday).  | 1                                 | 6 weeks   | 6 years  | N/A                    | Y               | Ν                               |  | 7/2/2018             |
| Vaccines     | 90702         | Diphtheria and tetanus toxoids<br>adsorbed (DT) when<br>administered to individuals<br>younger than 7 years, for<br>intramuscular use.   | 0.5 mL                     | 1/1/2000                | Diphtheria and<br>Tetanus Toxoids,<br>Adsorbed | diphtheria and tetanus<br>toxoids (DT), adsorbed, for<br>use in individuals younger<br>than seven years, for<br>intramuscular use.  | Indicated for active immunization against diphtheria and tetanus. Diphtheria and Tetanus Toxoids<br>Adsorbed is approved for use in children from 6 weeks through 6 years of age (prior to 7th birthday).   | 1                                 | 6 weeks   | 6 years  | N/A                    | Y               | N                               |  | 7/2/2018             |
| Vaccines     | 90707         | Measles, mumps and rubella<br>virus vaccine (MMR), live, for<br>subcutaneous use   | 0.5 mL                     | 1/1/2000                | M-M-R® II                                      | measles, mumps, and rubella<br>virus vaccine live suspension<br>for intramuscular or<br>subcutaneous injection  | Indicated for simultaneous vaccination against measles, mumps, and rubella in individuals 12 months of<br>age or older.   | 1                                 | 12 months   | N/A  | N/A                    | Y               | N                               | 10/2023: HCPCS Effective Date<br>updated from 1/1/2004 to<br>1/1/2000.                 | 10/27/2023           |
| Vaccines     | 90707         | Measles, mumps and rubella<br>virus vaccine (MMR), live, for<br>subcutaneous use   | 0.5 mL                     | 1/1/2000                | Priorix  | measles, mumps, and rubella<br>vaccine, live, suspension for<br>subcutaneous injection  | Indicated for active immunization for the prevention of measles, mumps, and rubella in individuals 12<br>months of age and older.   | 2                                 | 12 months   | N/A  | N/A                    | Y               | N                               |  | 8/16/2022            |
| Vaccines     | 90710         | Measles, mumps, rubella, and<br>varicella vaccine (MMRV), live,<br>for subcutaneous use  | 0.5 mL                     | 1/1/2000                | ProQuad®                                       | measles, mumps, rubella and<br>varicella virus vaccine live<br>suspension for intramuscular<br>or subcutaneous injection  | Indicated for active immunization for the prevention of measles, mumps, rubella, and varicella in children 12 months through 12 years of age.   | 1                                 | 12 months   | 12 years   | N/A                    | Y               | Ν                               |  | 3/16/2023            |
| Vaccines     | 90713         | Poliovirus vaccine, Inactivated<br>(IPV), for subcutaneous or<br>intramuscular use   | 0.5 mL                     | 7/1/2005                | IPOL*  | poliovirus vaccine,<br>inactivated  | Indicated for active immunization of infants (as young as 6 weeks of age), children and adults for the<br>prevention of poliomyelitis caused by poliovirus types 1, 2, and 3.   | 2                                 | 6 weeks   | N/A  | N/A                    | Y               | Ν                               |  | 9/21/2018            |
| Vaccines     | 90714         | Tetanus and diphtheria<br>toxoids adsorbed (†d),<br>preservative free, when<br>administered to individuals 7<br>years or older, for<br>intramuscular use   | 0.5 mL                     | 7/1/2005                | Tenivac®                                       | tetanus and diphtheria<br>toxoids, adsorbed,<br>suspension for intramuscular<br>injection   | Indicated for active immunization for the prevention of tetanus and diphtheria in persons 7 years of age<br>and older.  | 2                                 | 7 years   | N/A  | N/A                    | Y               | N                               |  | 7/3/2018             |
| Vaccines     | 90715         | Tetanus, diphtheria toxoids<br>and acellular pertussis vaccine<br>(Tdap), when administered to<br>individuals 7 years or older, for<br>intramuscular use   | 0.5 mL                     | 7/1/2005                | Adacel®,<br>Boostrix®                          | tetanus toxoid, reduced<br>diphtheria toxoid and<br>acellular pertussis vaccine<br>adsorbed, suspension for<br>intramuscular injection  | Adacet:<br>Indicated for:<br>• active booster immunization against tetanus, diphtheria and pertussis. Adacel is approved for use in<br>persons 10 through 64 years of age.<br>• immunization during the third trimester of pregnancy to prevent pertussis in infants younger than 2<br>months of age.<br>Boostrix:<br>Indicated for:<br>• active booster immunization against tetanus, diphtheria, and pertussis in individuals aged 10 years and | 1                                 | Min age restriction<br>updated at the<br>request of the State:<br>7 years | Product Specific<br>Age Restrictions<br>(see comments) | N/A                    | Y               | N                               | Product specific maximum age<br>restrictions:<br>• Adacel: 64 years<br>• Boostrix: N/A | 2/23/2023            |
| Vaccines     | 90716         | Varicella virus vaccine (VAR),<br>Live, for subcutaneous use   | 0.5 mL                     | 1/1/2000                | Varivax®                                       | varicella virus vaccine live<br>suspension for intramuscular<br>or subcutaneous injection   | Indicated for active immunization for the prevention of varicella in individuals 12 months of age and older.  | 2                                 | 12 months   | N/A  | N/A                    | Y               | N                               |  | 3/16/2023            |

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| edicaid/medic | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name  | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age  | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modifie<br>Date |
|---------------|---------------|---|----------------------------|-------------------------|---|---|---|-----------------------------------|--|-------------|------------------------|-----------------|---------------------------------|---|----------------------|
| Vaccines      | 90717         | Yellow fever vaccine, live, for subcutaneous use  | 0.5 mL                     | 1/1/2000                | YF-V∂x®   | yellow fever vaccine, for<br>subcutaneous use   | Indicated for active immunization for the prevention of yellow fever in persons 9 months of age and older<br>in the following categories:<br>Persons Living in or Traveling to Endemic Areas: While the actual risk for contracting yellow fever during<br>travel is probably low, variability of timeraries, behaviors and seasonal incidence of disease make it<br>diffuict to predict the actual risk for a given individual living in or traveling to a honove medmei co<br>epidemic area. Greater risk associated with living in or traveling to a ransove medmei co<br>areas of countries that do not officially reported at the time of travel and with traveling uotside the urban<br>areas of countries that do not officially report the disease but that lie in a yellow fever endemic zone.<br>Persons Traveling Internationally Through Countries with Yellow Zever: Some countries require an | 1                                 | 9 months   | N/A         | N/A                    | Y               | N                               |   | 10/27/2023           |
| Vaccines      | 90723         | Diphtheria, tetanus toxoids,<br>acellular pertussis vaccine,<br>hepatitis B, and inactivated<br>poliovirus vaccine,- (DTaP-<br>HepB-IPV) for intramuscular<br>use                                   | 0.5 mL                     | 1/1/2001                | Pediarix®   | diphtheria and tetanus<br>toxoids and acellular<br>pertussis adsorbed, hepatitis<br>b (recombinant) and<br>inactivated poliovirus<br>vaccine, suspension for<br>intramuscular injection | Indicated for active immunization against diphtheria, tetanus, pertussis, infection caused by all known<br>subtypes of hepatitis B virus, and poliomyelitis. Pediarix is approved for use as a three-dose series in<br>infants born of hepatitis B surface antigen (HBSAg)-negative mothers. Pediarix may be given as early as 6<br>weeks of age through 6 years of age (prior to the 7th birthday).  | 1                                 | 6 weeks  | 6 years     | N/A                    | Y               | N                               |   | 7/2/2018             |
| Vaccines      | 90732         | Pneumococcal polysaccharide<br>vaccine, 23-valent (PPSV23),<br>adult or immunosuppressed<br>patient dosage, for use in<br>individuals 2 years or older, for<br>subcutaneous or<br>intramuscular use | 0.5 mL                     | 1/1/2002                | Pneumovax <sup>®</sup> 23                               | pneumococcal vaccine<br>polyvalent sterile, liquid<br>vaccine for intramuscular or<br>subcutaneous injection  | <ul> <li>Indicated for active immunization for the prevention of pneumococcal disease caused by the 23<br/>serotypes contained in the vaccine (1, 2, 3, 4, 5, 68, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14, 15B, 17F, 18C, 19F,<br/>19A, 20, 22F, 23F, and 33F).</li> <li>Phenumovax 23 is approved for use in persons 50 years of age or older and persons aged greater than or<br/>equal to 2 years who are at increased risk for pneumococcal disease.</li> </ul>  | 1                                 | 2 years  | N/A         | N/A                    | Y               | N                               |   | 7/3/2018             |
| Vaccines      | 90734         | Meningococcal conjugate<br>vaccine, serogroups A, C, W, Y,<br>quadrivalent, diptheria toxoid<br>carrier (MenACWY-D) or<br>CRM197 carrier (MenACWY-<br>CRM), for intramuscular use                   | 0.5 mL                     | 1/1/2017                | Menactra®,<br>Menveo                                    | meningococcal (groups a, c,<br>γ, and w-135) polysaccharide<br>diphtheria toxoid conjugate<br>vaccine solution for<br>intramuscular injection   | Menactra:<br>indicated for active immunization to prevent invasive meningococcal disease caused by Neisseria<br>meningitidis serogroups A, C, Y and W-135. Menactra is approved for use in individuals 9 months through<br>55 years of age. Menactra does not prevent N meningitidis serogroup B disease.<br>Menveo:<br>midicated for active immunization to prevent invasive meningococcal disease caused by Neisseria   | 1                                 | Product Specific Age<br>Restrictions<br>(see comments) | 55 years    | N/A                    | Y               | N                               | Product specific age<br>restrictions:<br>• Menactra: 9 months through<br>55 years of age<br>• Menveo: 2 months through<br>55 years of age | 1/26/2024            |
| Vaccines      | 90738         | Japanese encephalitis virus<br>vaccine, inactivated, for<br>intramuscular use   | 0.5 mL                     | 7/1/2008                | lxiaro®   | Japanese encephalitis<br>vaccine, inactivated,<br>adsorbed suspension for<br>intramuscular injection  | Indicated for active immunization for the prevention of disease caused by Japanese encephalitis virus<br>(JEV). Ixiaro is approved for use in individuals 2 months of age and older.  | 2                                 | 2 months   | N/A         | N/A                    | Y               | N                               |   | 10/27/2023           |
| Vaccines      | 90739         | Hepatitis B vaccine (HepB),<br>CpG-adjuvanted, adult dosage,<br>2 dose or 4 dose schedule, for<br>intramuscular use   | 0.5 mL                     | 1/1/2013                | Heplisav-B <sup>®</sup>                                 | hepatitis b vaccine<br>(recombinant), adjuvanted<br>solution for intramuscular<br>injection   | Indicated for prevention of infection caused by all known subtypes of hepatitis B virus in adults 18 years of<br>age and older.   | 2                                 | 18 years   | N/A         | N/A                    | Y               | N                               |   | 6/6/2022             |
| Vaccines      | 90740         | Hepatitis B vaccine (HepB),<br>dialysis or immunosuppressed<br>patient dosage, 3-dose<br>schedule, for intramuscular<br>use   | 40 mcg                     | 1/1/2001                | Recombivax HB®<br>Dialysis<br>Formulation               | hepatitis b vaccine, dialysis<br>patient dosage (3 dose<br>schedule), for intramuscular<br>use  | Recombivax HB Dialysis Formulation is approved for use in adult predialysis and dialysis patients 18 years<br>of age and older for prevention of infection caused by all known subtypes of hepatitis B virus.   | 2                                 | 18 years   | N/A         | N/A                    | Y               | N                               |   | 10/31/2018           |
| Vaccines      | 90743         | Hepatitis B vaccine (HepB),<br>adolescent, 2-dose schedule,<br>for intramuscular use  | 1 mL                       | 1/1/2001                | Recombivax HB®  | hepatitis B vaccine<br>(recombinant) suspension for<br>intramuscular injection (2<br>dose schedule)   | indicated for prevention of infection caused by all known subtypes of hepatitis B virus. Recombivax HB is<br>approved for use in individuals of all ages.<br>Recombivax HB Dialysis Formulation is approved for use in predialysis and dialysis patients 18 years of age  | 1                                 | 11 years   | 15 years    | N/A                    | Y               | N                               |   | 9/28/2021            |
| Vaccines      | 90744         | Hepatitis B vaccine (HepB),<br>pediatric/adolescent dosage, 3<br>dose schedule, for<br>intramuscular use  | 0.5 mL                     | 1/1/2000                | Engerix B®<br>Pediatric,<br>Recombivax HB®<br>Pediatric | hepatitis b vaccine,<br>pediatric/adolescent dosage<br>(3 dose schedule), for<br>intramuscular use  | Hepatitis B vaccination is appropriate for people expected to receive human alpha-1 proteinase inhibitor<br>that is produced from heat-treated, pooled human plasma that may contain the causative agents of<br>hepatitis and other viral diseases.   | 2                                 | N/A  | 19 years    | N/A                    | Y               | N                               |   | 10/31/2018           |
| Vaccines      | 90746         | Hepatitis B vaccine (HepB),<br>adult dosage, 3 dose schedule,<br>for intramuscular use  | 1 mL                       | 1/1/2000                | Engerix B <sup>®</sup> ,<br>Recombivax HB <sup>®</sup>  | hepatitis b vaccine<br>(recombinant) suspension for<br>intramuscular injection for<br>adult use, 3 dose schedule  | Indicated for immunization against infection caused by all known subtypes of hepatitis B virus.   | 1                                 | 20 years   | N/A         | N/A                    | Y               | N                               |   | 9/21/2018            |

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| medicaid/medi<br>Category | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name  | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|---------------------------|---------------|--|----------------------------|-------------------------|-------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Vaccines                  | 90747         | Hepatitis B vaccine (HepB),<br>dialysis or immunosuppressed<br>patient dosage, 4-dose<br>schedule, for intramuscular<br>use  | 40 mcg                     | 1/1/2000                | Engerix B*  | hepatitis b vaccine, dialysis or<br>immunosuppressed patient<br>dosage (4 dose schedule), for<br>intramuscular use  | This schedule is designed for certain populations (e.g. dialysis patients, neonates born of hepatitis B-<br>infected mothers, others who have or might have been recently exposed to the virus, certain travelers to<br>high-risk areas) for immunization against infection caused by all known subtypes of hepatitis B virus.  | 2                                 | N/A         | N/A         | N/A                    | Y               | Ν                               |  | 10/31/2018            |
| Vaccines                  | 90750         | Zoster (shingles) vaccine,<br>(HZV), recombinant, sub-unit,<br>adjuvanted, for intramuscular<br>injection  | 0.5 mL                     | 1/1/2017                | Shingrix    | zoster vaccine recombinant,<br>adjuvanted, suspension for<br>intramuscular injection  | Indicated for prevention of herpes zoster (HZ) (shingles) in adults aged 50 years and older.<br>Indicated for prevention of herpes zoster (HZ) (shingles) in adults aged 18 years and older who are or will<br>be at increased risk of HZ due to immunodeficiency or immunosuppression caused by known disease or<br>therapy.<br>Limitations of Use:<br>• Shingrix is not indicated for prevention of primary varicella infection (chickenpox).   | 2                                 | 19 years    | N/A         | N/A                    | Y               | N                               | ACIP recommends for 2 19<br>years of age in<br>immunodeficient or<br>immunosuppressed adults | 11/4/2021             |
| Vaccines                  | 90759         | Hepatitis B vaccine (HepB), 3-<br>antigen (S, Pre-S1, Pre-S2), 10<br>mcg dosage, 3 dose schedule,<br>for intramuscular use   | 10 mcg                     | 1/1/2022                | PreHevbrio™ | hepatitis b vaccine<br>(recombinant) injectable<br>suspension, for intramuscular<br>use   | Indicated for prevention of infection caused by all known subtypes of hepatitis B virus in adults 18 years o<br>age and older.  | f 2                               | 18 years    | N/A         | N/A                    | Ŷ               | N                               |  | 3/30/2022             |
| Vaccines                  | 91304         | Severe acute respiratory<br>syndrome coronavirus 2 (SARS<br>CoV-2) (coronavirus disease<br>[COVID-19]) vaccine,<br>recombinant spike protein<br>nanoparticle, saponin-based<br>adjuvant, preservative free, 5  | 0.5 mL (5 mcg)             | 7/13/2022               | N/A         | Novavax COVID-19 Vaccine,<br>Adjuvanted suspension for<br>injection, for intramuscular<br>use (2023-2024 Formula)   | Emergency Use Authorization:<br>The U.S. Food and Drug Administration (FDA) has issued an Emergency Use Authorization (EUA) for the<br>emergency use of Novavax COVID-19 Vaccine, Adjuvanted (2023-2024 Formula) for active immunization<br>to prevent coronavirus disease 2021 (COVID-19) caused by severe acute respiratory syndrome coronaviru<br>2 (SARS-CoV-2) in individuals 12 years of age and older.   | s 2                               | 12 years    | N/A         | N/A                    | Ŷ               | N                               | 9/2023: Aligned procedure<br>code effective date with CMS<br>effective date.                 | 10/26/2023            |
| Vaccines                  | 91318         | Severe acute respiratory<br>syndrome coronavirus 2   | 0.3 mL (3 mcg)             | 9/11/2023               | N/A         | Pfizer-BioNTech COVID-19<br>Vaccine suspension for  | The U.S. Food and Drug Administration (FDA) has issued an Emergency Use Authorization (EUA) for the<br>emergency use of Pfizer-BioNTech COVID-19 Vaccine (2023-2024 Formula) for active immunization to   | 2                                 | 6 months    | 4 years     | N/A                    | Y               | N                               |  | 9/18/2023             |
| Vaccines                  | 91319         | [SARSCOV-2] (coronavirus<br>Severe acute respiratory<br>syndrome coronavirus 2<br>(SARSCOV-2) (coronavirus<br>disease [COVID-19]) vaccine,<br>mRNA-LNP, spike protein, 10<br>mcg/0.3 mL dosage, tris-<br>sucrose formulation, for<br>intramuscular use | 0.3 mL (10 mcg)            | 9/11/2023               | N/A         | injection, for intramuscular<br>Pfizer-BioNTech COVID-19<br>Vaccine suspension for<br>injection, for intramuscular<br>use - 5 years through 11<br>years of age (2023-2024<br>Formula) | prevent coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2<br>The U.S. Food and Drug Administration (FDA) has issued an Emergency Use Authorization (EUA) for the<br>emergency use of Pfizer-BioNTech COVID-19 Vaccine (2023-2024 Formula) for active immunization to<br>prevent coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2<br>(SARS-CoV-2) in individuals 5 years through 11 years of age. | 1                                 | 5 years     | 11 years    | N/A                    | Y               | N                               |  | 9/18/2023             |
| Vaccines                  | 91320         | Severe acute respiratory<br>syndrome coronavirus 2<br>(SARSCoV-2) (coronavirus<br>disease [COVID-19]) vaccine,   | 0.3 mL                     | 9/11/2023               | Comirnaty®  | Pfizer-BioNTech COVID-19<br>Vaccine, mRNA suspension<br>for injection, for<br>intramuscular use - 12 years  | Indicated for active immunization to prevent coronavirus disease 2019 (COVID-19) caused by severe acute<br>respiratory syndrome coronavirus 2 (SARS-CoV-2) in individuals 12 years of age and older.  | 2 1                               | 12 years    | N/A         | N/A                    | Y               | N                               |  | 9/18/2023             |
| Vaccines                  | 91321         | Severe acute respiratory<br>syndrome coronavirus 2 (SARS<br>CoV-2) (coronavirus disease<br>[COVID-19]) vaccine, mRNA-  | 0.25 mL                    | 9/11/2023               | N/A         | Suspension for injection, for<br>intramuscular use - 6 months   | The U.S. Food and Drug Administration (FDA) has issued an Emergency Use Authorization (EUA) for the<br>emergency use of Moderna COVID-19 Vaccine (2023-2024 Formula) for active immunization to prevent<br>coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-<br>CoV-2) in individuals 6 months through 11 years of age.  | 1                                 | 6 months    | 11 years    | N/A                    | Y               | N                               |  | 9/18/2023             |
| Vaccines                  | 91322         | Severe acute respiratory<br>syndrome coronavirus 2 (SARS<br>CoV-2) (coronavirus disease<br>[COVID-19]) vaccine, mRNA-<br>LNP, 50 mcg/0.5 mL dosage,<br>for intramuscular use   | 0.5 mL                     | 9/11/2023               | Spikevax™   | Moderna COVID-19 Vaccine,<br>mRNA Suspension for<br>injection, for intramuscular<br>use - 12 years of age and<br>older (2023-2024 Formula)  | Indicated for active immunization to prevent coronavirus disease 2019 (COVID-19) caused by severe acute<br>respiratory syndrome coronavirus 2 (SARS-CoV-2) in individuals 12 years of age and older.  | 2 1                               | 12 years    | N/A         | N/A                    | Y               | N                               |  | 9/18/2023             |

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| medicaid/med | icaid-ncci-ed | lit-files   |                            |                         |                           |   |  |                                   |   |             |                        |                 |                                 | *   |                       |
|--------------|---------------|---|----------------------------|-------------------------|---------------------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Drugs        | J0121         | Injection, omadacycline, 1 mg   | ; 1 mg                     | 10/1/2019               | Nuzyra™                   | omadacycline for injection,<br>for intravenous use                                    | Indicated for the treatment of adult patients with the following infections caused by susceptible<br>microorganisms:<br>• Community-acquired bacterial pneumonia (CABP)<br>• Acute bacterial skin and skin structure infections (ABSSSI)<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Nuzyra and other  | 1,500                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 9/27/2019             |
| Drugs        | J0122         | Injection, eravacycline, 1 mg   | 1 mg                       | 10/1/2019               | Xerava**                  | eravacycline for injection, for<br>intravenous use                                    | Indicated for the treatment of complicated intra-abdominal infections in patients 18 years of age and<br>older.<br>Limitations of Use:<br>Xerava is not indicated for the treatment of complicated urinary tract infections (cUTI).  | 7,000                             | 18 years  | N/A         | N/A                    | ¥               | Y                               |   | 9/27/2019             |
| Biologicals  | J0129         | Injection, abatacept, 10 mg   | 10 mg                      | 1/1/2007                | Orencia®                  | abatacept injection, for<br>intravenous use   | Treatment of:<br>• Adult Rheumatoid Arthritis (RA): moderately to severely active RA in adults. Orencia may be used as<br>monotherapy or concomitantly with DMARDs other than TNF antagonists.<br>• Juvenile Idiopathic Arthritis: moderately to severely active polyarticular juvenile idiopathic arthritis in<br>patients 2 years of age and older. Orencia may be used as monotherapy or concomitantly with<br>methotrexate.<br>• Active Psoroitik Arthritis (PAA) in adults.<br>Indicated for prophylaxis of:<br>• Acute graft versus host disease (aGVHD): in combination with a calcineurin inhibitor and methotrexate,<br>in adults and pediatric patients 2 years of age and older undergoing hematopoietic stem cell<br>transplantation (HSCT) from a matched or 1 allele-mismatched unrelated donor.<br>Important Limitations of Use:<br>• Should not be given concomitantly with TNF antagonists. | 400                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ÿ                               | Indication specific age<br>restrictions:<br>• RA and PA-X 18 years of age<br>and older<br>• JIA and aGVHD: 2 years of<br>age and older                                      | 1/14/2022             |
| Drugs        | J0133         | Injection, acyclovir, 5 mg  | 5 mg                       | 1/1/2006                | N/A                       | acyclovir sodium, for<br>injection, for intravenous<br>infusion                       | Indicated for:<br>• Herpes simplex infections in immunocompromised patients<br>• Initial episodes of herpes genitalis<br>• Herpes simplex encephalitis<br>• Neonatal herpes simplex virus infection<br>• Varicella-zoster infections in immunocompromised patients   | 8,400                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• Herpes Simplex Infections:<br>Mucosal and Cutaneous<br>Herpes Simplex (HSV-1 and<br>HSV-2) Infections in<br>Immunocompromised | 5/14/2019             |
| Drugs        | J0153         | Injection, adenosine, 1 mg,<br>(not to be used to report any<br>adenosine phosphate | 1 mg                       | 1/1/2015                | Adenocard®,<br>Adenoscan® | adenosine injection, for<br>intravenous use   | Adenoscan: Adjunct to thallium-201 myocardial perfusion scintigraphy in patients unable to exercise<br>adequately.   | 118                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Product specific age<br>restrictions:<br>Adenoscan: 18 years of age   | 5/6/2019              |
| Drugs        | J0171         | Injection, adrenalin,<br>epinephrine, 0.1 mg  | 0.1 mg                     | 1/1/2011                | Adrenalin®                | epinephrine injection, for<br>intramuscular or<br>subcutaneous use                    | Indicated for emergency treatment of allergic reactions (Type 1), including anaphylaxis  | N/A                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 10/26/2018            |
| Biologicals  | J0177         | Injection, aflibercept hd, 1 mg   | ; 1 mg                     | 4/1/2024                | Eylea® HD                 | aflibercept injection, for<br>intravitreal use  | Indicated for the treatment of patients with:<br>• Neovascular (Wet) Age-Related Macular Degeneration (nAMD)<br>• Diabetic Macular Gebma (DME)<br>• Diabetic Retinopathy (DR)  | 32                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 4/12/2024             |
| Biologicals  | J0178         | Injection, aflibercept, 1 mg  | 1 mg                       | 1/1/2013                | Eylea®                    | aflibercept injection for<br>intravitreal injection                                   | indicated for:<br>• Neovascular (Wet) Age-Related Macular Degeneration (AMD)<br>• Macular Edema Following Retinal Vein Occlusion (RVO)   | 8                                 | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | AMD, RVO, DME, DR: 18 years<br>of age and older<br>ROP: N/A   | 3/16/2023             |
| Biologicals  | J0179         | Injection, brolucizumab-dbll, 1<br>mg   | 1 mg                       | 1/1/2020                | Beovu®                    | brolucizumab-dbll injection,<br>for intravitreal injection                            | Indicated for the treatment of:<br>- Neovascular (Wet) Age-Related Macular Degeneration (AMD)<br>- Diabetic Macular Edema (DME)  | 24                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 6/9/2022              |
| Drugs        | J0180         | Injection, agalsidase beta, 1<br>mg   | 1 mg                       | 1/1/2005                | Fabrazyme®                | agalsidase beta injection,<br>powder, lyophilized for<br>solution for intravenous use | Indicated for treatment of adult and pediatric patients 2 years of age and older with confirmed Fabry<br>disease.  | 420                               | 2 years   | N/A         | N/A                    | Y               | Y                               |   | 4/26/2021             |

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|--------------------------|---------------|--|----------------------------|-------------------------|---|--|--|-----------------------------------|--|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Drugs                    | J0184         | Injection, amisulpride, 1 mg   | 1 mg                       | 1/1/2024                | Barhemsys®                                | amisulpride injection, for<br>intravenous use                                      | Indicated in adults for:<br>• Prevention of postoperative nausea and vomiting (PONV), either alone or in combination with an<br>antiemetic of a different class.<br>• Treatment of PONV in patients who have received antiemetic prophylaxis with an agent of a different<br>class or have not received prophylaxis.   | 50                                | 18 years   | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 12/22/2023            |
| Drugs                    | J0185         | Injection, aprepitant, 1 mg  | 1 mg                       | 1/1/2019                | Cinvanti™                                 | aprepitant injectable  | Indicated in adults, in combination with other antiemetic agents, for the prevention of:<br>• acute and delayed nausea and vomiting associated with initial and repeat courses of highly emetogenic  | 650                               | 18 years   | N/A         | N/A                    | Y               | Y                               | 9/2023: Max monthly units<br>updated from 390 units to 650 | 9/28/2023             |
| Biologicals              | J0202         | Injection, alemtuzumab, 1 mg   | g 1 mg                     | 1/1/2016                | Lemtrada*                                 | alemtuzumab injection, for<br>intravenous use                                      | Indicated for the treatment of patients with relapsing forms of multiple scierosis (MS).   | 60                                | 17 years   | N/A         | N/A                    | Y               | Ŷ                               |  | 7/2/2018              |
| Drugs                    | J0207         | Injection, amifostine, 500 mg  | : 500 mg                   | 1/1/2000                | Ethyol®                                   | amifostine for injection   | Indicated to:<br>• Reduce the incidence of moderate to severe xerostomia in patients undergoing postoperative radiation<br>treatment of head and neck cancer.<br>• Reduce the cumulative renal toxicity associated with repeated administration of cisplatin in patients<br>with advanced ovarian cancer, where the radiation port includes a substantial portion of the parotid<br>glands.  | 155                               | 18 years   | N/A         | N/A                    | Y               | Ŷ                               |  | 9/25/2018             |
| Drugs                    | J0208         | Injection, sodium thiosulfate<br>(pedmark), 100 mg                                   | 100 mg                     | 4/1/2023                | Pedmark <sup>®</sup>                      | sodium thiosulfate injection,<br>for intravenous use                               | Indicated to reduce the risk of ototoxicity associated with cisplatin in pediatric patients 1 month of age<br>and older with localized, non-metastatic solid tumors.<br>Limitations of Use:<br>The safety and efficacy of Pedmark have not been established when administered following cisplatin<br>infusions longer than 6 hours. Pedmark may not reduce the risk of ototoxicity when administered<br>following longer cisplatin infusions, because irreversible ototoxicity may have already occurred.  | 5,000                             | 1 month  | 18 years    | N/A                    | Y               | Ŷ                               |  | 3/22/2024             |
| Drugs                    | J0210         | Injection, methyldopate HCl,<br>up to 250mg  | 250 mg                     | 1/1/2000                | N/A                                       | methyldopate hydrochloride<br>injection  | Indicated for hypertension, when parenteral medication is indicated. The treatment of hypertensive crises<br>may be initiated with methyldopate HCI injection.   | 496                               | N/A  | N/A         | N/A                    | Y               | Y                               |  | 10/26/2018            |
| Biologicals              | J0217         | Injection, velmanase alfa-tycv<br>1 mg   | , 1 mg                     | 1/1/2024                | Lamzede®                                  | velmanase alfa-tycv for<br>injection, for intravenous use                          | Indicated for the treatment of non-central nervous system manifestations of alpha-mannosidosis in adult  | 700                               | N/A  | N/A         | N/A                    | Y               | Y                               |  | 12/21/2023            |
| Biologicals              | J0218         | Injection, olipudase alfa-rpcp,<br>1 mg  | 1 mg                       | 4/1/2023                | Xenpozyme™                                | olipudase alfa-rpcp for<br>injection, for intravenous use                          | Indicated for treatment of non-central nervous system manifestations of acid sphingomyelinase<br>deficiency (ASMD) in adult and pediatric patients.  | 1,260                             | N/A  | N/A         | N/A                    | Y               | Y                               |  | 3/16/2023             |
| Biologicals              | J0219         | Injection, avalglucosidase<br>alfa-ngpt, 4 mg  | 4 mg                       | 4/1/2022                | Nexviazyme™                               | avalglucosidase alfa-ngpt for<br>injection, for intravenous use                    | Indicated for the treatment of patients 1 year of age and older with late-onset Pompe disease (lysosomal<br>acid alpha-glucosidase (GAA) deficiency).  | 2,100                             | 1 year   | N/A         | N/A                    | Y               | Y                               |  | 3/17/2022             |
| Biologicals              | J0221         | Injection, alglucosidase alfa,<br>(Lumizyme), 10 mg                                  | 10 mg                      | 1/1/2012                | Lumizyme®                                 | alglucosidase alfa for<br>injection, for intravenous use                           | A hydrolytic lysosomal glycogen-specific enzyme indicated for patients with Pompe disease (GAA deficiency).  | 900                               | N/A  | N/A         | N/A                    | Y               | Y                               |  | 6/4/2019              |
| Drugs                    | J0222         | Injection, Patisiran, 0.1 mg   | 0.1 mg                     | 10/1/2019               | Onpattro™                                 | patisiran lipid complex<br>injection, for intravenous use                          | Indicated for the treatment of the polyneuropathy of hereditary transthyretin-mediated amyloidosis in<br>adults.   | 600                               | 18 years   | N/A         | N/A                    | Y               | Y                               |  | 9/27/2019             |
| Drugs                    | J0223         | Injection, givosiran, 0.5 mg   | 0.5 mg                     | 7/1/2020                | Givlaari™                                 | givosiran injection, for<br>subcutaneous use                                       | Indicated for the treatment of adults with acute hepatic porphyria (AHP).  | 1,512                             | 18 years   | N/A         | N/A                    | Y               | Y                               |  | 6/17/2020             |
| Drugs                    | J0224         | Injection, lumasiran, 0.5 mg   | 0.5 mg                     | 7/1/2021                | Oxlumo™                                   | lumasiran injection, for<br>subcutaneous use                                       | Indicated for the treatment of primary hyperoxaluria type 1 (PH1) to lower urinary and plasma oxalate<br>levels in pediatric and adult patients.   | 1,890                             | N/A  | N/A         | N/A                    | Y               | Y                               |  | 11/30/2022            |
| Drugs                    | J0225         | Injection, vutrisiran, 1 mg  | 1 mg                       | 1/1/2023                | Amvuttra™                                 | vutrisiran injection, for<br>subcutaneous use                                      | Indicated for the treatment of the polyneuropathy of hereditary transthyretin-mediated amyloidosis in<br>adults.   | 25                                | 18 years   | N/A         | N/A                    | Y               | Y                               |  | 12/6/2022             |
| Drugs                    | J0248         | Injection, remdesivir, 1 mg  | 1 mg                       | 12/23/2021              | Veklury®                                  | remdesivir injection, for<br>intravenous use                                       | Indicated for the treatment of coronavirus disease 2019 (COVID-19) in adults and pediatric patients (birth<br>to less than 18 years of age weighing at least 1.5 kg) who are:<br>• Hospitalized and have mild-to-moderate COVID-19, and are at high risk for progression to severe<br>COVID-19, including hospitalization or death.  | 400                               | Pediatric patients<br>from birth to less<br>than 28 days of age<br>weighing at least 1.5<br>kg |             | N/A                    | Y               | Y                               |  | 3/22/2024             |
| Biologicals              | J0256         | Injection, alpha 1-proteinase<br>inhibitor, human, 10 mg, not<br>otherwise specified | 10 mg                      | 1/1/2000                | Aralast NP®,<br>Prolastin-C®,<br>Zemaira® | alpha 1-proteinase inhibitor<br>(human) for intravenous use                        |  | 5,000                             | 18 years   | N/A         | N/A                    | Y               | Y                               |  | 6/6/2019              |
| Biologicals              | J0257         | Injection, alpha-1 proteinase<br>inhibitor (human), (Glassia),<br>10 mg              | 10 mg                      | 1/1/2012                | Glassia™                                  | alpha 1-proteinase inhibitor<br>(human) injection solution,<br>for intravenous use | Indicated for chronic augmentation and maintenance therapy in adults with clinically evident emphysema<br>due to severe hereditary deficiency of Alpha1-1P (lafpha1-anttryssin deficiency). Glassia increases<br>antigenic and functional (anti-neurophi leastase capacity, ANEC) serum levels and antigenic lung<br>epithelial lining fluid levels of alpha1-PI.<br>Umitations of USe:<br>• The effect of augmentation therapy with any Alpha1-PI, including Glassia, on pulmonary exacerbations<br>and on the progression of emphysema in alpha1-antitrypsin deficiency has not been conclusively<br>demonstrated in randomized, controlled clinical trails.<br>• Clinical data demonstrating the long-term effects of chronic augmentation and maintenance therapy of<br>individuals with classia are not available.<br>• Glassia is not indicated as therapy for lung disease in patients in whom severe Alpha1-PI deficiency has<br>not been established. | 4,200                             | 18 years   | N/A         | N/A                    | Y               | ¥                               |  | 9/25/2018             |

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| Category | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|----------|---------------|--|----------------------------|-------------------------|------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Drugs    | J0278         | Injection, amikacin sulfate,<br>100 mg                         | 100 mg                     | 1/1/2006                | N/A        | amikacin sulfate injection,<br>solution                                     | Indicated in the short-term treatment of serious infections due to susceptible strains of Gram-negative<br>bacteria, including Pseudomonas species, Echerichia coli, species of indole-positive and indole-negative<br>Protous, Providencia species, Klebsielia-Enterobacter-Serratia species, and Acinetobacter (Mima-Herellea)<br>species.<br>Clinical studies have shown amikacin sulfate injection to be effective in bacterial septicamia (including<br>neonatal sepsis); in serious infections of the respiratory tract, bones and joints, central nervous system<br>(including meningitis) and skin and soft tissue, intra-abdominal infections (including peritonitis); and in<br>burns and postoperative infections (including post-vascular surgery). Clinical studies have shown amikacin<br>also to be effective in serious complicated and recurrent unnary tract infections (including house and postoperative infections (including post-vascular surgery).   | 150                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 4/10/2019             |
| Drugs    | J0280         | Injection, aminophylline, up to<br>250mg                       | up to 250 mg               | 1/1/2000                | N/A        | aminophylline injection   | Indicated as an adjunct to inhaled beta-2 selective agonists and systemically administered corticosteroids<br>for the treatment of acute exacerbations of the symptoms and reversible airflow obstruction associated<br>with asthma and other chronic lung diseases, e.g., emphysema and chronic bronchitis.   | 217                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 9/25/2018             |
| Drugs    | J0285         | Injection, amphotericin B, 50<br>mg                            | 50 mg                      | 1/1/2000                | N/A        | amphotericin B for injection  | Amphotericin B for injection is specifically intended to treat potentially life-threatening fungal infections:<br>aspergliosis, cryptococcosis (torulosis), North American blastomycosis, systemic candidiasis,<br>coccidioidomycosis, histoplasmosis, zygomycosis including mucormycosis due to susceptible species of the<br>genera absidia, mucor and histopus, and infections due to related susceptible species for conditobolus and<br>basidiobolus, and sportorichosis. May be useful to treat American mucocutaneous leishmaniasis, but it is<br>not the drug of choice as primary therapy.  | 93                                | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 9/25/2018             |
| Drugs    | J0287         | Injection, amphotericin B lipid<br>complex, 10 mg              | 10 mg                      | 1/1/2003                | Abelcet®   | amphotericin B lipid complex<br>injection                                   | Indicated for the treatment of invasive fungal infections in patients who are refractory to or intolerant of<br>conventional amphotericin B therapy.   | 2,170                             | N/A   | N/A         | N/A                    | Y               | Y                               |   | 5/6/2019              |
| Drugs    | J0289         | Injection, amphotericin B<br>liposome, 10 mg                   | 10 mg                      | 1/1/2003                | AmBisome®  |   | Indicated for:<br>• Empirical therapy for presumed fungal infection in febrile, neutropenic patients<br>• Tratament of patients with Aspergillus species, Candida species, and/or Cryptococcus species infections<br>refractory to amphotericin B desoxycholate, or in patients where renal impairment or unacceptable<br>toxicity precludes the use of amphotericin B desoxycholate<br>• Treatment of Cryptococcal Menningtis in HVI-infected patients<br>• Treatment of visceral leishmaniasis. In immunocompromised patients with visceral leishmaniasis<br>treated with AmBisome, relapse rates were high following initial clearance of parasites.  | 2,604                             | 1 month   | N/A         | N/A                    | Y               | Ŷ                               |   | 4/10/2019             |
| Drugs    | J0290         | Injection, ampicillin sodium,<br>500 mg                        | 500 mg                     | 1/1/2000                | N/A        | ampicilin sodium for<br>injection, for intravenous or<br>intramuscular use  | Indicated in the treatment of infections caused by susceptible strains of the designated organisms in the<br>following conditions:<br>Bespiratory Treat Infections caused by Streptococcus pneumoniae, Staphylococcus aureus (penicillinase<br>and nonpenicillinase-producing), H. influenzae, and Group A beta-hemolytic streptococci.<br>• Bacterial Meningitis caused by E. coli, Group B streptococci, and other Gram-negative bacteria (Listeria<br>monocytogenes, N. meningitidis). The addition of an aminoglycoside with ampicillin may increase its<br>effectiveness against Gram-negative bacteria.<br>• Septicemia and Endocarititis caused by succeptible Gram-positive organisms including Streptococcus<br>spo, penicillin G-susceptible staphylococci, and enterococci. Gram-negative bacteria (Listeria<br>usually respond to intravenous therapy. The addition of an aminoglycoside may enhance the effectiveness<br>of ampicillin when treating streptococcuc ale dotamicillis.<br>• Urinary Tract Infections caused by sensoftites strains of E. coli and Proteus mirabilis.<br>• Gastrointestian Infections caused by Samonella to phyli (hybridi Greve), other Salmonella spp., and<br>Shigella spp. (dysentery) usually respond to oral or intravenous therapy.  | 1,736                             | N/A   | N/A         | N/A                    | Y               | Y                               |   | 4/10/2019             |
| Drugs    | J0291         | Injection, plazomicin, 5 mg                                    | 5 mg                       | 10/1/2019               | Zemdri™    | plazomicin injection, for<br>intravenous use                                | <ul> <li>Indicated for the treatment of patients 18 years of age or older with complicated urinary tract infections<br/>(LCUT) including pyelonephritis.</li> <li>As only limited clinical safety and efficacy data are available, reserve Zemdri for use in patients who<br/>have limited or no alternative treatment options.</li> <li>To reduce the development of drug-resistant bacteria and maintain effectiveness of Zemdri and other<br/>antibacterial drugs, Zemdri should be used only to treat infections that are proven or strongly suspected to<br/>be caused by susceptible microcorpanisms.</li> </ul>   | 2,940                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 10/3/2019             |
| Drugs    | J0295         | Injection, ampicillin<br>sodium/subactam sodium,<br>per 1.5 gm | per 1.5 gm                 | 1/1/2000                | Unasyn*    | ampicillin sodium and<br>subactam sodium injection,<br>powder, for solution | Indicated for the treatment of infection due to susceptible strains of the designated microorganisms in the<br>conditions listed below:<br>• Skin and skin structure infections caused by beta-lactamase producing strains of Staphylococcus aureus,<br>Echerichia coli, Klasbiella spo. (Including K. pneumoniae), Proteus mirabilis, Bacteroides fragilis,<br>Enterobacter spo., and Acinetobacter calcoaceticus.<br>• Intra-abdominal infections: caused by beta-lactamase producing strains of Escherichia coli, Klebsiella<br>spo. (Including K. pneumoniae), Bacteroides spo. (Including B. fragilis), and Enterobacter spo.<br>• Gynecological Infections: caused by beta-lactamase producing strains of Escherichia coli, and Bacteroides<br>spo. (Including K. fragilis).<br>• While Unasyn is indicated only for the conditions listed above, infections caused by ampicillin-<br>susceptible organisms are also amelable to treatment with Unasyn due to its ampicillin content.<br>Therefore, mixed infections: caused by ampicillin-susceptible organisms and beta-lactamase producing<br>organisms usuceptible to Unasyn should not require the addition of natorthar antibacterial.<br>• Appropriate culture and susceptibility tests should be performed before treatment in order to isolate<br>and identify the organisms causing infection and to determine their susceptibility to Unasyn. | 168                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | ¥               | ¥                               | Indication specific:<br>• Skin and skin structure<br>infections: 1 year of age and<br>older<br>• Intra-abdominal infections:<br>18 years of age and older | 6/7/2019              |
| Drugs    | J0300         | Injection, amobarbital, up to<br>125mg                         | up to 125 mg               | 1/1/2000                | Amytal®    | amobarbital sodium for<br>injection   | Indicated for use as a:<br>• Sedative<br>+ Nyportic, for the short-term treatment of insomnia, since it appears to lose its effectiveness for sleep<br>induction and sleep maintenance after 2 weeks<br>• Preanesthetic  | 112                               | 6 years   | N/A         | N/A                    | Y               | Y                               |   | 4/10/2019             |

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| medicaid/med | icaid-ncci-ed | it-files  |                            |                         |   |   |  |                                   |             |             |                        |                 |                                 |  |                       |
|--------------|---------------|---|----------------------------|-------------------------|---|---|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description                                     | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name  | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs        | J0330         | Injection, succinylcholine<br>chloride, up to 20mg    | up to 20 mg                | 1/1/2000                | Anectine <sup>®</sup> ,<br>Quelicin™                            | succinylcholine chloride<br>injection   | Indicated as an adjunct to general anesthesia, to facilitate tracheal intubation, and to provide skeletal<br>muscle relaxation during surgery or mechanical ventilation.   | 8                                 | N/A         | N/A         | N/A                    | Y               | Y                               |  | 9/21/2018             |
| Drugs        | J0349         | Injection, rezafungin, 1 mg                           | 1 mg                       | 10/1/2023               | Rezzayo™  | rezafungin for injection, for<br>intravenous use  | Indicated in patients 18 years of age or older who have limited or no alternative options for the treatment<br>of candidemia and invasive candidiasis.<br>Limitations of Use:<br>Rezzayo has not been studied in patients with endocarditis, osteomyelitis, and meningitis due to <i>Candida</i> .   | 1,000                             | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |  | 9/28/2023             |
| Drugs        | J0360         | Injection, hydralazine HCl, up<br>to 20mg             | up to 20 mg                | 1/1/2000                | N/A   | hydralazine hydrochloride<br>injection  | Indicated for severe essential hypertension when the drug cannot be given orally or when there is an<br>urgent need to lower blood pressure.   | 75                                | N/A         | N/A         | N/A                    | Y               | Y                               |  | 6/4/2019              |
| Drugs        | J0401         | Injection, aripiprazole (abilify maintena), 1 mg      | 1 mg                       | 1/1/2014                | Abilify Maintena*   | aripiprazole extended-release<br>injectable suspension, for<br>intramuscular use                                    | Indicated for the treatment of schizophrenia in adults.<br>Indicated for maintenance monotherapy treatment of bipolar I disorder in adults.  | 800                               | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 6/24/2024             |
| Drugs        | J0402         | Injection, aripiprazole (abilify asimtufii), 1 mg     | 1 mg                       | 1/1/2024                | Abilify Asimtufii®  | aripiprazole extended-release<br>injectable suspension, for<br>intramuscular use                                    | Indicated:<br>• for the treatment of schizophrenia in adults<br>• as maintenance monotherapy treatment of bipolar I disorder in adults   | 960                               | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 12/21/2023            |
| Drugs        | J0456         | Injection, azithromycin, 500<br>mg                    | 500 mg                     | 1/1/2000                | Zithromax®  | azithromycin for intravenous<br>infusion  | Indicated for mild to moderate infections caused by designated, susceptible bacteria in community-<br>acquired pneumonia in adults and pelvic inflammatory disease.  | 10                                | 16 years    | N/A         | N/A                    | Ŷ               | Y                               |  | 9/25/2018             |
| Drugs        | J0461         | Injection, atropine sulfate,<br>0.01 mg               | 0.01 mg                    | 1/1/2010                | N/A   | atropine sulfate injection for<br>intravenous, intramuscular,<br>subcutaneous, intraosseous,<br>or endotracheal use | Indicated for temporary blockade of severe or life threatening muscarinic effects.   | 27,900                            | N/A         | N/A         | N/A                    | Y               | Ŷ                               |  | 10/4/2018             |
| Drugs        | J0470         | Injection, dimercaprol, per<br>100mg                  | per 100 mg                 | 1/1/2000                | BAL in oil™   | dimercaprol injection   | Indicated in the treatment of:<br>• Arsenic, gold and mercury poisoning.<br>• Acute lead poisoning when used concomitantly with Edetate Calcium Disodium Injection.<br>Dimercaprol is effective for use in acute poisoning by mercury salts if therapy is begun within one or two<br>hours following ingestion. It is not very effective for chronic mercury poisoning. Dimercaprol is of<br>questionable value in poisoning by other heavy metals such as antimony and bismuth. It should not be<br>used in iron, cadmium, or selenium poisoning because the resulting dimercaprol-metal complexes are<br>more toxic than the metal alone, especially to the kidneys. | 252                               | N/A         | N/A         | N/A                    | ¥               | Ŷ                               |  | 6/7/2019              |
| Drugs        | J0475         | Injection, baclofen, 10 mg                            | 10 mg                      | 1/1/2000                | Gablofen®,<br>Lioresal®<br>Intrathecal                          | baclofen injection  | Indicated for use in the management of severe spasticity of cerebral or spinal origin in adult and pediatric<br>patients age 4 years and above.<br>Baddrein intrahecal should be reserved for patients unresponsive to oral baclofen therapy, or those who<br>experience intolerable central nervous system side effects at effective doses.<br>• Patients should first respond to a screening dose of intrathecal baclofen prior to consideration for long<br>term infusion via an implantable pump.<br>• Spasticity due to traumatic brain injury: wait at least one year after injury before considering baclofen<br>intrathecal therapy.                           | 8                                 | 4 years     | N/A         | N/A                    | Ŷ               | Ŷ                               | 5/2023: NC Suggested Max<br>Monthly Units updated to align<br>with NCTracks, which has been<br>set to 8 units/month since<br>9/1/2018. |                       |
| Drugs        | J0476         | Injection, baclofen, 50 mcg,<br>for intrathecal trial | 50 mcg                     | 1/1/2000                | Gablofen <sup>®</sup> ,<br>Lioresal <sup>®</sup><br>Intrathecal | baclofen injection, for<br>intrathecal trial  | Management of severe spasticity caused by spinal cord lesions or multiple sclerosis. Baclofen also is used<br>intrathecally in patients with spasticity of cerebral origin, including those with cerebral palsy and acquired<br>brain injury. Baclofen injection is designated an orphan drug by the FDA for the management of spasticity<br>in patients with cerebral palsy.  | 5                                 | N/A         | N/A         | N/A                    | Y               | Ŷ                               |  | 5/21/2019             |
| Biologicals  | J0485         | Injection, belatacept, 1 mg                           | 1 mg                       | 1/1/2013                | Nulojix®  | belatacept for injection, for<br>intravenous use  | Prophydaxs of organ rejection in adult patients receiving a kidney transplant. Use in combination with<br>basilikimab induction, mycophenolate mofetil, and corticosteroids.<br>Limitations of Use:<br>- Use only in patients who are EBV seropositive.<br>- Use has not been established for the prophylaxis of organ rejection in transplanted organs other than the<br>kidney.  | 6,000                             | 18 years    | N/A         | N/A                    | Y               | ¥                               |  | 6/6/2019              |
| Biologicals  | J0490         | Injection, belimumab, 10 mg                           | 10 mg                      | 1/1/2012                | Benlysta*   | belimumab injection, for<br>intravenous use   | Indicated for the treatment of patients aged 5 years and older with active, autoantibody-positive,<br>systemi clupus erythematosus who are receiving standard therapy.<br>Indicated for the treatment of patients aged 5 years and older with active lupus nephritis who are<br>receiving standard therapy.<br>Limitations of Use:<br>The efficacy of Benlysta has to been evaluated in patients with severe active central nervous system<br>lupus. Use of Benlysta has to recommended in this situation.   | 420                               | 5 years     | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 8/16/2022             |
| Biologicals  | J0491         | Injection, anifrolumab-fnia, 1<br>mg                  | 1 mg                       | 4/1/2022                | Saphnelo™   | anifrolumab-fnia injection,<br>for intravenous use  | Indicated for the treatment of adult patients with moderate to severe systemic lupus erythematosus<br>(SLE), who are receiving standard therapy.<br>Limitations of Use: The efficacy of Saphnelo has not been evaluated in patients with severe active lupus<br>nephritis or severe active central nervous system lupus. Use of Saphnelo is not recommended in these<br>situations.  | 600                               | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |  | 3/21/2022             |
| Drugs        | J0500         | Injection, dicyclomine HCl, up<br>to 20mg             | up to 20 mg                | 1/1/2000                | Bentyl®   | dicyclomine hydrochloride<br>injection for intramuscular<br>use   | Indicated for the treatment of functional bowel/irritable bowel syndrome.  | 8                                 | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Drugs        | J0515         | Injection, benztropine<br>mesylate, per 1 mg          | 1 mg                       | 1/1/2000                | Cogentin®   | benztropine mesylate<br>injection   | Indicated:<br>- for use as an adjunct in the therapy of all forms of parkinsonism.<br>- for use in the control of extrapyramidal disorders (except tardive dyskinesia) due to neuroleptic drugs<br>(e.g., phenothiazines).   | 248                               | 3 years     | N/A         | N/A                    | Y               | Y                               |  | 11/17/2021            |

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• Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name    | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|---------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Drugs       | 10558         | Injection, penicillin G<br>benzathine and penicillin G<br>procaine, 100,000 units                              | 100,000 units              | 1/1/2011                | Bicillin* C-R | penicillin G benzathine and<br>penicillin G procaine<br>injectable suspension                   | Indicated for the treatment of moderately severe infections due to penicillin G-susceptible<br>microorganisms that are susceptible to serum levels common to this particular dosage form. Therapy<br>should be guided by bacteriological studies (including susceptibility testing) and by clinical response.<br>Biollin C-R is indicated in the treatment of the following in adults and pediatric patients:<br>• Moderately severe to severe infections of the upper-respiratory tract, scartef tever, erysipelas, and skin<br>and soft-tissue infections due to susceptible steptococci. NOTE: Streptococci in Groups A, C, G, H, L, and<br>M are very sensitive to pencillin G. Other groups, including Group D (enterococci), are resistant. Penicillin<br>G sodium or potassium is recommended for streptococcal infections with bacteremia.<br>• Moderately severe pneumonia and ottis media due to susceptible Streptococcus pneumoniae. NOTE:<br>Severe pneumonia, empyrema, bacteremia, pericardits, meningits, peritonits, and arthritis of<br>pneumococcal etology are better treated with penicillin G sodium or potassium during the acute stage.<br>• When high, sustained serum levels are required, penicillin G sodium or potassium, either IM or IV, should<br>be used. This drug should not be used in the treatment of venereal diseases, including syphilis, gonorrhea,<br>yaws, bejel, and pinta.  | 96                                | N/A   | N/A         | N/A                    | Y               | Y                               |   | 8/24/2018             |
| Drugs       | J0561         | Injection, penicillin G<br>benzathine, 100,000 units   | 100,000 units              | 1/1/2011                | Bicillin® L-A | penicillin G benzathine<br>injectable suspension  | Indicated for the treatment of infections due to penicillin G-sensitive microorganisms that are susceptible<br>to the low and very prolonged serum levels common to this particular dosage form. Therapy should be<br>guided by bacteriological studies (Including sensitivity tests) and by clinical response. The following<br>infections will usually respond to adequate dosage of intramuscular penicillin 6 benzathine: mild to<br>moderate upper respiratory infections due to susceptible streptococci, venereal infections (syphills, yaws,<br>beg), and pina) and prophysical of rheumatic (ever and chorea.  | 96                                | N/A   | N/A         | N/A                    | Y               | Y                               |   | 8/24/2018             |
| Biologicals | J0565         | Injection, bezlotoxumab, 10<br>mg  | 10 mg                      | 1/1/2018                | Zinplava™     | bezlotoxumab injection, for<br>intravenous use  | Indicated to reduce recurrence of <i>Clostridioides difficile</i> infection (CDI) in adult and pediatric patients 1<br>year of age or older who are receiving antibacterial drug treatment for CDI and are high risk for CDI<br>recurrence.<br>Limitation of use: Zinplava is not indicated for the treatment of CDI. Zinplava is not an antibacterial drug.<br>Zinplava should only be used in conjunction with antibacterial drug treatment of CDI.   | 140                               | 1 year  | N/A         | N/A                    | Y               | Y                               |   | 6/19/2023             |
| Biologicals | J0567         | Injection, cerliponase alfa, 1<br>mg   | 1 mg                       | 1/1/2019                | Brineura®     | cerliponase alfa injection, for<br>intraventricular use   | Indicated to slow the loss of ambulation in symptomatic pediatric patients 3 years of age and older with<br>late infantile neuronal ceroid lipofuscinosis type 2 (CLN2), also known as tripeptidyl peptidase 1 (TPP1)<br>deficiency.  | 900                               | 3 years   | N/A         | N/A                    | Y               | Y                               |   | 7/2/2018              |
| Drugs       | J0577         | Injection, buprenorphine<br>extended-release (brixadi),<br>less than or equal to 7 days of<br>therapy          | f 1 syringe                | 4/1/2024                | Brixadi™      | buprenorphine extended-<br>release injection for<br>subcutaneous use CIII<br>(weekly)           | Indicated for the treatment of moderate to severe opioid use disorder in patients who have initiated<br>treatment with a single dose of a transmucosal buprenorphine product or who are already being treated<br>with buprenorphine.<br>Brixadi should be used as part of a complete treatment plan that includes counseling and psychosocial<br>support.   | 5                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 3/22/2024             |
| Drugs       | J0578         | Injection, buprenorphine<br>extended-release (brixadi),<br>greater than 7 days and up to<br>28 days of therapy | 1 syringe                  | 4/1/2024                | Brixadi™      | buprenorphine extended-<br>release injection for<br>subcutaneous use CIII<br>(monthly)          | Indicated for the treatment of moderate to severe opioid use disorder in patients who have initiated<br>treatment with a single dose of a transmucosal buprenorphine product or who are already being treated<br>with buprenorphine.<br>Brixadi should be used as part of a complete treatment plan that includes counseling and psychosocial<br>support.   | 2                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 3/22/2024             |
| Biologicals | J0584         | Injection, burosumab-twza 1<br>mg  | 1 mg                       | 1/1/2019                | Crysvita®     | burosumab-twza injection,<br>for subcutaneous use   | Indicated for:<br>• The treatment of X-linked hypophosphatemia (XLH) in adult and pediatric patients 6 months of age and<br>older.<br>• The treatment of FGF23-related hypophosphatemia in tumor-induced osteomalacia (TIO) associated<br>with phosphaturic mesenchymai tumors that cannot be curatively resected or localized in adult and<br>pediatric patients 2 years of age and older.   | 540                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• XLH: 6 months of age and<br>older<br>• TIO: 2 years of age and older  | 7/28/2020             |
| Biologicals | J0585         | injection,<br>onabotulinumtoxinA, 1 unit   | 1 unit                     | 1/1/2000                | Botox*        | onabotulinumtoxinA for<br>injection, for intramuscular,<br>intradetrusor, or intradermal<br>use | Indicated for:<br>• Treatment of overactive bladder (OAB) with symptoms of urge urinary incontinence, urgency, and<br>frequency, in adults who have an inadequate response to or are intolerant of an anticholinergic<br>medication<br>• Treatment of urinary incontinence due to detrusor overactivity associated with a neurologic condition<br>[e.g., spinal cord injury (SCI), multiple sclerosis (MSI) in adults who have an inadequate response to or are<br>intolerant of an anticholinergic medication<br>• Treatment of neurogenic detrusor overactivity (NDO) in pediatric patients 5 years of age and older who<br>have an inadequate response to or are intolerant of anticholinergic medication.<br>• Prophylaxis of headaches in adult patients with chronic migraine (215 days per month with headache<br>lasting a hours ad qor longer)<br>• Treatment of severe axillary hyperhidrosis that is inadequately managed by topical agents in adult<br>patients<br>• Treatment of severe axillary hyperhidrosis that is inadequately managed by topical agents in adult<br>patients<br>• Treatment of severe axillary hyperhidrosis that is inadequately managed by topical agents in adult<br>patients<br>• Treatment of trabismus in patients 12 years of age and older<br>• Treatment of trabismus in patients 12 years of age and older<br>• Treatment of trabismus in patients 12 years of age and older<br>• Treatment of hours. Safety and effectiveness of Botox have not been established for:<br>• Prophylaxis of episodic migraine [14 headache days or fewer per month]<br>• Treatment of hyperhidrosis in body areas other than axillary | 600 in 90 day interval            | N/A   | N/A         | N/A                    | Y               | ¥                               | 1/2023: NC Suggested Max<br>Monthly Units updated to align<br>with NCTracks, which has been<br>set to 600 units in 90 days<br>since 1/1/2019.<br>9/2023: NC Suggested Max<br>Monthly Units updated from 3<br>month interval to 9 day<br>interval to align with<br>NCTracks.<br>11/2023: Edited 1/2023 and<br>9/2023 comments for clarity. |                       |

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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                    | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modifie<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|-------------------------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|----------------------|
| Biologicals | J0586         | implant, 1 microgram   | 5 units                    | 1/1/2010                | Dysport*                      | abobotulinumtoxinA for<br>injection, for intramuscular<br>use   | <ul> <li>Treatment of adults with cervical dystonia.</li> <li>The temporary improvement in the appearance of moderate to severe glabellar lines associated with procerus and corrugator muscle activity in adult patients &lt;65 years of age.</li> <li>Treatment of spasticity in patients 2 years of age and older.</li> </ul>   | 300                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific<br>recommendations.<br>• Cervical Dystonia: 18 years<br>of age and older<br>• Glabellar Lines: 18 years of<br>age and older<br>• Upper Limb Spasticity: 2<br>years of age and older<br>Lower Limb Spasticity: 2<br>years of age and older  | 8/25/2020            |
| Biologicals | J0587         | Injection,<br>rimabotulinumtoxinB, 100<br>units  | 100 units                  | 1/1/2002                | Myobloc <sup>®</sup>          | rimabotulinumtoxin B<br>injection   | Indicated for:<br>- Treatment of adult patients with cervical dystonia to reduce the severity of abnormal head position and<br>neck pain associated with cervical dystonia.<br>- Treatment of chronic sialor/hea in adults.  | 100                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/27/2019            |
| Biologicals | J0588         | Injection,<br>incobotulinumtoxinA, 1 unit  | 1 unit                     | 1/1/2012                | Xeomin <sup>®</sup>           | incobotulinumtoxinA for<br>injection, for intramuscular or<br>intraglandular use                        | Indicated for the treatment or improvement of:<br>• Chronic sialorrhea in patients 2 years of age and older<br>• Upper limb spasticity in adults<br>• Upper limb spasticity in pediatric patients 2 to 17 years of age, excluding spasticity caused by cerebral<br>palsy<br>• Cervical dystonia in adults<br>• Blepharospasm in adults   | 600 in a 12-week<br>interval      | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Ŷ               | Ŷ                               | Indication specific age<br>restrictions:<br>Cervical dystonia and<br>blepharospasm: 18 years of<br>age and older<br>Upper limb spasticity and<br>chronic sialorthea: 2 years of<br>age and older<br>1/2023: KC Suggested Max<br>Monthly Units updated to align<br>with MUE values. (Previously<br>set to 400 units.) | 9/13/2023            |
| Drugs       | J0594         | Injection, busulfan, 1 mg  | 1 mg                       | 1/1/2007                | Busulfex®                     | busulfan injection for<br>intravenous use   | Indicated for use in combination with cyclophosphamide as a conditioning regimen prior to allogeneic<br>hematopoletic progenitor cell transplantation for chronic myelogenous leukemia (CML).  | 1,312                             | N/A   | N/A         | N/A                    | Y               | Y                               | <ul> <li>Upper Limb Spasticity: Safety<br/>and effectiveness in pediatric<br/>patients below the age of 2<br/>years have not been<br/>established.</li> </ul>  | 9/27/2018            |
| Drugs       | J0595         | Injection, butorphanol<br>tartrate, 1mg  | 1 mg                       | 1/1/2004                | N/A                           | butorphanol tartrate<br>injection   | Indicated:<br>• As a properative or pre-anesthetic medication<br>• As a supeoperative or pre-anesthetic medication<br>• For the relief of pain during labor, and<br>• For the management of pain severe enough to require an opioid analgesic and for which alternative<br>treatments are inadequate<br>Limitations of Use:<br>• Because of the risks of addiction, abuse, and misuse, with opioids, even at recommended dozes, reserve<br>butorphanol tartrate for us in patients for whom alternative treatment option (e.g. non-opioid analgesics):<br>- Have nor been tolerated, or at not expected to be tolerate<br>- Have nor provide adequate analgesia, or are not expected to provide adequate analgesia | 992                               | 18 years  | N/A         | N/A                    | Y               | Y                               | Lower Limb Spasticity: Safety<br>and effectiveness in pediatric<br>patients below the age of 2<br>years have not been<br>established.  | 9/27/2018            |
| Biologicals | J0596         | Injection, c-1 esterase<br>inhibitor (recombinant),<br>Ruconest, 10 units                                | 10 units                   | 1/1/2016                | Ruconest®                     | c1 esterase inhibitor<br>(recombinant) for<br>intravenous use, lyophilized<br>powder for reconstitution | Indicated for treatment of acute attacks in adult and adolescent patients with hereditary angloedema<br>(HAE).   | 3,360                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019            |
| Biologicals | J0597         | Injection, C-1 esterase<br>inhibitor (human), Berinert, 10<br>units                                      | 10 units                   | 1/1/2011                | Berinert®                     | c1 esterase inhibitor (human)<br>for intravenous use  | Treatment of acute abdominal, facial, or laryngeal hereditary angioedema (HAE) attacks in adult and<br>pediatric patients.   | 1,120                             | N/A   | N/A         | N/A                    | Y               | Ŷ                               |  | 4/10/2019            |
| Biologicals | J0598         | Injection, C1 esterase inhibitor<br>(human), Cinryze, 10 units   | 10 units                   | 1/1/2010                | Cinryze®                      | for intravenous use   | Indicated for routine prophylaxis against angioedema attacks in adults, adolescents and pediatric patients<br>(6 years of age and older) with hereditary angioedema (HAE).   | 2,750                             | 6 years   | N/A         | N/A                    | Y               | Y                               |  | 7/26/2018            |
| Drugs       | J0600         | Injection, edetate calcium<br>disodium, up to 1000 mg  | up to 1000 mg              | 1/1/2000                | Calcium Disodium<br>Versanate | edetate calcium disodium<br>injection for intravenous or<br>intramuscular use                           | Indicated for the reduction of blood levels and depot stores of lead in lead poisoning (acute and chronic)<br>and lead encephalopathy in both pediatric populations and adults.  | 15                                | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/10/2018           |
| Drugs       | 10606         | Injection, etelcalcetide, 0.1 mg   | 0.1 mg                     | 1/1/2018                | Parsabiv™                     | etelcalcetide injection, for<br>intravenous use   | Indicated for secondary hyperparathyroidism (HPT) in adult patients with chronic kidney disease (CKD) on<br>hemodialysis.<br>Limitations of Use:<br>Parsabiv has not been studied in adult patients with parathyroid carcinoma, primary hyperparathyroidism<br>or with CKD who are not on hemodialysis and is not recommended for use in these populations.  | 2,250                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 6/4/2019             |
| Drugs       | J0612         | Injection, calcium gluconate,<br>not otherwise specified, 10 mg  | 10 mg                      | 4/1/2023                | N/A                           | calcium gluconate injection,<br>for intravenous use   | Indicated for pediatric and adult patients for the treatment of acute symptomatic hypocalcemia.<br>Limitations of Use:<br>The safety of calcium gluconate injection for long term use has not been established.  | 124,000                           | N/A   | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 3/22/2024            |
| Drugs       | J0613         | Injection, calcium gluconate<br>(wg critical care), not<br>therapeutically equivalent to<br>j0612, 10 mg | 10 mg                      | 4/1/2023                | N/A                           | calcium gluconate injection,<br>for intravenous use (WG<br>Critical Care)                               | Calcium Gluconate in Sodium Chloride Injection is a form of calcium indicated for pediatric and adult<br>patients for the treatment of acute symptomatic hypocalcemia.<br>Limitations of Use: The safety of Calcium Gluconate Injection for long term use has not been established.  | 24,800                            | N/A   | N/A         | N/A                    | Ŷ               | Y                               |  | 3/22/2024            |
| Drugs       | J0636         | Injection, calcitriol, 0.1 mcg   | 0.1 mcg                    | 1/1/2003                | N/A                           | calcitriol injection  | Indicated in the management of hypocalcemia in patients undergoing chronic renal dialysis. It has been<br>shown to significantly reduce elevated parathyroid hormone levels. Reduction of PTH has been shown to<br>result in an improvement in real osteodystrophy.  | 560                               | 13 years  | N/A         | N/A                    | Y               | Y                               |  | 9/27/2018            |

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| medicaid/medi |               |  | 1                          |                         | 1   |   |   | 1                                 |  |             |                        | 1 1             | Rebating                        |   |                       |
|---------------|---------------|--|----------------------------|-------------------------|---|---|---|-----------------------------------|--|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                                    | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age  | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Biologicals   | J0638         | Injection, canakinumab, 1 mg   | 1 mg                       | 1/1/2011                | llaris*                                       | canakinumab injection, for<br>subcutaneous use                              | Indicated for the treatment of:<br>- Periodic Fever Syndromes:<br>- Cropoprin Associated Periodic Syndromes (CAPS), in adults and children 4 years of age and older<br>including: Familial Cold Autoinflammatory Syndrome (FCAS) and MucKle-Wells Syndrome (MWS).<br>- Turnor Necrossis Factor Receptor Associated Periodic Syndrome (TRAS) in adult and pediatric patients.<br>- Hypertimmunoglobulin D Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD) in adult and pediatric<br>patients.<br>- Familial Nediterranean Fever (FMF) in adult and pediatric patients.<br>- Active Systemic Livenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Livenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Livenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Livenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Livenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Active Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Adult Const Still's Disease:<br>- Adult Sing Systemic Juvenile Idiopathic Arthritis (SIIA) in patients aged 2 years and older.<br>- Adult Const Still's Disease (ASD) | 600                               | Indication Specific<br>Age Restrictions<br>(see comments)  | N/A         | N/A                    | ¥               | ¥                               | Indication specific age<br>restrictions:<br>• SIIA, AOSO, TRAPS,<br>HIDS/MKD, and FMF: 2 years<br>of age and older<br>• CAPS (FCAS and MWS): 4<br>years of age and older<br>• Gout flares: 18 years of age<br>and older | 9/28/2023             |
| Drugs         | J0640         | Injection, leucovorin calcium,<br>per 50 mg  | 50 mg                      | 1/1/2000                | N/A   | leucovorin calcium for<br>injection for intravenous or<br>intramuscular use | Indicated:<br>• After high dose methotrexate therapy in osteosarcoma.<br>• To diminish the toxicity and counteract the effects of impaired methotrexate elimination and of<br>inadvertent overdosages of folic acid antagonists.<br>• In the treatment of megalobasics anemias due to folic acid deficiency when oral therapy is not feasible.<br>• For use in combination with 5-fluorouracil to prolong survival in the palliative treatment of patients with<br>advanced colorectal cancer. Leucovorin should not be mixed in the same infusion as 5-fluorouracil because<br>a precipitate may form.   | 80                                | N/A  | N/A         | N/A                    | Y               | Y                               |   | 7/2/2018              |
| Drugs         | J0641         | Injection, levoleucovorin, not otherwise specified, 0.5 mg   | 0.5 mg                     | 1/1/2009                | Fusilev®                                      | levoleucovorin injection<br>solution for intravenous use                    | Indicated for:<br>• Rescue after high-dose methotrexate therapy in octeosarcoma.<br>• Oriminishing the toxicity and counteracting the effects of impaired methotrexate elimination and of<br>inadvertent overdosage of folic acid antagonists.<br>• Use in combination chemotherapy with 5-fluorouracil in the palliative treatment of patients with<br>advanced metastatic colorectal cancer.<br>Umitations of Use:  | 10,000                            | N/A  | N/A         | N/A                    | Y               | Y                               |   | 10/3/2019             |
|               |               |  |                            |                         |   |   | Fusilev is not approved for pernicious anemia and megaloblastic anemias. Improper use may cause a<br>hematologic remission while neurologic manifestations continue to progress.  |                                   |  |             |                        |                 |                                 |   |                       |
| Drugs         | J0642         | Injection, levoleucovorin<br>(khapzory), 0.5 mg  | 0.5 mg                     | 10/1/2019               | Khapzory™                                     | levoleucovorin for injection,<br>for intravenous use                        | Initiated for:<br>Initiated for:  | 4,800                             | N/A  | N/A         | N/A                    | Y               | Y                               |   | 10/3/2019             |
| Drugs         | 10665         | Injection, bupivacaine, not<br>otherwise specified, 0.5 mg   | 0.5 mg                     | 7/1/2023                | Marcaine™,<br>Sensorcaine®                    | injection, for infiltration,  | Bupivaciane hydrochloride injection:<br>• Indicated in adults for the production of local or regional anesthesia or analgesia for surgery, dental and<br>oral surgery procedures, diagnostic and therapeutic procedures, and for obstetrical procedures. For each<br>type of block indicated to produce local or regional anesthesia or analgesia, specific concentrations and<br>presentations are recommended.<br>• Limitations of Use: Not all blocks are indicated for use with bupivacaine given clinically significant risks  | 4,000                             | Formulation-specific<br>age restrictions (see<br>comments) | N/A         | N/A                    | Y               | ¥                               | Formulation-specific age<br>restrictions:<br>Bupivaciane hydrochloride<br>injection: 12 years of age and<br>older<br>Bupivaciane hydrochloride in<br>dextrose injection: 18 years of<br>age and older                   | 10/26/2023            |
| Drugs         | J0670         | Injection, mepivacaine<br>hydrochloride, per 10 mL   | 10 mL                      | 1/1/2000                | Carbocaine™,<br>Polocaine®,<br>Polocaine® MPF | mepivacaine hydrochloride<br>injection                                      | Carbocaine, Polocaine and Polocaine MPF: Indicated for production of local or regional analgesia and<br>anesthesia by local infiltration, peripheral nerve block techniques, and central neural techniques including<br>epidural and caudal blocks.   | 50                                | N/A  | N/A         | N/A                    | Y               | Y                               |   | 4/10/2019             |
| Drugs         | J0687         | Injection, cefazolin sodium (wg<br>critical care), not<br>therapeutically equivalent to<br>j0690, 500 mg | 500 mg                     | 7/1/2024                | N/A   | cefazolin for injection, for<br>intravenous use (WG Critical<br>Care)       | Cefazolin for injection is indicated for perioperative prophylaxis in adult patients.<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Cefazolin for<br>injection and other antibacterial drugs, Cefazolin for injection should be used only to treat or prevent<br>infections that are proven or strongly suspected to be caused by bacteria.   | 496                               | 18 years   | N/A         | N/A                    | Y               | Ŷ                               |   | 6/24/2024             |

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| ategory | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modifie<br>Date |
|---------|---------------|---|----------------------------|-------------------------|------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|----------------------|
| Drugs   | J0688         | Injection, cefazolin sodium<br>(hikma), not therapeutically<br>equivalent to j0690, 500 mg  | 500 mg                     | 1/1/2024                | N/A        | cefazolin for injection, for<br>intravenous use (Hikma)                     | Cefazolin for injection is a cephalosporin antibacterial indicated for:<br>• Treatment of the following infections caused by susceptible isolates of the designated microorganisms in<br>adult and pediatir patients 1 month of age and older for whom appropriate dosing with this formulation<br>can be achieved:<br>0 Bespiratory tract infections<br>0 Urinary tract infections<br>0 Skin and skin structure infections<br>0 Bilary tract infections<br>0 Bone and joint infections<br>0 Genital infections<br>0 Genital infections<br>0 Septemaia<br>0 Endocarditis<br>• Perioperative prophylaxis in adult patients<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of cefazolin for<br>infections that are proven or strongly suspected to be caused by bacteria.  | 496                               | 1 month   | N/A         | N/A                    | ¥               | Y                               |  | 6/25/2024            |
| Drugs   | 10689         | Injection, cefazolin sodium<br>(baxter), not therapeutically<br>equivalent to j0690, 500 mg | 500 mg                     | 1/1/2023                | N/A        | cefazolin in dextrose<br>injection, for intravenous use<br>(Baxter)         | Indicated for:<br>• Treatment of the following infections caused by susceptible isolates of the designated microorganisms in<br>adult and pediatric patients for whom appropriate dosing with this formulation can be achieved:<br>0 Respiratory tract infections<br>0 Urinary tract infections<br>0 Skin and skin structure infections<br>0 Bone and joint infections<br>0 Genital infections<br>0 Genital infections<br>0 Septicemia<br>0 Endocarditis<br>• Perioperative prophylaxis in adults and pediatric patients aged 10 to 17 years old for whom appropriate<br>dosing with this formulation can be achieved.<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of cefazolin injection<br>and other antibacterial drugs, cefazolin injection should be used only to treat or prevent infections that<br>are proven or strongly suspected to be caused by bacteria.   | 496                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Treatment of infections<br>caused by usceptible isolates<br>of the designated<br>microogramisms: 1 month and<br>older<br>• Perioperative prophylaxis: 10<br>years of age and older | 6/25/2024            |
| ugs     | 10690         | Injection, cefazolin sodium,<br>500 mg  | 500 mg                     | 1/1/2000                | N/A        | cefazolin sodium for injection  | Indicated for the treatment of the following serious interctions when due to susceptione organisms:<br>Respiratory Tract Infections: Due to 5 pneuroniais, Rebeidla species, N. Enforcas, 2. Surveis (penicillin-<br>resistive and penicillin-resistant), and group A beta-hemolytic streptococci. Injectable benzathine<br>penicillin is considered the drug of choice in treatment and prevention of streptococci. Injectable benzathine<br>penicillin is considered the drug of choice in treatment and prevention of streptococci. Injectable benzathine<br>penicillin is considered the drug of choice in treatment and prevention of streptococci. Injectable benzathine<br>the asopharyns; however, data establishing the efficacy of celaziolin in the subsequent prevention of<br>thematic fever a ront available at present.<br>• Urinary Tract Infections: Due to E. coli, P. mirabilis, Klebsiella species, and some strains of enterobacter<br>and enterococci.<br>• Skin and Skin Structure Infections: Due to S. aureus (penicillin-sensitive and penicillin-resistant), group A<br>beta-hemolytic streptococci, and other strains of streptococci.<br>• Bilary Tract Infections: Due to S. aureus.<br>• Genital Infections: Due to S. aureus.<br>• Septemar: Due to S. preumoniae, S. aureus (penicillin-sensitive and penicillin-resistant), P. mirabilis, E.<br>• Signardina Species.<br>• Septemar: Due to S. aureus (penicillin-sensitive and penicillin-resistant), P. mirabilis, E.<br>• Septemar: Due to S. aureus (penicillin-sensitive and penicillin-resistant), P. mirabilis, E.<br>• Septemar: Due to S. aureus (penicillin-sensitive and penicillin-resistant), P. mirabilis, E.<br>• Septemar: Due to S. aureus (penicillin-resistant and group A betahemolytic<br>streptococci.<br>Perioperative Prophylaxis: The prophylactic administration of cefazolin preoperatively, intraoperatively,<br>and postoperatively may reduce the incidence of certain postoperative infections in patients undergoing<br>surgical procedures which are classified as contaminated or potentially contaminated (e.g., vaginal<br>hysterectomy, and cholecystec | 495                               | 1 month   | N/A         | N/A                    | Y               | Ÿ                               |  | 6/25/2024            |
| ıgs     | J0691         | Injection, lefamulin, 1 mg  | 1 mg                       | 7/1/2020                | Xenleta™   | lefamulin injection, for<br>intravenous use                                 | Indicated for the treatment of adults with community-acquired bacterial pneumonia (CABP) caused by the<br>following susceptible microorganisms: Streptococcus pure pneumoniae, Staphylococcus aureus (methicillin-<br>susceptible isolates), Haemophilus influenzae, Legionella pneumophila, Mycoplasma pneumoniae, and<br>Chiamydophila pneumoniae.<br>To reduce the development of drug resistant bacteria and maintain the effectiveness of Xenleta and other<br>antibacterial drugs, Xenleta should be used only to treat or prevent infections that are proven or strongly  | 2,100                             | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 6/17/2020            |
| rugs    | J0692         | Injection, cefepime HCl, 500<br>mg  | 500 mg                     | 1/1/2002                | Maxipime™  | cefepime hydrochloride<br>injection for intravenous or<br>intramuscular use | suspected to be caused by bacteria.<br>Indicated for the treatment of the following infections caused by susceptible strains of the designated<br>microorganisms:<br>• Moderate to severe pneumonia<br>• Empiric therapy for febrile neutropenic patients<br>• Uncomplicated and complicated urinary tract infections (including pyelonephritis)<br>• Uncomplicated skin and skin structure infections<br>• Complicated thra-abdominal infections (used in combination with metronidazole) in adults   | 120                               | 2 months  | N/A         | N/A                    | Y               | Ŷ                               |  | 8/5/2021             |

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 The HCPCS Code effective date represents the date the HCPCS code was established

• Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

| medicaid/med | HCPCS<br>Code | HCPCS Description                                    | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|--------------|---------------|--|----------------------------|-------------------------|------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Drugs        | J0694         | injection, cefoxitin sodium, 1<br>gram               | lg                         | 1/1/2000                | N/A        | cefoxitin for injection   | Indicated for the freatment of serious interctions caused by susceptible strains of the designated<br>microorganisms in the disease listed below.<br>• Lower respiratory tract infections: including pneumonia and lung abscess, caused by Streptococcus<br>pneumoniae, other streptococci (excluding enterococci, e.g., Enterococcus faecalis (formerly<br>streptococcus faecalis), Staphylococcus aureus (including penticilinase-producing strains), Escherichia<br>streptococcus faecalis), Staphylococcus aureus (including penticilinase-producing strains), Escherichia<br>coli, Ribesiela species, Haemophilus influenzae, and Bacteroides species.<br>Hintra-abdomiani Infections (auced by Escherichia coli, Ribesiela species, Proteus mirabilis, Morganella<br>morgani, Proteus vulgaris and Providencia species (including P. rettgeri).<br>• Intra-abdomiani Infections, including penticnitis and intra-abdominal abscess, caused by Escherichia coli,<br>Klebsiela species, Bacteroides species including Bacteroides fragilis, and Clostridium species.<br>• Gymecological infections, including entomettista public callulitis, and pelvic inflammatory disease auced<br>by Escherichia coli, Neisseria gonorrhoese (including penticilinase-producing strains), Bacteroides species<br>including 8. fragilis. Clostridium species, Peptococcus niger, Peptostreptococcus<br>species, and Streptococcus agalaciae. Cefontin, like cephalosporins, has no activity against Chlamydia<br>trachomatis. Therefore, when cefontin is used in the treatment of patients with pelvic inflammatory<br>disease and C. trachomatis is one of the suspected pathogens, appropriate anti-Chamydia coverage<br>should be added.<br>• Solin and skin structure infections: caused by Staphylococcus aureus (including pencillinase producing strains).<br>• Skin and skin structure infections: caused by Staphylococcus aureus (including pencillinase producing<br>strains). Staphylococcus (pathemidis, Streptococcus and guerical<br>infaciated for the prophylaxis of infection in patients undergoing uncontaminated gastrointestinal surg                       | 372                               | 3 months  | N/A         | N/A                    | Y               | Y                               |   | 9/27/2018             |
| Drugs        | J0695         | Injection, ceftolozane 50 mg<br>and tazobactam 25 mg | 75 mg                      | 1/1/2016                | Zerbaxa®   | ceftolozane and tazobactam<br>for injection, for intravenous<br>use | Indirial Hustanstonia, shakanias hustanstonia, secana a carsina -<br>Indicated in patients 13 years or older for the treatment of the following infections caused by designated<br>susceptible microorganisms:<br>• Complicated intra-abdominal infections (cIAI), used in combination with metronidazole.<br>• Complicated unray tract infections (cIAII), including pyelonephritis.<br>• Hospital-acquired Bacterial Pneumonia and Ventilator-associated Bacterial Pneumonia (HABP/VABP)<br>indicated in pediatric patients (birth to less than 18 years old) for the treatment of the following infections<br>caused by designated susceptible microorganisms:  | 1,680                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | cIAI and cUTI: N/A<br>HABP/VABF: 18 years of age<br>and older | 5/9/2022              |
|              |               |  |                            |                         |            |   | <ul> <li>Complicated Intra-abdominal Infections (cIAI), used in combination with metronidazole</li> <li>Complicated Urinary Tract Infections (cUTI), including pyelonephritis</li> <li>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Zerbaxa and other<br/>antibacterial drugs, Zerbaxa should be used only to treat or prevent infections that are proven or strongly<br/>suspected to be caused by bacteria.</li> </ul>   |                                   |   |             |                        |                 |                                 |   |                       |
| Drugs        | J0696         | Injection, ceftriaxone sodium<br>per 250 mg          | 250 mg                     | 1/1/2000                | Rocephin*  | ceftriaxone sodium injection  | Indicated for the treatment of the following impections when caused by susceptible organisms:<br>- Lower Regristrony Tract Infections: Caused by Streptococcus perumoniae, Staphylococcus aureus,<br>Haemophilus influenzae, Haemophilus parainfluenzae, Klebsiella pneumoniae, Escherichia coli,<br>Enterobacter acogenes, Proteus minibilis or Serraria marcescens.<br>- Acute Bacterial Otitis Media: Caused by Streptococcus perumoniae, Haemophilus influenzae (including<br>beta-latamase producing strains) or Moraella catanthalis (including beta-latamase producing strains).<br>- Skin and Sin Structure Infections. Caused by Staphylococcus aureus, Staphylococcus apreus,<br>Streptococcus progenes, Virdians group streptococc). Escherichia coli, Enterobacter cloace, Klebsiella<br>oxytca, Klebsiella pneumoniae, Proteus minibilis, Morganella morganii. Pseudomans as eruginosa,<br>Serratia marcescens, Acinetobacter calcoaceticus, Bacteroides fragilis or Peptostreptococcus species.<br>- Unionay Tract Infections: Caused by Staphylococcus aureus, Staphylococcus aureus,<br>Serratia marcescens, Acinetobacter calcoaceticus, Bacteroides fragilis or Peptostreptococcus<br>protein in Klebsiella pneumoniae.<br>- Uncomplicated Gonorhea (cervical/urethral and rectal): Caused by Neisseria goomrhoeae, including<br>both pencillinase- and nonpenicillinase-producing strains, and pharyngeal gonorhea caused by<br>nonpencillinase- producing strains of Neisseria gonorrhoeae. Cefinaxone sodium, like other<br>cephalosporins, has no activity against Chlamydia trachomatis. Therefore, when cephalosporins are used<br>in the treatment of patients with pelvic inflammatory disease and Chlamydia trachomatis is one of the<br>suspected pathogens, appropriate antichystaphylococcus aureus, Streptococcus pneumoniae, Escherichia coli,<br>Haemophilus influenzae or Klebsiella pneumoniae<br>- Intra-abdomila Infections: Caused by Staphylococcus aureus, Streptococcus pneumoniae, Escherichia<br>coli, Proteus mirabilis, Klebsiella pneumoniae or Enterobacter species.<br>- Intra-abdomila Infections: Caused by Staphylococc | 496                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | v                               | See package insert for specific neonate contraindication.     | 10/4/2018             |

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| Category | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name                                       | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler | Comments   | Last Modified<br>Date |
|----------|---------------|---|----------------------------|-------------------------|------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------|--|-----------------------|
| Drugs    | J0697         | Injection, sterile cefuroxime<br>sodium, per 750 mg   | 750 mg                     | 1/1/2000                | Zinacef*   | cefuroxime for injection                           | Indicated for the treatment of patients with infections caused by susceptible strains of the designated<br>organisms in the following diseases:<br>Lower Respiratory Tract Infections: including pneumonia, caused by Streptococcus pneumoniae,<br>Haemophilus influenzae (including ampicillin-resistant strains), Klebsiella spp., Staphylococcus aureus<br>(pericillinase- and non-pericillinase-producing strains), Streptococcus pyogenes, and Escherichia coli.<br>Urinary Tract Infections: caused by Staphylococcus aureus (penicillinase- and non-penicillinase producing strains), Streptococcus pyogenes, Scherichia coli, Messiella spp., and Enterobacter spp.<br>• Sein and Skin-Structure Infections: caused by Staphylococcus aureus (penicillinase- and non-penicillinase), superioducing strains), Streptococcus pyogenes, Scherichia coli, Messiella spp., and Enterobacter spp.<br>• Septicemais: caused by Staphylococcus aureus (penicillinase- and non-penicillinase-<br>strains), and Klebsiella spp.<br>• Meningti: caused by Staphylococcus aureus (penicillinase- and non-penicillinase-producing strains),<br>strains), Neisseria meningtiklis, and Staphylococcus aureus (penicillinase- and non-penicillinase-producing strains),<br>• Gonorrhoeae: Uncomplicated and disseminated gonococcal infections due to Neisseria gonorrhoeae (penicillinase- and non-penicillinase-producing strains), in oth males and females.<br>• Bone and Joint Infections: caused by Staphylococcus aureus (penicillinase- and non-penicillinase-<br>producing strains).  | 372                               | 3 months  | N/A         | N/A                    | Y               | Required            |  | 10/4/2018             |
| Drugs    | 90698         | Cefotaxime sodium, per gram   | lg                         | 1/1/2000                | Claforan®  | cefotaxime for injection                           | Indicated for the treatment of patients with serious intections caused by susceptible strains of the<br>designated microorganisms in the diseases listed below.<br>Lower registrony tract infections: including penumonia, caused by Streptococcus pneumoniae (formerly<br>Diplococcus pneumoniae). Streptococcus progenes" (Group A streptococci) and other streptococci<br>(excluding entropy tract infections: Staphylococcus areaus (geneillinase and non-<br>penidlinase producing). Escherichia coll, Klebsiella species, Haemophilus influenzae (including ampicillin<br>resistant strains). Haemophilus parinfluenzae, Proteus mirabilis, Serata macrescens". Enterobacter<br>species, indole positive Proteus and Pseudomonas species (including P. aeruginosa).<br>Cenitourinary infections: Urinary tract infections caused by Enterococcus species, Staphylococcus<br>apidemilis, Staphylococcus aureus", (pencillinase and non-pencillinase producing). Ctrobacter species,<br>Interobacter species, Escherichia coll, Klebsiella species, Proteus mirabilis, Protesu valgeris", Providencia<br>stuarti, Morganella morgani", Providencia rettgeri ", serratia marcescens", and Pseudomonas species<br>(including P. aeruginosa). Also, uncomplicated gonorrhae, endometritis and pelva delluitis caused by<br>Staphylococcus egidermidis, Streptococcus species, Enterobacter species, fueltorbacter species,<br>Indevisiel species, escherichia coli, iProtesu mirabilis, Bacteroides species, (Induding Bacteroides traglis'),<br>Costridium species, and anarobic coccu (Induding Pedotstreptococcus species) and Pelvacoccus species and Cristophilosporiane aureu el in the trastment of patients<br>with pelvic inflammatory disease and C. trachomatis is one of the suspected pathogens, appropriate anti-<br>chiamydial coverage should be added. | 372                               | N/A   | N/A         | N/A                    | ¥               | ¥                   |  | 5/20/2019             |
| Drugs    | J0699         | Injection, cefiderocol, 10 mg   | 10 mg                      | 10/1/2021               | Fetroja®   | cefiderocol for injection, for<br>intravenous use  | Indicated in patients 18 years of age or older for the treatment of complicated urinary tract infections<br>(CUTT), including pyelonephritis caused by the following susceptible Gram-negative microorganisms:<br>Escherichia culi, Klebsiella pneumoniae, Proteus mirabilis, Pseudomonas aeruginosa and Enterobacter<br>cloacae complex.  | 11,200                            | 18 years  | N/A         | N/A                    | Y               | Ŷ                   |  | 9/29/2021             |
| Drugs    | J0701         | Injection, cefepime<br>hydrochloride (baxter), not<br>therapeutically equivalent to<br>maxipime, 500 mg | 500 mg                     | 1/1/2023                | N/A        | cefepime injection for<br>intravenous use (Baxter) | suspected to be caused by bacteria.<br>Indicated in the treatment of the following infections caused by susceptible isolates of the designated<br>microorganisms: pneumonia; empiric therapy for febrile neutropenic patients; uncomplicated and<br>complicated uninary tract infections; uncomplicated skin and skin structure infections; and complicated<br>intra-abdominal infections (used in combination with metronidazole).<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Cefepime<br>injection and other antibacterial drugs, Cefepime injection should be used only to treat or prevent<br>infections that are proven or strongly suspected to be caused by bacteria.  | 120                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                   | Indication specific age<br>restrictions:<br>• Complicated intra-abdominal<br>infections: 17 years of age and<br>older<br>• All other indications: 2<br>months of age and older | 12/19/2022            |

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| Category | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name              | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|----------|---------------|--|----------------------------|-------------------------|-------------------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Drugs    | J0702         | Injection, betamethasone<br>acetate 3 mg and<br>betamethasone sodium<br>phosphate 3 mg                   | 1 mL                       | 1/1/2000                | Celestone*<br>Soluspan* | betamethasone sodium<br>phosphate and<br>betamethasone acetate<br>injectable suspension | When oral therapy is not reasible, the intramuscular use or cleastone Soluspan is indicated as follows:<br>- Allergic States: Control of severe or incapacitating allergic conditions intractable to adequate trails of<br>conventional treatment in asthma, atopic dermatitis, contact dermatitis, drug hypersensitivity reactions,<br>perennial or sessional allergic infinitions; strendsulon reactions.<br>• Dermatologic Diseases: Sullous dermatitis herpetformis, scholative erythroderma, mycosis fungoides,<br>pemphigus, severe erythema multilitis seum sichness, transfusion reactions.<br>• Endocrine Disorders: Congenital adrenal hyperplasia, hypercalcemia associated with cancer,<br>nonsuppurative thyroiditis. Hydrocortisone or cortisone is the drug of choice in primary or secondary<br>appinguis, severe erythema multilitie analogs may be used in conjunction with mineraloutchoids where<br>appicable; in infancy mineralocorticoid supplementation is of particular importance.<br>• Gastrointestinal Diseases: To the the patient over a critical period of the disease in regional enteritis and<br>ulcerative colitis.<br>• Hematologic Disorders: Acquired (autoimmune) hemolytic anemia, Diamond-Blackfan anemia, pure red<br>• Alleselases: For pallater unavolgic or myocardial involvement, tuberculous chemotherapy.<br>• Neoplastic Diseases: For pallater unavolgic or myocardial involvement, tuberculous chemotherapy.<br>• Neoplastic Diseases: For pallater unavolgic or myocardial involvement, tuberculous chemotherapy.<br>• Neoplastic Diseases: For pallater company and the appropriate antituberculous chemotherapy.<br>• Neoplastic Diseases: For pallater controlowy.<br>• Ophthalmic Diseases: To induce duresis or remission of proteinuria in idiopathic nephrotic syndrome or that<br>due to lous erythematous.<br>• Reprivatory Diseases: Sympathetic ophthalmia, temporal arteritis, uveitis and ocular inflammatory<br>conditions unresponsive to topical corticosterodis.<br>• Reprivatory Diseases: Sympathetic ophthalmia, temporal arteritis, uveitis and ocular inflammatory<br>conditions unresponsive to           | 155                               | N/A   | N/A         | N/A                    | ¥               | ¥                               |   | 9/25/2018             |
| Drugs    | J0703         | Injection, cefepime<br>hydrochloride (b braun), not<br>therapeutically equivalent to<br>maxipime, 500 mg | 500 mg                     | 1/1/2023                | N/A                     | cefepime for injection and<br>dextrose injection for<br>intravenous use (B. Braun)      | Indicated in the treatment of the following infections caused by susceptible strains of the designated<br>microorganisms:<br>• Pneumonia<br>• Empiric therapy for febrile neutropenic patients<br>• Uncomplicated and complicated urinary tract infections<br>• Uncomplicated skin and skin structure infections<br>• Complicated intra-abdominal infections (used in combination with metronidazole)<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Cefepime for<br>injection and Dextrose injection and other antibacterial drugs, Cefepime for injection and Dextrose<br>Injection should be used only to treat or prevent infections that are proven or strongly suspected to be<br>caused by bacteria.  | 120                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Ŷ               | ¥                               | Indication-specific age<br>restrictions:<br>Complicated intra-abdomial<br>infections: 17 years of age and<br>older<br>- All other indications: 2<br>months of age and older | 12/12/2022            |
| Drugs    | J0712         | Injection, ceftaroline fosamil,<br>10 mg   | 10 mg                      | 1/1/2012                | Teflaro®                | ceftaroline fosamil for<br>injection, for intravenous use                               | Indicated for the treatment of the following infection caused by designated susceptible bacteria:<br>• Community-acquired bacterial pneumonia (CABP) in adult and pediatric patients 2 months of age and<br>other   | 1,680                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific:<br>CABP: 2 months of age and<br>older<br>ABSSSI: 34 weeks gestational<br>age and 12 days postnatal age<br>and older                                    | 10/28/2019            |
| Drugs    | J0713         | Injection, ceftazidime, per 500<br>mg  | per 500 mg                 | 1/1/2000                | Tazicef®                | ceftazidime for injection, for<br>intravenous or intramuscular<br>use                   | Indicated for the treatment of patients with infections caused by susceptible strains of the designated<br>organisms in the following diseases:<br>- Lower Regarizatory Tract Infections: including pneumonia, caused by Pseudomonas aeruginosa and other<br>Pseudomonas spp: Haemophilus influenzae, including ampcillin-resistant strains; Klebsiella spp;<br>Enterobacter spp: Forteus minibilis; Scherkich acity, Serratia spp; Cirbobacter spp; Streptococcus<br>pneumoniae; and Stan-Phylococcus aureus (methicillin-susceptible strains).<br>• Skin and Skin-Structure Infections: caused by Pseudomonas areuginosa; Klebsiella spp; Escherichia coli;<br>Proteus spp. Indiuding Proteus minibilis and indioe-positive Proteus; Entrobacter spp; Serratia spp;<br>Staphylococcus aureus (methicillin-susceptible strains); and Streptococcus progenes (group A beta-<br>hemolyici streptococci).<br>• Urinary Tract Infections: both complicated and uncomplicated, caused by Pseudomonas aeruginosa;<br>Enterobacter spp; Proteus spp, including Proteus mirabilis and indiole-positive Proteus; Klebsiella spp; and<br>Escherichia coli.<br>• Bacterial Sept; Proteus spp, Jroteus pp, Streptococcus aureus<br>(methicillinsusceptible strains).<br>• Soma and Joint Infections: caused by Pseudomonas aeruginosa, Klebsiella spp, Haeroobatter spp, and<br>Staphylococcus aureus (methicillin-susceptible strains).<br>• Gyneologic Intelfoctions: including endometritis, pevic cellulitis, and other infections of the female genital<br>tract caused by Escherichia coli.<br>• Intra-abdomisma and Bacteriol Septies spp, and<br>Staphylococcus aureus (methicillin-susceptible strains).<br>• Intra-abdomisma and Bacteriol Septies spi, and Staphylococcus aureus<br>(Methicillinsus aerus).<br>• Intra-abdomisma and Bacterioles spp. (amay strains of Bacterioles fraglis are resistant).<br>• Central Infections: including peritonitis caused by Escherichia coli, Klebsiella spp, and<br>Staphylococcus aureus (methicillin-susceptible strains) and polymicrobial infections caused by as<br>spatiant of the spatian and Bacterio septicant and states of<br>men | 372                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 5/21/2019             |
| Drugs    | J0714         | Injection, ceftazidime and avibactam, 0.5 g/0.125 g  | 0.625 g                    | 1/1/2016                | Avycaz*                 | ceftazidime and avibactam<br>for injection, for intravenous<br>use                      | Indicated for the treatment of the following infections caused by designated susceptible Gram-negative<br>microorganisms in adult and pediatric patients (at least 31 weeks gestational age):<br>• Complicated Intra-abdominal infections (clAI), used in combination with metronidazole<br>• Complicated Uninary Tract Infections (clAI), including Pyelonephitis<br>• Hospital-acquired Bacterial Pneumonia and Ventilator-associated Bacterial Pneumonia (HABP/VABP)   | 168                               | 31 weeks gestational age                                  | N/A         | N/A                    | Ŷ               | Y                               |   | 2/27/2024             |

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|--------------|---------------|--|----------------------------|-------------------------|-----------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name            | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Biologicals  | J0716         | Injection, centruroides<br>immune f(ab)2, up to 120<br>milligrams  | up to 120 mg (1 vial)      | 1/1/2013                | Anascorp®             | centruroides (scorpion)<br>immune F(ab') <sup>2</sup> (equine)<br>injection lyophilized for<br>solution, for intravenous use<br>only                        | Antivenom indicated for treatment of clinical signs of scorpion envenomation.   | N/A                               | N/A         | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Biologicals  | J0717         | Injection, certolizumab pegol,<br>1 mg   | 1 mg                       | 1/1/2014                | Cimzia*               | certolizumab pegol for<br>injection, for subcutaneous<br>use  | Indicated for:<br>* Reducing signs and symptoms of Crohn's disease and maintaining clinical response in adult patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.<br>* Treatment of adult satients with active positiat carthritis.<br>* Treatment of adult satients with active positiat carthritis.<br>* Treatment of adults with active positiat carthritis.<br>* Treatment of adults with active ankylosing spondylitis.<br>* Treatment of adults with moderate-to-severe plaque psoriasis who are candidates for systemic therapy<br>or phototherapy.<br>* Treatment of adults with active non-radiographic axial spondyloarthritis who have objective signs of<br>Inflammation.   | 1,200                             | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 5/1/2019              |
| Drugs        | J0720         | Injection, chloramphenicol<br>sodium succinate, up to 1 g  | up to 1 g                  | 1/1/2000                | N/A                   | chloramphenicol sodium<br>succinate for injection, for<br>intravenous administration  | **Chioramphenicol must be used only in those serious infections for which less potentially dangerous<br>drugs are ineffective or contraindicated. (See package insert for recommendations and warnings<br>associated with chioramphenicol.)<br>Indicated for:<br>Acute infections caused by Salmonella typhi. In treatment of typhoid fever some authorities<br>recommend that chioramphenicol be administered at<br>therapeutic levels for 8 to 10 days after the patient has become afebrile to lessen the possibility of<br>relapse. It is not recommended for the routine treatment of the typhoid carrier state.<br>Serious infections caused by susceptible strains in accordance with the concepts expressed in the<br>package insert:<br>- Salmonella species<br>- H, influenzae, specifically meningeal infections<br>- Rickettsia<br>- Lymphogranuloma-psiltacosis group<br>- Various gram-negative bacteria causing bacteremia, meningitis or other serious gram-negative<br>infections.<br>- Other susceptible organisms which have been demonstrated to be resistant to all other appropriate<br>antimicrobial agents.<br>- Cystic fibrosis regimens | 217                               | N/A         | N/A         | N/A                    | Y               | Y                               |  | 10/4/2018             |
| Biologicals  | J0725         | Injection, chorionic<br>gonadotropin, per 1,000 USP<br>units   | 1,000 USP units            | 1/1/2000                | Novarel®,<br>Pregnyl® | chorionic gonadotropin for<br>injection   | Indicated for:<br>• Prepubertal cryptorchidism not due to anatomic obstruction. In general, HCG is thought to induce<br>testicular descent in situations when descent would have occurred at puberty. HCG thus may help to<br>predict whether or not orchiopeya will be needed in the future. Although, in some cases, descent following<br>HCG administration is permanent, in most cases the response is temporary. Therapy is usually instituted<br>between the ages of 4 and 9.<br>• Selected cases of hypogonadotropic hypogonadism (hypogonadism secondary to a pituitary deficiency)<br>in males.<br>• Induction of ovulation and pregnancy in the anovulatory, infertile woman in whom the cause of<br>anovulation is secondary and not due to primary ovarian failure, and who has been appropriately<br>pretreated with human menotropins.  | 60                                | 4 years     | N/A         | N/A                    | Y               | Y                               |  | 6/19/2023             |
| Drugs        | J0735         | Injection, clonidine<br>hydrochloride, 1 mg  | 1 mg                       | 1/1/2000                | Duracion®             | clonidine hydrochloride<br>injection solution   | Indicated in combination with opiates for the treatment of severe pain in cancer patients that is not<br>adequately relieved by opioid analgesics alone. Epidural clonidine is more likely to be effective in patients<br>with neuropathic pain than somatic or visceral pain.  | See Comments                      | N/A         | N/A         | N/A                    | Y               | Y                               | Maximum daily and monthly<br>doses are individualized and<br>patient specific. | 10/4/2018             |
| Drugs        | J0739         | Injection, cabotegravir, 1 mg,<br>FDA approved prescription,<br>only for use as HIV pre-<br>exposure prophylaxis (not for<br>use as treatment for HIV) | 1 mg                       | 1/1/2000                | Apretude              | cabotegravir extended-<br>release injectable suspension,<br>for intramuscular use   | Indicated in at-risk adults and adolescents weighing at least 35 kg for PrEP to reduce the risk of sexually<br>acquired HIV-1 infection.  | 1,200                             | 12 years    | N/A         | N/A                    | Y               | Y                               |  | 1/4/2024              |
| Drugs        | J0740         | Injection, cidofovir, 375 mg   | 375 mg                     | 1/1/2000                | Vistide*              | cidofovir injection for<br>intravenous infusion   | Indicated for the treatment of cytomegalovirus (CMV) retinitis in patients with acquired<br>immunodeficiency syndrome (AIDS).   | 6                                 | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 9/27/2018             |
| Drugs        | J0741         | Injection, cabotegravir and rilpivirine, 2mg/3mg   | 2 mg/3 mg                  | 10/1/2021               | Cabenuva™             | cabotegravir extended-<br>release injectable suspension;<br>rilpivirine extended-release<br>injectable suspension, co-<br>packaged for intramuscular<br>use | Indicated as a complete regimen for the treatment of HIV-1 infection in adults and adolescents 12 years of<br>age and older and weighing at least 35 kg to replace the current antiretroviral regimen in those who are<br>virologically suppressed (HIV-1 RNA less than 50 copies per mI) on a stable antiretroviral regimen with no<br>history of treatment failure and with no known or suspected resistance to either cabotegravir or<br>ripivirine.   | 600                               | 12 years    | N/A         | N/A                    | Y               | Y                               |  | 4/21/2022             |
| Drugs        | J0742         | Injection, imipenem 4 mg,<br>cilastatin 4 mg and relebactam<br>2 mg  | n 10 mg                    | 7/1/2020                | Recarbrio™            | imipenem, cilastatin, and<br>relebactam for injection, for<br>intravenous use   | Indicated in patients 18 years of age and older who have limited or no alternative treatment options, for<br>the treatment of the following infections caused by susceptible gram-negative bacteria:<br>• Complicated urinary tract infections, including pyelonephritis (UTI)<br>• Complicated urinary tract infections (IAI)<br>• Hospital-acquired bacterial pneumonia and ventilator-associated bacterial pneumonia (HABP/VABP)<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Recarbrio and<br>other antibacterial drugs, Reacribrio should be used only to treat or prevent infections that are proven or<br>strongly suspected to be caused by bacteria.  | 7,000                             | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 7/28/2020             |

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| Image: Property in the propert        | ategory    | HCPCS<br>Code | HCPCS Description              | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name    | Generic Name                           | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
|---|------------|---------------|--------------------------------|----------------------------|-------------------------|---------------|--|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| u       kin   | Drugs      | J0743         |                                | 250 mg                     | 1/1/2000                | Primaxin®     |  | Lower respiratory tract infections Udmay tract infections Intra-abdominal infections Gynecologic infections Bacterial septicemia Bone and joint infections Solin and skin structure infections Solin and skin structure infections Endocarditis Uminitations of Use: Not indicated in patients with meningitis because safety and efficacy have not been established. Not recommended in patientic patients with CNS infections because of the risk of seizures.  | 496                               | N/A         | N/A         | N/A                    | Y               | ¥                               |          | 9/27/2018             |
| One       Main       Main      <  | Drugs      | J0744         |                                | 200 mg                     | 1/1/2002                | Cipro IV®     |  | bacteria and in pediatric patients where indicated:<br>Skin and skin structure infections<br>Bone and joint infections<br>Complicated intra-abdominal infections<br>Nosocomial penemonia<br>Empirical therapy for febrile neutropenic patients<br>Inhalational antrax post-exposure in adult and pediatric patients<br>Plague in adult and pediatric patients<br>Chronic bacterial prostatis<br>Lower respiratory tract infections<br>- Acute exacerbation of chronic bronchitis<br>- Urinary tract infections:<br>- Complicated UT and pyelonephrits in pediatric patients | 186                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 4/9/2019              |
| v constraint   | Drugs      | J0770         |                                | up to 150 mg               | 1/1/2000                | Coly-Mycin® M | colistimethate for injection           | bacilli. Particularly indicated when the infection is caused by sensitive strains of P. aeruginosa. Clinically<br>effective in treatment of infections due to the following gram-negative organisms: Enterobacter   |                                   | N/A         | N/A         | N/A                    | Ŷ               | Y                               |          | 6/4/2019              |
| Opp         Opp         Oppo         O   | iologicals | J0775         | clostridium histolyticum, 0.01 | 0.01 mg                    | 1/1/2011                | Xiaflex®      |  | <ul> <li>Treatment of adult men with Peyronie's disease with a palpable plaque and curvature deformity of at<br/>least 30 degrees at the start of therapy.</li> </ul>   | 360                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/6/2019              |
| Mode  | Drugs      | J0780         |                                | up to 10 mg                | 1/1/2000                | N/A           |  | has not been shown effective in the management of behavioral complications in patients with mental  | 124                               | 2 years     | N/A         | N/A                    | Y               | Y                               |          | 8/24/2018             |
| Data       Indication contractorphic (status<br>gs), up to 40 units       10/1/2020       Actus*ed       Actus*ed       Indicate of the following formation is addesses: thematic, calling, dematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic, alling, calmades; teter,<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all<br>who is used anti-out is addesses: thematodic alling; teter, teter, all who is used anti-out is addesses of the same of t | iologicals | J0791         |                                | 5 mg                       | 7/1/2020                | Adakveo®      |  |   | 280                               | 16 years    | N/A         | N/A                    | Y               | Y                               |          | 6/17/2020             |
| Drugs         N802         Injection, corticotropin (an),<br>up to 40 units         up to 40 units         10/1/2023         Purified         repository of short-term administration (to tide the patient over a<br>latere spoke) of short-term administration (to tide the patient over a<br>latere spoke)  | Drugs      | J0801         |                                | up to 40 units             | 10/1/2023               | Acthar® Gel   | injection, gel for<br>intramuscular or | age.<br>• Indicated for the treatment of exacerbations of multiple sclerosis in adults.<br>• May be used for the following disorders and diseases: rheumatic, collagen, dermatologic, allergic states,<br>ophthalmic, respiratory, and edematous state.   | 63                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 7/29/2024             |
| Injection, cosyntropin, 0.25  | Drugs      | J0802         |                                | up to 40 units             | 10/1/2023               |               |  |   | 63                                | N/A         | N/A         | N/A                    | Y               | v                               |          | 9/28/2023             |
| Drugs J0834 mileculor, do yn colin, d. 2 0.25 mg 1/1/2010 Cortrosyn <sup>w</sup> drianosin method no de as a diagraphica gen in the scening of patients pestiline to nave autonocolical 3 N/A N/A N/A Y Y 2/4/20:   | Drugs      | J0834         | Injection, cosyntropin, 0.25   | 0.25 mg                    | 1/1/2010                | Cortrosyn™    | cosyntropin injection for              | Ontic neuritic     Intended for use as a diagnostic agent in the screening of patients presumed to have adrenocortical  | 3                                 | N/A         | N/A         | N/A                    | Y               | Y                               |          | 2/4/2019              |

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|-------------|---------------|---|----------------------------|-------------------------|------------|--|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Biologicals | J0840         | Injection, crotalidae polyvalent<br>immune fab (Ovine), up to 1<br>gram   | up to 1 g (1 vial)         | 1/1/2012                | CroFab®    | crotalidae polyvalent immun<br>fab (ovine) lyophilized<br>powder for solution for<br>intravenous injection     | e Indicated for the management of adult and pediatric patients with North American crotalid<br>envenomation. The term crotalid is used to describe the Crotalinae subfamily (formerly known as<br>Crotalidae) of venomous snakes which includes rattlesnakes, copperheads and cottonmouths/water<br>mocicasins.  | N/A                               | N/A         | N/a         | N/A                    | Y               | N                               |          | 1/4/2019              |
| Biologicals | J0841         | Injection, crotalidae immune<br>f(ab')2 (equine), 120 mg  | 120 mg                     | 1/1/2019                | Anavip®    | crotalidae immune f(ab')2<br>(equine), lyophilized powder<br>for solution for injection for<br>intravenous use | Indicated for the management of adult and pediatric patients with North American rattlesnake   | N/A                               | N/A         | N/A         | N/A                    | Ŷ               | Y                               |          | 12/28/2018            |
| Drugs       | J0872         | Injection, daptomycin (xellia),<br>unrefrigerated, not<br>therapeutically equivalent to<br>j0878 or j0873, 1 mg | 1 mg                       | 7/1/2024                | N/A        | daptomycin for injection, fo<br>intravenous use (Xella) –<br>unerfigment Storage<br>permitted                  | Daptomycin for injection is indicated for the treatment of:<br>• Complicated skin and skin structure infections (cSSSI) in adult and pediatric patients (1 to 17 years of age) and,<br>• Staphylococcus aureus bloodstream infections (bacteremia), in adult patients including those with right-<br>sided infective endocardits,<br>• Staphylococcus aureus bloodstream infections (bacteremia) in pediatric patients (1 to 17 years of age).<br>Limitations of Use:<br>• Daptomycin for injection is not indicated for the treatment of pneumonia.<br>• Daptomycin for injection is not indicated for the treatment of left-sided infective endocarditis due to S.<br>aureus.<br>• Daptomycin for injection is not recommended in pediatric patients younger than one year of age due to<br>the risk of potential effects on muscular, neuromuscular, and/or nervous systems (either peripheral<br>and/or central) observed in neonatal dogs.<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Daptomycin for<br>injection and other antibacterial drugs, Daptomycin for injection should be used to treat or prevent<br>infections that are proven or strongly suspected to be caused by bacteria.   | 31,000                            | 1 year      | N/A         | N/A                    | Ŷ               | ¥                               |          | 6/24/2024             |
| Drugs       | J0873         | Injection, daptomycin (xellia)<br>not therapeutically equivalent<br>to j0878, 1 mg                              | 1 mg                       | 1/1/2024                | N/A        | daptomycin for injection, fo<br>intravenous use (Xellia) -<br>refrigerated storage required                    | Daptomycin for injection is not indicated for the treatment of pheumonia.  | 31,000                            | 1 year      | N/A         | N/A                    | Y               | Y                               |          | 6/25/2024             |
| Drugs       | J0874         | Injection, daptomycin (baxter),<br>not therapeutically equivalent<br>to j0878, 1 mg                             | 1 mg                       | 10/1/2023               | N/A        | daptomycin in sodium<br>chloride injection, for<br>intravenous use (Baxter)                                    | Indicated for the treatment of:<br>• Complicated skin and skin structure infections (c555) in adult and pediatric patients (1 to 17 years of<br>age) for whom appropriate dosing can be achieved and,<br>• Staphylococcus aurcus biodostream infections (bacteremia), in adult patients for whom appropriate<br>dosing can be achieved, including those with right-sided infective endocarditis,<br>• Staphylococcus aurcus biodostream infections (bacteremia) in pediatric patients (1 to 17 years of age)<br>for whom appropriate dosing can be achieved.<br>Limitations of Use:<br>• Daptomycin in Sodium Chloride injection is not indicated for the treatment of left-sided infective<br>endocarditis due to <i>S. aurcus</i> .<br>• Daptomycin in Sodium Chloride injection is not indicated for the treatment of left-sided infective<br>endocarditis due to <i>S. aurcus</i> .<br>• Daptomycin in Sodium Chloride injection is not indicated for the treatment of left-sided infective<br>endocarditis due to <i>S. aurcus</i> .<br>• Daptomycin in Sodium Chloride injection is not recommended in pediatric patients younger than one<br>year of age due to the risk of potential effects on muscular, neuromuscular, and/or nervous systems<br>(either peripheral and/or central) observed in neonatal dogs.<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Daptomycin in<br>Sodium Chloride injection and other antibacterial drugs, Daptomycin in Sodium Chloride Injections should<br>be used to treat or prevent infections that are proven or strongly suspected to be caused by bacteria. | 31,000                            | 1 year      | N/A         | N/A                    | Ŷ               | Y                               |          | 9/28/2023             |
| Drugs       | J0875         | Injection, dalbavancin, 5 mg  | 5 mg                       | 1/1/2016                | Dalvance®  | dalbavancin for injection, fo<br>intravenous use   | Indicated for the treatment of:<br>- adult patients with acute bacterial skin and skin structure infections (ABSSSI) caused by designated<br>susceptible strains of Gram-positive microorganisms.<br>- pediatric patients with acute bacterial skin and skin structure infections (ABSSSI) caused by designated<br>susceptible strains of Gram-positive microorganisms.  | 300                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 8/25/2021             |

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|-------------|---------------|--|----------------------------|-------------------------|------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
|             |               |  |                            |                         |            |   | Indicated for the treatment of:<br>• Complicated skin and skin structure infections (cSSSI) in adult patients<br>• Staphylococcus aureus bloodstream infections (bacteremia), in adult patients including those with right-<br>sided infective endocarditis<br>Limitations of Use:   |                                   |   |             |                        |                 |                                 |   |                       |
| Drugs       | J0877         | Injection, daptomycin<br>(hospira), not therapeutically<br>equivalent to j0878, 1 mg | 1 mg                       | 1/1/2023                | N/A        | daptomycin for injection, for<br>intravenous use (Hospira)                                  | This Daptomycin for injection product is not approved for use in pediatic patients.     Daptomycin for injection is not indicated for the treatment of pneumonia.     Daptomycin for injection is not indicated for the treatment of left-sided infective endocarditis due to 5.     aureus.     Daptomycin for Injection is not recommended in pediatric patients younger than one year of age due to     the risk of potential effects on muscular, neuromuscular, and/or nervous systems (either peripheral     and/or central) observed in neoratial dogs.   | 27,900                            | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 6/25/2024             |
|             |               |  |                            |                         |            |   | To reduce the development of drug-resistant bacteria and maintain the effectiveness of Daptomycin for<br>Injection and other antibacterial drugs, Daptomycin for Injection should be used to treat or prevent<br>infections that are proven or strongly suspected to be caused by bacteria.  |                                   |   |             |                        |                 |                                 |   |                       |
| Drugs       | J0878         | Injection, daptomycin, 1 mg  | 1 mg                       | 1/1/2005                | Cubicin®   | daptomycin injection, for<br>intravenous use  | Indicated for the treatment of:<br>- Complicated skin and skin structure infections (cSSSI) in adult and pediatric patients (1 to 17 years of<br>age).<br>- Staphylococcus aureus bioodstream infections (bacteremia), in adult patients including those with right-<br>sided infective endocardits.<br>- Indicated for the treatment of Staphylococcus aureus bioodstream infections (bacteremia) in pediatric<br>patients (1 to 17 years of age).<br>Limitations of Use:<br>- Cubicin is not indicated for the treatment of pneumonia.<br>- Cubicin is not indicated for the treatment of left-sided infective endocarditis due to 5. aureus.<br>- Cubicin is not indicated for the treatment of left-sided infective endocarditis due to the risk of  | 31,000                            | 1 year  | N/A         | N/A                    | Y               | Y                               |   | 6/25/2024             |
|             |               |  |                            |                         |            |   | potential effects on muscular, neuromuscular, and/or nervous systems (either peripheral and/or central)<br>observed in neonatal dogs.<br>Indicated for the treatment of moderate-to-severe pruritus associated with chronic kidney disease (CKD-   |                                   |   |             |                        |                 |                                 |   |                       |
| Drugs       | J0879         | Injection, difelikefalin, 0.1<br>microgram, (for esrd on<br>dialysis)                | 0.1 mcg                    | 4/1/2002                | Korsuva™   | difelikefalin injection, for<br>intravenous use   | aP) in adults undergoing hemodialysis (HD).<br>Limitation of Use: Korsuva has not been studied in patients on peritoneal dialysis and is not recommended<br>for use in this population.  | 19,500                            | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 4/21/2022             |
| Biologicals | J0881         | Injection, darbepoetin alfa, 1<br>microgram (non-ESRD use)                           | 1 mcg                      | 1/1/2006                | Aranesp®   | darbepoetin alfa injection, foi<br>intravenous of subcutaneous<br>use (non-ESR0 use)        | Indicated for the treatment of anemia due to:<br>• Chronic Kidney Disease (CKD) in patients on dialysis and patient not on dialysis.<br>• The effects of concomitant myelosuppressive chemotherapy, and upon initiation, there is a minimum of<br>two additional months of planned chemotherapy.<br>Umitations of Use: Aranesp has not been shown to improve quality of life, fatigue, or patient well-being.  | 1,575                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | ¥                               | Indication specific age<br>restrictions:<br>• CRD: None<br>• Cancer: 18 years of age and<br>older | 4/10/2019             |
| Biologicals | J0882         | Injection, darbepoetin alfa, 1<br>microgram (for ESRD on<br>dialysis)                | 1 mcg                      | 1/1/2006                | Aranesp®   | darbepoetin alfa injection, fo<br>intravenous or subcutaneous<br>use (ESRD use on dialysis) | Indicated for the treatment of anemia due to:<br>Chronic Kidney Disease (CRD) in patients on dialysis and patients not on dialysis.<br>• The effects of concomitant myelosuppressive chemotherapy, and upon initiation, there is a minimum of<br>two additional months of planned<br>chemotherapy.<br>Limitations of Use: Aranesp has not been shown to improve quality of life, fatigue, or patient well-being.<br>Aranesp is not indicated for use:<br>In patients with cancer receiving hormonal agents, biologic products, or radiotherapy, unless also<br>receiving concomitant myelosuppressive<br>chemotherapy.<br>In patients with cancer receiving myelosuppressive chemotherapy when the anticipated outcome is<br>cure.<br>In patients with cancer receiving myelosuppressive chemotherapy when the anticipated outcome is<br>cure.<br>In patients with cancer receiving myelosuppressive chemotherapy in whom the anemia can be managed<br>by transfusion. | 315                               | N/A   | N/A         | N/A                    | Y               | ¥                               |   | 4/10/2019             |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
 The HCPCS Code effective date represents the date the HCPCS code was established

• Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

| Category    | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name        | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modifie<br>Date |
|-------------|---------------|---|----------------------------|-------------------------|-------------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|----------------------|
| Biologicals | J0885         | Injection, epoetin alfa, (for<br>non-ESRD use), 1000 units                              | 1,000 units                | 1/1/2006                | Epogen®, Procrit® | epoetin alfa for injection, for<br>intravenous or subcutaneous<br>use (for non ESRD use)                                 |   | 630                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• CKD not on dialysis: 1 month<br>of age and older<br>• Anemia due to concomitant<br>myelosuppressive<br>chemotherapy: 5 years of age<br>and older<br>2 idovudine-treated, anemia,<br>patients with HIV infection: 8<br>months and older | 1/12/2022            |
| Biologicals | J0887         | Injection, epoetin beta, 1<br>microgram, (for ESRD on<br>dialysis)                      | 1 mcg                      | 1/1/2015                | Mircera®          | methoxy polyethylene glycol-<br>epoetin beta injection, for<br>intravenous or subcutaneous<br>use (for ESRD on dialysis) | Indicated for the treatment of anemia associated with chronic kidney disease (CKD) in:<br>• adult patients on dialysis and adult patients not on dialysis.<br>• pediatric patients 3 months to 17 years of age on dialysis or not on dialysis who are converting from<br>another ESA after their hemoglobin level was stabilized with an ESA.<br>Limitations of Use:<br>Mircrear is not indicated and is not recommended for use:<br>• In the treatment of anemia due to cancer chemotherapy<br>• As a substitute for RBC transfusions in patients who require immediate correction of anemia.<br>Mircrear has not been shown to improve quality of life, fatigue, or patient well-being.   | 720                               | Indication Specific<br>Age Restrictions (see<br>comments) | N/A         | N/A                    | Y               | ¥                               | Patients converting from<br>another ESA after their<br>hemoglobin level was<br>stabilized with an ESA: 3<br>months of age and older<br>Patients not converting from<br>another ESA after their<br>hemoglobin level was<br>stabilized with an ESA: 1<br>years of age and older        | 5/23/2024            |
| Biologicals | 3888U         | Injection, epoetin beta, 1<br>microgram, (for non-ESRD use)                             | 1 mcg                      | 1/1/2015                | Mircera®          | methoxy polyethylene glycol-<br>epoetin beta injection, for<br>intravenous or subcutaneous<br>use (for non-ESRD use)     | Indicated for the treatment of anemia associated with chronic kidney disease (CKD) in:<br>• adult patients on dialysis and adult patients not on dialysis.<br>• pediatric patients 3 months to 17 years of age on dialysis or not on dialysis who are converting from<br>another ESA after their hemoglobin level was stabilized with an ESA.<br>Limitations of Use:<br>Mircera is not indicated and is not recommended for use:<br>• In the treatment of anemia due to cancer chemotherapy<br>• As a substitute for RBC transfusions in patients who require immediate correction of anemia.<br>Mircera has not been shown to improve quality of life, fatigue, or patient well-being.   | 720                               | Indication Specific<br>Age Restrictions (see<br>comments) | N/A         | N/A                    | Y               | ¥                               | Patients converting from<br>another ESA after their<br>hemoglobin level was<br>stabilized with an ESA: 3<br>months of age and older<br>Patients not converting from<br>another ESA after their<br>hemoglobin level was<br>stabilized with an ESA: 18<br>years of age and older       | 5/23/2024            |
| Drugs       | J0893         | Injection, decitabine (sun<br>pharma), not therapeutically<br>equivalent to j0894, 1 mg | 1 mg                       | 1/1/2023                | N/A               | decitabine for injection, for<br>intravenous use (Sun<br>Pharma)   | Indicated for treatment of adult patients with myelodysplastic syndromes (MDS) including previously<br>treated and untreated, de novo and secondary MDS of all French-American-British subtypes (refractory<br>anemia, refractory anemia with ringed sideroblasts, refractory anemia with excess blasts, refractory<br>anemia with excess blasts in transformation, and chronic myelomonocytic leukemia) and intermediate-1,<br>intermediate-2, and high-risk International Prognostic Scoring System groups.   | 450                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 12/6/2022            |
| Drugs       | J0894         | Injection, decitabine, 1 mg   | 1 mg                       | 1/1/2007                | N/A               | decitabine for injection, for<br>intravenous infusion  | Indicated for treatment of patients with myelodysplastic syndromes (MDS) including previously treated<br>and untreated, de novo and secondary MDS of all French-American-British subtypes (refractory anemia,<br>refractory anemia with ringed sideroblasts, refractory anemia with excess blasts, refractory anemia with<br>excess blasts in transformation, and chronic myelomoncytic leukemia) and intermediate-1, intermediate-<br>2, and high-risk international Prognostic Scoring System groups.   | 450                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/4/2018            |
| Drugs       | J0895         | Injection, deferoxamine<br>mesylate, 500 mg   | 500 mg                     | 1/1/2000                | Desferal®         | deferoxamine mesylate for<br>injection   | Indicated for the treatment of acute iron intoxication and of chronic iron overload due to transfusion-<br>dependent anemias.   | 372                               | 3 years   | N/A         | N/A                    | Y               | Y                               |  | 10/4/2018            |
| Biologicals | 10896         | Injection, luspatercept-aamt,<br>0.25 mg  | 0.25 mg                    | 7/1/2020                | Reblozy!®         | luspatercept-aamt for<br>injection, for subcutaneous<br>use  | Thickated for the treatment of:<br>* anemia in adult patients with beta thalassemia who require regular red blood cell (RBC) transfusions.<br>* anemia failing an erythropoiesis stimulating agent and requiring 2 or more RBC units over 8 weeks in<br>adult patients with very low- to intermediate-risk myelodysplastic syndromes with ring sideroblasts (MDS-<br>RS) or with myelodysplastic/myeloproliferative neoplasm with ring sideroblasts and thrombocytosis<br>(MDS/MPN-RS-T).<br>* anemia without previous erythropoiesis stimulating agent use (ESA-naïve) in adult patients with very low<br>to intermediate-risk myelodysplastic syndromes (MDS) who may require regular red blood cell (RBC)<br>transfusions of Use:<br>Rebloavi is not indicated for use as a substitute for RBC transfusions in patients who require immediate<br>correction of anemia. |                                   | 18 years  | N/A         | N/A                    | ¥               | Ŷ                               |  | 9/28/2023            |

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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit                                      | HCPCS<br>Effective Date | Brand Name                               | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modifie<br>Date |
|-------------|---------------|--|---|-------------------------|--|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|----------------------|
| Biologicals | J0897         | Injection, denosumab, 1 mg<br>(Xgeva, Prolia)  | 1 mg  | 1/1/2012                | Prolia <sup>®</sup> , Xgeva <sup>®</sup> | denosumab injection, for<br>subcutaneous use  | Prolia<br>Indicated for:<br>The treatment in postmenopausal women with osteoporosis at high risk for fracture<br>The treatment to increase bone mass in men with osteoporosis at high risk for fracture<br>The treatment to increase bone mass in women at high risk for fracture receiving androgen deprivation<br>therapy for nonmetastatic prostate cancer<br>The treatment to increase bone mass in women at high risk for fracture receiving adjuvant aromatase<br>inhibitor therapy for breast cancer.<br>The treatment of glucocorticoid-induced osteoporosis in men and women at high risk for fracture.<br>Xgeva<br>Indicated for:<br>The grevention of skeletal-related events in patients with multiple myeloma and in patients with bone<br>metastases from solid tumors<br>The treatment of adults and skeletally mature adolescents with glant cell tumor of bone that is<br>unresectable or where surgical resections i likely to result in severe motholity<br>The treatment of hypercalceming of malignarcy refractory to bisphosphonate therapy  | 360                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Product/indication specific age<br>restrictions:<br>• Prolia: 18 years of age and<br>older<br>• Sgevs: indication specific.<br>o Giant cell tumor of bone:<br>Only use in skeletally mature<br>adolescents.<br>o All other indications: 18<br>years of age and older | 10/31/2018           |
| Drugs       | J0911         | Instillation, taurolidine 1.35<br>mg and heparin sodium 100<br>units (central venous catheter<br>lock for adult patients<br>receiving chronic<br>hemodialysis) | taurolidine 1.35 mg<br>and heparin sodium<br>100 units (0.1 mL) | 7/1/2024                | DefenCath®                               | taurolidine and heparin<br>catheter lock solution, for<br>central venous catheter<br>instillation use | Taurolidine and heparin catheter lock solution is indicated to reduce the incidence of catheter-related<br>bloodstream infections (RBSI) in adult patients with kidney failure receiving chronic hemodiahysis (HD)<br>through a central venous catheter (CVC). This drug is indicated for use in a limited and specific population<br>of patients.<br>Limitations of Use<br>The safety and effectiveness of DefenCath have not been established for use in populations other than<br>adult patients with kidney failure receiving chronic HD through a CVC.  | 700                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 7/29/2024            |
| Drugs       | J1000         | Injection, depo-estradiol<br>cypionate, up to 5 mg   | up to 5 mg  | 1/1/2000                | Depo®-Estradiol                          | estradiol cypionate injection   | Indicated in the treatment of hypoestrogenism caused by hypogonadism and moderate to severe<br>vasomotor symptoms associated with the menopause.<br>Indicated as follows when the oral route is not Teasible:  | 2                                 | 18 years  | N/A         | Females Only           | Y               | Y                               |  | 10/4/2018            |
| Drugs       | 11010         | Injection, methylprednisolone<br>acetate, 1 mg   | 1 mg  | 4/1/2024                | Depo-Medrol*                             | methylprednisolone acetate<br>injection, suspension, USP  | Intramucular Administration  Allergis States: Control of severe or incapacitating allergic conditions intractable to adequate trials of  conventional treatment in asthma, atopic dermatitis, contact dermatitis, drug hypersensitivity reactions,  serum sichness; transfusion reactions.  Dermatologic Diseases: Bullous dermatitis herpetiformis, enfoliative dermatitis, mycosis fungoides,  pemphigus, severe erythema multiforme (Stevens-Johnson syndrome).  C indicating Disorders: Primary or secondary adenocortical insufficiency (hydrocortisone or cortisone is  the drug of choice; synthetic analogs may be used in conjunction with mineralocorticoids where  applicable, in infanzy, mineralocorticid supplementations is of particular importance), congenital adrenal  hyperplaia, hypercalcemia associated with cancer, nonsupportive thyroidits. (Sastrointestinal Diseases: To tick the patient over a critical period of the disease in regional entertits  (systemi: therapy) and ulcerative colitis.  Hematologic Disorders: Agrim equil (autoimmune) hemolytic anemia, congenital (erythroid) hypoplastic  | 800                               | N/A   | N/A         | N/A                    | ¥               | ¥                               |  | 3/22/2024            |
| Drugs       | J1050         | Injection,<br>medroxyprogesterone acetate,<br>1 mg   | 1 mg  | 1/1/2013                | Depo-Provera®                            | medroxyprogesterone<br>acetate, injectable<br>suspension  | Indicated for prevention of pregnancy in females and adjunctive therapy and palliative treatment of<br>inoperable, recurrent, and metastatic endometrial or renal carcinoma.   | 5,000                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Ŷ               | Ŷ                               | Indication specific age<br>restrictions:<br>• Endometrial and renal<br>carcinoma: 18 years and older<br>• Prevention of pregnancy: Use<br>after menarche.  |                      |
| Drugs       | J1071         | Injection, testosterone<br>cypionate, 1 mg   | 1 mg  | 1/1/2015                | Depo®-<br>Testosterone                   | testosterone cypionate<br>injection, USP  | Indicated for replacement therapy in the male in conditions associated with symptoms of deficiency or<br>absence of endogenous testosterone.<br>J. Primary hypogenadism (congenital or acquired)-testicular failure due to cryptorchidism, bilateral<br>torsion, orchits, vanishing testis syndrome; or archidectomy.<br>J. Hypogenadism <sup>(congenital or acquired)</sup> -genadotropin or LHRH deficiency, or<br>pltutary-hypothalamic injury from tumors, trauma, or radiation.<br>Safety and efficacy of Depo-Testosterone (testosterone cypionate) in men with "age-related<br>hypogenadism <sup>(congenital or active states or the pogenadism<sup>(congenital or active states or the entert states of the st</sup></sup> | 1,200                             | 12 years  | N/A         | Males Only             | Y               | Ŷ                               |  | 4/10/2019            |
| Drugs       | J1095         | Injection, dexamethasone 9<br>percent, intraocular, 1<br>microgram   | 1 mcg   | 1/1/2019                | Dexycu™                                  | dexamethasone intraocular<br>suspension 9%, for<br>intraocular administration                         | Indicated for the treatment of postoperative inflammation.   | 1,034                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 3/26/2019            |
| Drugs       | J1096         | Dexamethasone, lacrimal<br>ophthalmic insert, 0.1 mg   | 0.1 mg  | 10/1/2019               | Dextenza®                                | dexamethasone ophthalmic<br>insert 0.4 mg, for<br>intracanalicular use                                | Indicated for:<br>• The treatment of ocular inflammation and pain following ophthalmic surgery.<br>• The treatment of ocular itching associated with allergic conjunctivitis.  | 8                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 11/17/2021           |

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| Category | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|----------|---------------|--|----------------------------|-------------------------|------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Drugs    | J1097         | phenylephrine 10.16 mg/ml<br>and ketorolac 2.88 mg/ml<br>ophthalmic irrigation solution,<br>1 ml | 1 mL                       | 10/1/2019               | Omidria®   | phenylephrine and ketorolac<br>intraocular solution, 1%<br>/0.3%, for addition to ocular<br>irrigating solution |   | 8                                 | N/A   | N/A         | N/A                    | Y               | Y                               |  | 9/27/2019             |
| Drugs    | J1100         | Injection, dexamethasone<br>sodium phosphate, 1 mg   | 1 mg                       | 1/1/2000                | N/A        | dexamethasone sodium<br>phosphate injection   | Intravenous or intramuscular Administration: when oral interapy is not reasine and the strength, obsige<br>form, and route of administration of the drug reasonably lend the preparation to the transment of the<br>condition, those products labeled for intravenous or intramuscular use are indicated as follows:<br>Endocrine Disorders: Primary or secondary administration is of particular importance). Acute<br>ademicontical insufficiency (hydrocortisone or cortisone is the drug of choice, mineralocorticoid<br>supplicable; in infancy, mineralocorticoid supplementation is of particular importance). Acute<br>ademicortical insufficiency (hydrocortisone or cortisone is the drug of choice, mineralocorticoid<br>supplementation may be necessary, particularly when synthetic analogs are used), Prosperatively, and in<br>the event of serious trauma or illnes, in patients with horown adrenal insufficiency or when adrencocrtical<br>reserve is doubtful, shock unresponsive to conventional therapy if adrencocrtical insufficiency when adrencocrtical<br>reserve is doubtful, shock unresponsive to conventional threapy if adrencocrtical insufficiency exists or is<br>suspected. Congenital adrenal hyperplasia, Nonsuppurative thyroiditis, synovitis of osteoarthritis, rheumatoid<br>arthritis including juvelie heumatotical arthritis, specificated cases may require low-dose maintenance<br>therapy, acute and subacute bursitis, epicondylitis, acute nonspecific tenosynovitis, acute gouty arthritis,<br>poriatic arthritis, and akylosing spondylitis.<br>• Collegen Diseases: During an exacerbation or as maintenance therapy in selected cases of systemic lupus<br>erythematosus and acute fluensatic carditis.<br>• Jermatologic biosesses: Partinguis, severe erythema multiforme (Stevers-Johnson Syndrome),<br>edollative dermatitis, bulous dermatitis, therpetiformis, severe seborchic dermatitis, severe portasis, and<br>mycosis fungides.<br>• Allengic States: control of severe or incapacitating allergic conditions intractable to adequate trials of<br>conventional treatment in thronchial asthma, contact dermatitis, sexport | 310                               | N/A   | N/A         | N/A                    | ¥               | Ÿ                               |  | 10/4/2018             |
| Drugs    | J1105         | Dexmedetomidine, oral, 1 mcg   | 1 mcg                      | 1/1/2024                | Igalmi™    | dexmedetomidine sublingual<br>film, for sublingual or buccal<br>use   |   | 1,800                             | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 12/22/2023            |
| Drugs    | J1110         | Injection, dihydroergotamine<br>mesylate, per 1 mg   | 1 mg                       | 1/1/2000                | DHE 45®    | dihydroergotamine mesylate<br>injection   | Indicated for the acute treatment of migraine headaches with or without aura and the acute treatment of<br>cluster headache episodes.   | 30                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/10/2018            |
| Drugs    | J1120         | Injection, acetazolamide<br>sodium, up to 500 mg   | up to 500 mg               | 1/1/2000                | Diamox®    | acetazolamide sodium<br>injection, powder,<br>lyophilized, for solution   | Indicated for the adjunctive treatment of:<br>• Edema due to congestive heart failure<br>• Drug-induced edema<br>• Centrencephalic epilepsies (petit mal, unlocalized seizures)<br>• Chronic simple (open-angle) glaucoma<br>• Secondary glaucoma<br>• Preoperatively in acute angle-closure glaucoma where delay of surgery is desired in order to lower<br>intraocular pressure   | 62                                | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 10/31/2018            |
| Drugs    | J1160         | Injection, digoxin, up to 0.5 mg   | up to 0.5 mg               | 1/1/2000                | Lanoxin®   | digoxin injection, for<br>intravenous or intramuscular<br>use   | Indicated for:  | 1 35                              | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Mild to moderate heart<br>failure and control of resting<br>ventricular rate in chronic<br>atrial fibrillation: 18 years of<br>age and older<br>• Increasing myocardial<br>contractility: None | 10/10/2018            |
| Drugs    | J1165         | Injection, phenytoin sodium,<br>per 50 mg  | per 50 mg                  | 1/1/2000                | N/A        | phenytoin sodium injection,<br>for intravenous or<br>intramuscular use  | Indicated for the treatment of generalized tonic-clonic status epilepticus and prevention and treatment of<br>seizures occurring during neurosurgery. Intravenous phenytoin can also be substituted, as short-term use,<br>for oral phenytoin. Parenteral phenytoin should be used only when oral phenytoin administration is not<br>possible.  |                                   | N/A   | N/A         | N/A                    | Y               | Ŷ                               |  | 6/8/2019              |
| Drugs    | J1170         | Injection, hydromorphone, up<br>to 4 mg  | up to 4 mg                 | 1/1/2000                | Dilaudid®  | hydromorphone<br>hydrochloride for<br>intravenous, intramuscular,<br>and subcutaneous use                       | Indicated for the management of pain severe enough to require an opioid analgesic and for which<br>alternate treatments are inadequate.<br>L'imitations of Use: Because of the risks of addiction, abuse, and misuse with opioids, even at<br>recommende dosse, reserve hydromorphone injection for use in patients for whom alternative<br>treatment options [e.g., nonopioid analgesics or opioid combination products]:<br>• Have not been tolerrated, or are not expected to be tolerated<br>• Have not provide adequate analgesia, or are not expected to provide adequate analgesia   | 186                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 10/26/2018            |

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 The HCPCS Code effective date represents the date the HCPCS code was established

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| Category    | HCPCS<br>Code | HCPCS Description                                   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name            | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions   | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|-------------|---------------|---|----------------------------|-------------------------|-----------------------|---|---|-----------------------------------|---|-------------|--|-----------------|---------------------------------|--|-----------------------|
| Drugs       | J1190         | Injection, dexrazoxane<br>hydrochloride, per 250 mg | 250 mg                     | 1/1/2000                | Totect®,<br>Zinecard® | dexrazoxane for injection                                       | Zinecard: Indicated for reducing the incidence and severity of cardiomyopathy associated with doxorubicin<br>administration in women with metastatic breast cancer who have received a cumulative doxorubicin dose<br>of 300 mg/m <sup>2</sup> and who will continue to receive doxorubicin therapy to maintain tumor control. Do not use<br>with doxorubicin initiation.<br>Totect: Indicated for the treatment of extravasation resulting from IV anthracycline chemotherapy.<br>• Reducing the incidence and severity of cardiomyopathy associated with doxorubicin administration in<br>women with metastatic breast cancer who have received a cumulative doxorubicin administration in<br>doxorubicin initiation.   | 20                                | 18 years  | N/A         | Zinecard: Females<br>Only<br>Totect:<br>Extravasation:<br>N/A<br>Cardiomyopathy:<br>Females only | Y               | Y                               |  | 12/28/2020            |
| Drugs       | J1200         | Injection, diphenhydramine<br>HCI, up to 50 mg      | 50 mg                      | 1/1/2000                | N/A                   | diphenhydramine<br>hydrochloride injection                      | Diphenhydramine in the injectable form is effective in adults and pediatric patients, other than premature<br>infants and neonates, for the following conditions when diphenhydramine in the oral form is impractical:<br>Anthistaminic: For amelioration of allergic reactions to blood or plasma, in anaphysicsia sa na dijunct to<br>epinephrine and other standard measures after the acute symptoms have been controlled, and for other<br>uncomplicated allergic conditions of the immediate type when oral therapy is impossible or<br>contraindicated. A standard measures after the acute symptoms have been controlled, and for other<br>contraindicated. A standard measures after the acute symptoms have been controlled, and for other<br>antiparkinsonism. For use in parkinsonism, when or and therapy is impossible or contraindicated, as<br>follows: parkinsonism in othe inderly who are unable to tolerate more potent agents; mild cases of<br>parkinsonism in other age groups, and in other cases of parkinsonism in combination with centrally acting<br>anticholinergic agents. | 248                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A  | Y               | Y                               | Contraindicated in newborns<br>or premature infants.   | 10/4/2018             |
| Drugs       | J1202         | Miglustat, oral, 65 mg                              | 65 mg                      | 4/1/2024                | Opfolda™              | miglustat capsules, for oral<br>use                             | Miglustat capsule is indicated, in combination with Pombiliti, for the treatment of adult patients with late-<br>onset Pompe disease (lysosomal acid alpha-glucosidase [GAA] deficiency) weighing ≥40 kg and who are<br>not improving on their current enzyme replacement therapy (ERT).  | 12                                | 18 years  | N/A         | N/A  | Y               | Y                               |  | 3/22/2024             |
| Biologicals | J1203         | Injection, cipaglucosidase alfa-<br>atga, 5 mg      | 5 mg                       | 4/1/2024                | Pombiliti™            | cipaglucosidase alfa-atga for<br>injection, for intravenous use | onset Pompe disease (lysosomal acid alpha-glucosidase (GAA) deficiency) weighing >40 kg and who are   | 1,701                             | 18 years  | N/A         | N/A  | Y               | Y                               |  | 3/22/2024             |
| Drugs       | J1205         | Injection, chlorothiazide<br>sodium, per 500 mg     | 500 mg                     | 1/1/2000                | N/A                   | chlorothiazide sodium for<br>injection                          | Indicated as adjunctive therapy in edema associated with congestive heart failure, hepatic cirrhosis, and<br>corticosteroid and estrogen therapy.   | 100                               | 18 years  | N/A         | N/A  | Y               | Y                               |  | 9/27/2018             |
| Drugs       | J1212         | Injection, DMSO, dimethyl<br>sulfoxide, 50%, 50 mL  | 50 mL                      | 1/1/2000                | RIMSO-50®             | dimethyl sulfoxide (DMSO)<br>irrigation                         | Indicated for symptomatic relief of patients with interstitial cystitis.  | 3                                 | N/A   | N/A         | N/A  | Y               | Y                               |  | 10/4/2018             |
| Drugs       | J1230         | Injection, methadone HCl, up<br>to 10 mg            | up to 10 mg                | 1/1/2000                | N/A                   | methadone hydrochloride<br>injection                            | Indicated for:<br>• The management of pain severe enough to require an opioid analgesic and for which alternative<br>treatment options are inadequate.<br>Limitations of Use: Because of the risks of addiction, abuse, and misuse with opioids, even at<br>recommended doses, reserve methadone injection for use in patients for whom alternative treatment<br>options (e.g. non-opioid analgesics or opioid combination products):<br>• O Have not been tolerated, or are not expected to be tolerated.<br>• Use in temporary treatment of opioid dependence in patients unable to take oral medication.<br>Limitations of Use: Injectable methadone products are not approved for the outpatient treatment of<br>opioid dependence. In this patient population, parenteral methadone is to be used only for patients<br>unable to take oral medication, such as hospitalized patients.  | 93                                | 18 years  | N/A         | N/A  | Y               | Y                               |  | 10/26/2018            |
| Drugs       | J1240         | Injection, dimenhydrinate, up<br>to 50 mg           | up to 50 mg                | 1/1/2000                | N/A                   | dimenhydrinate injection  | Indicated for prevention and treatment of nausea, vomiting and vertigo of motion sickness.  | 372                               | N/A   | N/A         | N/A  | Y               | Y                               |  | 6/10/2019             |
| Drugs       | J1245         | Injection, dipyridamole, per 10<br>mg               | per 10 mg                  | 1/1/2000                | N/A                   | dipyridamole injection  | As an alternative to exercise in thallium myocardial perfusion imaging for the evaluation of coronary<br>artery disease in patients who cannot exercise adequately.   | 6                                 | 18 years  | N/A         | N/A  | Y               | Y                               |  | 6/10/2019             |
| Drugs       | J1250         | Injection, dobutamine<br>hydrochloride, per 250 mg  | 250 mg                     | 1/1/2000                | N/A                   | dobutamine injection  | Indicated:<br>• When parenteral therapy is necessary for inotropic support in the short-term treatment of adults with<br>cardiac decompensation due to depressed contractility resulting either from organic heart disease or from<br>cardiac surgical procedures.<br>• In patients who have atrial fibrillation with rapid ventricular response, a digitalis preparation should be<br>used prior to institution of therapy with dobutamine.  | 930                               | 18 years  | N/A         | N/A  | Y               | Y                               |  | 10/4/2018             |
| Drugs       | J1265         | Injection, dopamine<br>hydrochloride, 40 mg         | 40 mg                      | 1/1/2006                | N/A                   | dopamine hydrochloride  | Indicated for the correction of hemodynamic imbalances present in the shock syndrome due to myocardial<br>infarction, trauma, andotoxic septicemia, open-heart surgery, renal failure, and chronic cardiac<br>decompensation as in congestive failure.  | 6,355                             | 18 years  | N/A         | N/A  | Y               | Y                               |  | 10/4/2018             |
| Drugs       | J1270         | Injection, doxercalciferol, 1<br>mcg                | 1 mcg                      | 1/1/2002                | Hectorol®             | doxercalciferol injection                                       | Indicated for the treatment of secondary hyperparathyroidism in adult patients with chronic kidney<br>disease on dialysis.  | 90                                | 18 years  | N/A         | N/A  | Y               | Y                               |  | 10/4/2018             |
| Drugs       | J1290         | Injection, ecallantide, 1 mg                        | 1 mg                       | 1/1/2011                | Kalbitor®             | ecallantide injection for<br>subcutaneous use                   | Indicated for treatment of acute attacks of hereditary angioedema in patients 12 years of age and older.  | 120                               | 12 years  | N/A         | N/A  | Y               | Y                               |  | 10/10/2018            |
| Biologicals | J1300         | Injection, eculizumab, 10 mg                        | 10 mg                      | 1/1/2008                | Soliris®              | eculizumab injection, for<br>intravenous use                    | Indicated for:<br>• Treatment of patients with paroxysmal nocturnal hemoglobinuria (PNH) to reduce hemolysis.<br>• Treatment of patients with atypical hemolytic uremic syndrome (aHUS) to inhibit complement-mediated<br>thrombotic microangiopathy.<br>• Treatment of adult patients with generalized Myasthenia Gravis (gMG) who are anti-acetyicholine<br>receptor (ARb) antibody positive.<br>• Treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult patients who are anti-aquaporin-<br>4 (AQP4) antibody positive.   | 480                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A  | Y               | ¥                               | Indication specific age<br>restrictions:<br>• PNH: 18 years of age and<br>older<br>• aHUS: None<br>• Myasthenia Gravis: 18 years<br>of age and older | 7/26/2019             |
|             |               |   |                            |                         |                       | edaravone injection, for  | Limitation of Use: Soliris is not indicated for the treatment of patients with Shiga toxin E. coli related<br>hemolytic uremic syndrome (STEC-HUS).   |                                   |   |             |  |                 |                                 |  |                       |
| Drugs       | J1301         | Injection, edaravone, 1 mg                          | 1 mg                       | 1/1/2019                | Radicava®             | intravenous use   | Indicated for the treatment of amyotrophic lateral sclerosis (ALS).   | 1,020                             | 18 years  | N/A         | N/A  | Y               | Y                               |  | 10/10/2018            |

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| medicaid/med | icaid-ncci-eo | dit-files   |                            |                         |                   |  |  |                                   |   |             |                        |                 |                                 |  |                       |
|--------------|---------------|---|----------------------------|-------------------------|-------------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description                                   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name        | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Biologicals  | J1302         | Injection, sutimlimab-jome, 10<br>mg                | 10 mg                      | 10/1/2022               | Enjaymo™          | sutimlimab-jome injection,<br>for intravenous use              | Indicated for the treatment of hemolysis in adults with cold agglutinin disease (CAD).   | 2,310                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 2/23/2023             |
| Biologicals  | J1303         | Injection, ravulizumab-cwvz,<br>10 mg               | 10 mg                      | 10/1/2019               | Ultomiris™        | ravulizumab-cwvz injection,<br>for intravenous use             | Indicated for:<br>- the treatment of adult and pediatric patients one month of age and older with paroxysmal nocturnal<br>hemoglobinuria (RMH).<br>- the treatment of adults and pediatric patients one month of age and older with atypical hemolytic<br>uremic syndrome (aHUS) to inhibit complement-mediated thrombotic microangiopathy (TMA).<br>Limitations of Use:<br>Ultomiris is not indicated for the treatment of patients with Shiga toxin E. coli related hemolytic uremic<br>syndrome (STEC-HUS).<br>- the treatment of adult patients with generalized myasthenia gravis (gMG) who are anti-acetylcholine<br>receptor (ACNR) antibody-positive.  | 660                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Ŷ               | ¥                               | PNH and aHUS: 1 month of age<br>and older<br>gMG and NMOSD: 18 years of<br>age and older | 5/3/2024              |
| Drugs        | J1304         | Injection, tofersen, 1 mg                           | 1 mg                       | 1/1/2024                | Qalsody™          | tofersen injection, for<br>intrathecal use                     | Indicated for the treatment of amyotrophic lateral sclerosis (ALS) in adults who have a mutation in the<br>superoxide dismutase 1 (SOD1) gene.   | 300                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/22/2023            |
| Biologicals  | J1305         | Injection, evinacumab-dgnb,<br>5mg                  | 5 mg                       | 10/1/2021               | Evkeeza™          | evinacumab-dgnb injection,<br>for intravenous use              | Indicated as an adjunct to other low-density lipoprotein-cholesterol (LDL-C) lowering therapies for the<br>treatment of adult and pediatric patients, aged 5 years and older, with homozygous familial<br>hypercholesterolemia (HoFH).<br>Limitations of Use:<br>- The safety and effectiveness of Evkeeza have not been established in patients with other causes of<br>hypercholesterolemia, including those with heterozygous familial hypercholesterolemia (HeFH).<br>• The effects of Evkeeza on cardiovascular morbidity and mortality have not been determined.   | 894                               | 5 years   | N/A         | N/A                    | Y               | Ŷ                               |  | 4/25/2023             |
| Drugs        | J1306         | Injection, inclisiran, 1 mg                         | 1 mg                       | 1/1/2000                | Leqvio*           | inclisiran injection, for<br>subcutaneous use                  | Indicated as an adjunct to diet and maximally tolerated statin therapy for the treatment of adults with<br>heterozygous familial hypercholesterolemia (HeFH) or clinical atherosclerotic cardiovascular disease<br>(ASCVD), who require additional lowering of low-density lipoprotein cholesterol (LDL-C).  | 284                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/13/2023             |
| Biologicals  | J1322         | Injection, elosulfase alfa, 1 mg                    | 1 mg                       | 1/1/2015                | Vimizim®          | elosulfase alfa injection, for<br>intravenous use              | Indicated for patients with Mucopolysaccharidosis type IVA (MPS IVA; Morquio A syndrome).  | 1,400                             | 5 years   | N/A         | N/A                    | Y               | Y                               |  | 6/8/2019              |
| Biologicals  | J1323         | Injection, elranatamab-bcmm,<br>1 mg                | 1 mg                       | 4/1/2024                | Elrexfio™         | elranatamab-bcmm injection,<br>for subcutaneous use            | agent, and an anti-CD38 monoclonal antibody.   | 380                               | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 4/12/2024             |
| Drugs        | J1325         | Injection, epoprostenol, 0.5<br>mg                  | 0.5 mg                     | 1/1/2000                | Flolan®, Veletri® | epoprostenol for injection,<br>for intravenous use             | Indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group I) to improve exercise<br>capacity. Studies establishing effectiveness included predominantly (97%) patients with NYHA Functional<br>Class III-IV symptoms and etiologies of idiopathic or heritable PAH (49%) or PAH associated with<br>connective tissue diseases (51%).   | 248                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 6/4/2019              |
| Drugs        | J1335         | Injection, ertapenem sodium,<br>500 mg              | 500 mg                     | 1/1/2004                | Invanz®           | ertapenem injection for<br>intravenous or intramuscular<br>use | Indicated in adult patients and pediatric patients (3 months of age and older) for the treatment of the<br>following moderate to severe infections caused by susceptible bacteria:<br>- Complicated strins abdominal infections.<br>- Complicated strin and strin structure infections, including diabetic foot infections without osteomyelitis.<br>- Complicated uniany tract infections including pyelonephritis.<br>- Complicated uniany tract infections including postpartum endomyometritis, septic abortion and post surgical<br>genecologic infections.   | 28                                | 3 months  | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 10/10/2018            |
| Drugs        | J1364         | Injection, erythromycin<br>lactobionate, per 500 mg | 500 mg                     | 1/1/2000                | Erythrocin™       | erythromycin lactobionate<br>for injection                     | Indicated in the treatment of infections caused by susceptible strains of the designated organisms in the<br>diseases listed below when oral administration is not possible or when the severity of the infection<br>requires immediate high serum levels of erythromycin. Intravenous therapy should be replaced by oral<br>administration at the appropriate time.<br>• Upper respiratory tract infections of mild to moderate degree caused by Streptococcus progenes (Group<br>A beta-hemolytic streptococci). Streptococcus penumoniae (Diplococcus penumoniae). Heamophilus<br>influenzae (when used concomitantly with adequate doses of sulfonamides, since many strains of H.<br>influenzae (when used concomitantly with adequate doses of sulfonamides, since many strains of H.<br>influenzae (when used concomitantly with adequate doses of sulfonamides, since many strains of H.<br>influenzae (when used concomitantly with adequate doses of sulfonamides, since many strains of H.<br>influenzae (when used concomitantly with adequate doses of sulfonamides, since many strains of H.<br>influenzae (are nearboy)tic streptococcu). Streptococcus penumoniae (Diplococcus preumoniae).<br>• Sulfan and sins instructure infections of mild to moderate severity caused by Streptococcus progenes<br>and Staphylococcus aureus (resistant staphylococci may emerge during treatment).<br>• Diphtheria: As an adjunct to anticosin infections due to Conyrebacterium diphtheriae to prevent<br>establishment of carriers and to eardicate the organism in carriers.<br>• Erythrama: in the treatment of infections due to Conyrebacterium minutissimum.<br>• Acute pavic inflammatory disease caused by Nesseria gonorrhoeae: Erythromic lactobionate-IV<br>(erythromycin lactobionate for injection, USP) followed by enthromycin stearate or erythromycin base<br>orially, as an alternative drug in treatment of acute peric inflamatory disease caused by Nesseria gonorrhoeae<br>in female patients with a history of sensitivity to penicillin.<br>• Before treatment of gonorrhae, patients who are suspected of also having syphilis should h | 248                               | N/A   | N/A         | N/A                    | Y               | ¥                               |  | 10/10/2018            |

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| medicaid/medi | caid-ncci-eo  | lit-files   | r.                         |                         |                          |   |   |                                   |   |             |                        |                 |                                 |   |                       |
|---------------|---------------|---|----------------------------|-------------------------|--------------------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name               | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Drugs         | J1380         | Injection, estradiol valerate,<br>up to 10 mg   | up to 10 mg                | 1/1/2000                | Delestrogen®             | estradiol valerate injection  | Indicated in the treatment of:<br>• Moderate-to-severe vasomotor symptoms associated with the menopause<br>• Hypoestrogenism caused by hypogonadism, castration or primary ovarian failure<br>• Advanced androgen-dependent carcinoma of the prostate (for palliation only)<br>• Uvluvi and vaginal atrophy associated with the menopause. When prescribing solely for the treatment<br>of symptoms of vulvar and vaginal atrophy, topical vaginal products should be considered.   | 20                                | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 6/10/2019             |
| Drugs         | J1410         | Injection, estrogens,<br>conjugated, per 25 mg  | 25 mg                      | 1/1/2000                | Premarin <sup>®</sup> IV |   | Indicated in the treatment of abnormal uterine bleeding caused by hormonal imbalance in the absence of<br>organic pathology. Indicated for short-term use only, to provide a rapid and temporary increase in<br>estrogen levels.  | 62                                | N/A   | N/A         | Females Only           | Y               | Ŷ                               |   | 10/10/2018            |
| Drugs         | J1437         | Injection, ferric derisomaltose,<br>10 mg   | 10 mg                      | 10/1/2020               | MonoFerric™              | ferric derisomaltose injection,<br>for intravenous use  | Indicated for the treatment of iron deficiency anemia in adult patients:<br>• who have intolerance to oral iron or have had unsatisfactory response to oral iron.<br>• who have non-hemodialysis dependent chronic kidney disease.  | 100                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 12/28/2020            |
| Drugs         | J1439         | Injection, ferric<br>carboxymaltose, 1 mg   | 1 mg                       | 1/1/2015                | Injectafer*              | ferric carboxymaltose<br>injection, for intravenous use   | Indicated for the treatment of iron deficiency anemia (IDA) in adult patients:<br>- Who have intolerance to oral iron or have had unsatisfactory response to oral iron.<br>- Who have non-dialysis dependent chronic könney disease.<br>- With heart failure and New York Heart Association class I/UII to improve exercise capacity.<br>Indicated for the treatment of iron deficiency anemia in pediatric patients 1 year of age to 17 years of age<br>who have either intolerance to oral iron or an unsatisfactory response to oral iron.                             | 1,500                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Ą               | Ŷ                               | Indication specific age<br>restrictions:<br>IDA in patients who have<br>either intolerance to oral iron<br>or an unsatisfactory response<br>to oral iron: 1 year of age and<br>older<br>• IDA in patients who have non-<br>dialysis dependent chronic<br>kidney disease, iron deficiency<br>in adult patients with heart<br>failure and New York Heart<br>Association class I/III to<br>improve exercise capacity: 13<br>years of age and older | - 6/19/2023           |
| Biologicals   | J1440         | Fecal microbiota, live - jslm, 1<br>ml  | 1 mL                       | 7/1/2023                | Rebyota™                 | fecal microbiota, live - jslm<br>suspension, for rectal use   | Indicated for the prevention of recurrence of Clostridioides difficile infection (CDI) in individuals 18 years<br>of age and older, following antibiotic treatment for recurrent CDI.<br>Limitation of Use: Rebyota is not indicated for treatment of CDI.  | 150                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 6/22/2023             |
| Biologicals   | J1442         | Injection, filgrastim (G-CSF),<br>excludes biosimilars, 1<br>microgram  | 1 mcg                      | 1/1/2016                | Neupogen*                | filgrastim injection, for<br>subcutaneous or intravenous<br>use   | Indicated to:<br>• Decrease the incidence of infection, as manifested by febrile neutropenia, in patients with nonmyeloid<br>malignancies receiving myelosuppressive<br>anti-cancer drugs associated with a significant incidence of severe neutropenia with fever.<br>• Reduce the time to neutrophil recovery and the duration of fever, following induction or consolidation<br>chemotherapy treatment of patients with acute<br>myeloid leukemia (AML).<br>• Reduce the duration of neutropenia and neutropenia-related clinical sequelae, e.g., febrile neutropenia, | 59,520                            | N/A   | N/A         | N/A                    | Ŷ               | Ŷ                               |   | 6/6/2019              |
| Drugs         | J1443         | Injection, ferric pyrophosphate<br>citrate solution (triferic), 0.1<br>mg of iron   | 0.1 mg of iron             | 10/1/2021               | Triferic®                | ferric pyrophosphate citrate<br>solution, for hemodialysis<br>use, and powder for solution,<br>for hemodialysis use | Indicated for the replacement of iron to maintain hemoglobin in adult patients with hemodialysis-<br>dependent chronic kidney disease (HDD-CKD),<br>Limitations of Use:<br>• Trifferic is not intended for use in patients receiving peritoneal dialysis.<br>• Trifferic has not been studied in patients receiving home hemodialysis.  | 38,080                            | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 9/29/2021             |
| Drugs         | J1444         | Injection, ferric pyrophosphate<br>citrate powder, 0.1 mg of iron<br>(This code would be used with<br>the "JE" modifier, when<br>administered via dialysate.) | 0.1 mg                     | 7/1/2019                | Triferic®                | ferric pyrophosphate citrate<br>powder packet for<br>hemodialysis use   | Indicated for the replacement of iron to maintain hemoglobin in adult patients with hemodialysis-<br>dependent chronic kidney disease (HDD-CKD).<br>Limitations of Use:<br>- Trifferic is not intended for use in patients receiving peritoneal dialysis.<br>- Trifferic has not been studied in patients receiving home hemodialysis.  | 38,080                            | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 7/26/2019             |
| Biologicals   | J1447         | Injection, tbo-filgrastim, 1<br>microgram   | 1 mcg                      | 1/1/2016                | Granix*                  | tbo-filgrastim injection, for subcutaneous use  | Indicated in adult and pediatric patients 1 month and older for reduction in the duration of severe<br>neutropenia in patients with non-myeloid malignancies teckving myelosuppressive anti-cancer drugs<br>associated with a clinically significant incidence of febrile neutropenia.  | 10,920                            | 1 month   | N/A         | N/A                    | Y               | Y                               |   | 5/20/2019             |
| Drugs         | J1448         | Injection, trilaciclib, 1mg   | 1 mg                       | 10/1/2021               | Cosela*                  | trilaciclib for injection, for intravenous use  | Indicated to decrease the incidence of chemotherapy-induced myelosuppression in adult patients when<br>administered prior to a platinum/etoposide-containing regimen or topotecan-containing regimen for<br>extensive-stage small cell lung cancer.   | 9,000                             | 18 years  | N/A         | N/A                    | Y               | Ŷ                               | 12/2023: NC Suggested Max<br>Monthly updated from 1,200<br>units to 9,000 units effective<br>5/1/2023 at DHB request.   | 12/1/2023             |
| Biologicals   | J1449         | Injection, eflapegrastim-xnst,<br>0.1 mg  | 0.1 mg                     | 4/1/2023                | Rolvedon™                | eflapegrastim-xnst injection,<br>for subcutaneous use   | Indicated to decrease the incidence of infection, as manifested by febrile neutropenia, in adult patients<br>with nonmyeloid malignancies receiving myelosuppressive anti-cancer drugs associated with clinically<br>significant incidence of febrile neutropenia.<br>Limitations of Use:<br>Rolvedon is not indicated for the mobilization of peripheral blood progenitor cells for hematopoietic stem<br>cell transplantation.  | 396                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 3/16/2023             |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
 The HCPCS Code effective date represents the date the HCPCS code was established

• Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

| edicaid/medi<br>Category | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                  | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modifie<br>Date |
|--------------------------|---------------|---|----------------------------|-------------------------|-----------------------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|----------------------|
| Drugs                    | J1453         | Injection, fosaprepitant, 1 mg  | 1 mg                       | 1/1/2009                | Emend®                      | fosaprepitant for injection,<br>for intravenous use                                  | Indicated in adults and pediatric patients 6 months of age and older, in combination with other antiemetic<br>agents, for the prevention of:<br>• acute and delayed nausea and vomiting associated with initial and repeat courses of highly emetogenic<br>cancer chemotherapy (HEC). Including high-dose cisplatin.<br>• delayed nausea and vomiting associated with initial and repeat courses of moderately emetogenic<br>cancer chemotherapy (MEC).  | 750                               | 6 months  | N/A         | N/A                    | Y               | Y                               | 9/2023: NC Suggested Max<br>Monthly Units updated from<br>600 units to 750 units<br>effective 1/1/2023 at DHB<br>request  | 9/28/2023            |
|                          |               |   |                            |                         |                             |  | Limitations of Use: Emend has not been studied for treatment of established nausea and vomiting.   |                                   |   |             |                        |                 |                                 |   |                      |
| Drugs                    | J1454         | Injection, fosnetupitant 235<br>mg and palonosetron 0.25 mg   | 235.25 mg (1 vial)         | 1/1/2019                | Akynzeo®                    | fosnetupitant and<br>palonosetron for injection,<br>for intravenous use              | Indicated in combination with dexamethasone in adults for the prevention of acute and delayed nausea<br>and vomiting associated with initial and repeat courses of highly emetogenic cancer chemotherapy.<br>Limitations of Use:<br>Akyrace for injection has not been studied for the prevention of nausea and vomiting associated with<br>anthracycline plus cyclophosphamide chemotherapy.  | 5                                 | 18 years  | N/A         | N/A                    | Y               | Y                               | 9/1/2023: NC Suggested Max<br>Monthly Units updated to align<br>with NCTracks, which has been<br>set to 5 units/month since<br>1/1/2019.  | 9/13/2023            |
| Drugs                    | J1455         | Injection, foscarnet sodium,<br>per 1,000 mg  | 1,000 mg                   | 1/1/2000                | Foscavir®                   | foscarnet sodium injection   | Indicated for the treatment of:<br>• CMV retinitis in patients with acquired immunodeficiency syndrome (AIDS). Combination therapy with<br>Foscavir and gancicolowi is indicated for patients who have relapsed after monotherapy with either drug.<br>Safety and efficacy of foscavir have not been established for treatment of other CMV infections (e.g.<br>perumonitis, gastroenteritis); congenital or neonatal CMV disease, or nonimmunocompromised<br>individuals.<br>• Acyclovir-resistant muccourtaneous HSV infections in immunocompromised patients. Safety and efficacy<br>of Foscavir have not been established for treatment of other HSV infections (e.g. retinitis, encephalitis),<br>congenital or neonatal HSV disease, or HSV in nonimmunocompromised individuals. | 996                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 6/4/2019             |
| Drugs                    | J1456         | Injection, fosaprepitant (teva),<br>not therapeutically equivalent<br>to j1453, 1 mg                  | 1 mg                       | 1/1/2023                | N/A                         | fosaprepitant for injection,<br>for intravenous use (Teva)                           | Indicated in adults, in combination with other antiemetic agents, for the prevention of:<br>• acute and delayed nauses and vomiting associated with initial and repeat courses of highly emetogenic<br>cancer chemotherapy (HEC) including high-dose cipatiani.<br>• delayed nauses and vomiting associated with initial and repeat courses of moderately emetogenic<br>cancer chemotherapy (MEC).<br>Limitations of Use:  | 750                               | 18 years  | N/A         | N/A                    | Y               | Y                               | 9/2023: NC Suggested Max<br>Monthly Units updated from<br>600 units to 750 units<br>effective 1/1/2023 at DHB<br>request  | 9/28/2023            |
|                          |               |   |                            |                         |                             |  | Fosaprepitant for Injection has not been studied for treatment of established nausea and vomiting.   |                                   |   |             |                        |                 |                                 |   |                      |
| ologicals                | J1458         | Injection, galsulfase, 1 mg   | 1 mg                       | 1/1/2007                | Naglazyme*                  | galsulfase injection for<br>intravenous use  | Indicated for patients with Mucopolysaccharidosis VI (MPS VI; Maroteaux-Lamy syndrome). Naglazyme<br>has been shown to improve walking and stair-climbing capacity.  | 700                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 7/2/2018             |
| nmune<br>lobulins        | J1459         | Injection, immune globulin<br>(Privigen), intravenous, non-<br>lyophilized (e.g., liquid), 500<br>mg  | 500 mg                     | 1/1/2009                | Privigen*                   | immune globulin intravenous<br>(human), 10% liquid                                   | Indicated for the treatment of:<br>• Primary humoral immunodeficiency (PI)<br>• Chronic immune thrombocytopenic purpura (ITP) in patients age 15 years and older<br>• Chronic inflammatory demyelinating polyneuropathy (CIDP) in adults<br>Limitations of Use:<br>Privigen maintenance therapy in CIDP has not been studied beyond 6 months.  | 840                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | ¥                               | Indication specific age<br>erstrictions:<br>• Primary Humoral<br>Immunodeficiency: 3 years of<br>age and older<br>• Chronic Immune<br>Thrombocytopenic Purpura: 51<br>years of age and older<br>• Chronic Inflammatory<br>• Chronic Inflammatory<br>• Demyelinating<br>Polyneuropathy: 18 years of<br>age and older | 7/3/2018             |
| imune<br>obulins         | J1460         | Injection, gamma globulin,<br>intramuscular, 1 cc   | 1 cc                       | 1/1/2000                | GamaSTAN® S/D,<br>GamaSTAN® | immune globulin (human),<br>solution for intramuscular<br>injection, less than 10 cc | Indicated:<br>• For prophylaxis following exposure to hepatitis A.<br>• To prevent or modify measles in a susceptible person exposed fewer than 6 days previously.<br>• To modify varicella.<br>• To modify vubella in exposed women who will not consider a therapeutic abortion.<br>• Not indicated for routine prophylaxis or treatment of viral hepatitis type B, rubella, poliomyelitis, mumps<br>or varicella.   | 10                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 10/25/201            |
| nmune<br>lobulins        | J1554         | Injection, immune globulin<br>(asceniv), 500 mg   | 500 mg                     | 4/1/2021                | Asceniv™                    | immune globulin intravenous,<br>human – slra 10% liquid                              | Indicated for the treatment of primary humoral immunodeficiency (PI) in adults and adolescents (12 to 17 years of age).  | 460                               | 12 years  | N/A         | N/A                    | Y               | Y                               |   | 3/25/2021            |
| mmune<br>ilobulins       | J1555         | Injection, immune globulin<br>(Cuvitru), 100 mg   | 100 mg                     | 1/1/2018                | Cuvitru                     | immune globulin<br>subcutaneous (human), 20%<br>solution                             | Indicated as replacement therapy for primary humoral immunodeficiency (PI) in adult and pediatric<br>patients two years of age and older.  | 14,880                            | 2 years   | N/A         | N/A                    | Y               | Y                               |   | 9/12/2018            |
| mmune<br>ilobulins       | J1556         | Injection, immune globulin<br>(Bivigam), 500 mg   | 500 mg                     | 1/1/2014                | Bivigam®                    | immune globulin intravenous<br>(human), 10% liquid                                   | Indicated for the treatment of adults and pediatric patients 2 years of age and older with primary humoral<br>immunodeficiency (PI).   | 480                               | 2 years   | N/A         | N/A                    | Y               | Y                               |   | 2/16/2024            |
| mmune<br>lobulins        | J1557         | Injection, immune globulin,<br>(Gammaplex), intravenous,<br>non-lyophilized, (e.g. liquid),<br>500 mg | 500 mg                     | 1/1/2012                | Gammaplex®                  | immune globulin intravenous<br>(human), 5% and 10% liquid,<br>for intravenous use    | Gammaplex 5%: Indicated for the treatment of:<br>• Chronic immune thrombocytopenic purpurg (ITP).<br>• Primary humoral immundeficiency (P) in adults and pediatric patients 2 years of age and older.<br>Gammaplex 10%: Indicated for the treatment of:<br>• Primary humoral immundeficiency (PI) in adults.<br>• Chronic immune thrombocytopenic purpurg (ITP) in adults.   | 560                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Product specific age<br>restrictions:<br>Gammaplex 5%: 2 years of age<br>and older<br>Gammaplex 10%: 18 years of<br>age and older   | 9/21/2018            |
| Immune<br>Globulins      | J1558         | Injection, immune globulin<br>(xembify), 100 mg   | 100 mg                     | 7/1/2020                | Xembify®                    | immune globulin<br>subcutaneous, human – klhw<br>20% solution                        | Indicated for treatment of Primary Humoral Immunodeficiency (PI) in patients 2 years of age and older.   | 14,880                            | 2 years   | N/A         | N/A                    | Y               | Y                               |   | 6/17/2020            |
| -                        | 1             | 1   | 1                          |                         | 1                           | 2070 301001011   | 1  | 1                                 | 1   | I           | 1                      | I               |                                 | 1   | <u>ا</u>             |

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| medicaid/medi       | icaid-ncci-ed | lit-files   |                            | 1                       | 1                              |   |   | 1   | 1   |             |                        | ,               |                                 |   |                       |
|---------------------|---------------|---|----------------------------|-------------------------|--------------------------------|---|---|---|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category            | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                     | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units   | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Immune<br>Globulins | J1559         | Injection, immune globulin<br>(Hizentra), 100 mg  | 100 mg                     | 1/1/2011                | Hizentra®                      | immune globulin<br>subcutaneous (human), 20%<br>liquid  | <ul> <li>Indicated as replacement therapy for primary immunodeficiency (PI) in adults and pediatric patients 2<br/>years of age and older. This includes, but is not limited to, the humoral immune defect in congenital<br/>agarmagiobulinemia, common avaitable immunodeficiency, X-linked agammagiobulinemia, Wiskott-<br/>Aldrich syndrome and severe combined immunodeficiencies.</li> <li>Indicated as maintenance therapy for the treatment of adult patients with chronic inflammatory<br/>demyelinating polyneuropathy (CIDP) to prevent relapse of neuromusular disability and impairment.</li> </ul>   | 2,800   | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• PI - 2 years of age and older<br>• CDIP - 18 years of age and<br>older  | 7/16/2018             |
| Immune<br>Globulins | J1560         | Injection, gamma globulin,<br>intramuscular, over 10 cc<br>(always use for any amount<br>injected over 10cc and place<br>number of units) | 10 cc                      | 1/1/2000                | GamaSTAN® S/D,<br>GamaSTAN®    | immune globulin (human),<br>solution for intramuscular<br>injection greater than 10 cc  | Indicated:<br>• For prophylaxis following exposure to hepatitis A.<br>• To prevent or modify measles in a susceptible person exposed fewer than 6 days previously.<br>• To modify varicella.<br>• To modify rubella in exposed women who will not consider a therapeutic abortion.<br>• Not indicated for routine prophylaxis or treatment of viral hepatitis type B, rubella, poliomyelitis, mump:<br>or varicella.  | 17  | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 9/21/2018             |
| Immune<br>Globulins | J1561         | Injection, immune globulin,<br>(Gamunex-C/Gammaked), non<br>lyophilized (e.g. liquid), 500<br>mg  | 500 mg                     | 1/1/2013                | Gammaked™,<br>Gamunex®-C       | immune globulin injection<br>(human), 10%<br>caprylate/chromatography<br>purified   | Gamunex-C is indicated for:<br>• Primary Humoral Immunodeficiency (PI) in patients 2 years of age and older<br>• Idiopathic Thrombocytopenic Purpura (ITP) in adults and children<br>• Chronic Inflammatory Demyelinating Polyneuropathy (CIDP) in adults<br>Gammaked is indicated for:<br>• Primary Humoral Immunodeficiency (PI) in patients 2 years of age and older<br>• Idiopathic Thrombocytopenic Purpura (ITP)<br>• Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)  | 840   | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Primary Humoral<br>Immunodeficiency (PI): 2 years<br>of age and older<br>• Idiopathic Thrombocytopenic<br>Purpura (ITP): None<br>• Chronic Inflammatory<br>Demyelinating Polyneuropathy<br>(CIDP): 18 years of age and<br>older | :<br>9/12/2018        |
| Immune<br>Globulins | J1566         | Injection, immune globulin,<br>intravenous, lyophilized (e.g.<br>powder), not otherwise<br>specified, 500mg                               | 500 mg                     | 1/1/2006                | Carimune NF*,<br>Gammagard S/D | (human), lyophilized,<br>nanofiltered - Carimune NF<br>immune globulin intravenous<br>(human), solvent detergent<br>treated - Gammagard S/D | Carimune NF: Indicated for the maintenance treatment of patients with primary immunodeficiencies<br>(PID), e.g., common variable immunodeficiency, X-linked agammaglobulinemia, severe combined<br>immunodeficiency.<br>Gammagard S/D: Indicated for the treatment of Primary Immunodeficiency (PI) in adults and pediatric<br>patients two years of a ge or older, prevention of Daterial infections in hypoganmaglobulinemia and/or<br>recurrent bacterial infections associated with B-cell Chronic Lymphocytic Leukemia (CLL), prevention<br>and/or control of bieeding in adult Chronic Idiopathic Thrombocytopenic Purpura (ITP) patients and<br>prevention of coronary artery aneurysms associated with Kawasaki syndrome in pediatric patients. | 952   | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>Carimune NF:<br>- PID: None<br>- Gammagard S/D:<br>- PI: 2 years of age and older<br>- Chronic ITP: 38 years of age<br>and older<br>- Kawasaki Disease: None<br>- CLL: None   | 9/8/2021              |
| Immune<br>Globulins | J1568         | Injection, immune globulin,<br>(Octagam), intravenous, non-<br>lyophilized (e.g. liquid), 500<br>mg                                       | 500 mg                     | 1/1/2008                | Octagam®                       | immune globulin intravenous<br>(human) liquid solution for<br>intravenous administration  | Octagam 5%: Indicated for the treatment of primary humoral immunodeficiency.<br>Octagam 10%: Indicated for the treatment of:<br>• Chronic immune thrombocytopenic purpura (ITP) in adults.<br>• Dermatomyositis (DM) in adults.   | <ul> <li>Octagam 5%: 336<br/>units</li> <li>Octagam 10%: 1,120<br/>units</li> </ul> | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Product specific age<br>restrictions:<br>• Octagam 5%: 6 years of age<br>and older.<br>• Octagam 10%: 18 years of<br>age and older.   | 8/25/2021             |
| Immune<br>Globulins | J1569         | Injection, immune globulin,<br>(Gammagard liquid), non-<br>lyophilized, (e.g. liquid), 500<br>mg  | 500 mg                     | 1/1/2008                | Gammagard<br>Liquid            | immune globulin infusion<br>(human), 10% solution, for<br>intravenous and<br>subcutaneous administration                                    | Indicated as a:<br>- replacement therapy for primary humoral immunodeficiency (PI) in adult and pediatric patients two<br>years of age or older<br>- maintenance therapy to improve muscle strength and disability in adult patients with Multifocal Motor<br>Neuropathy (MMN)<br>- therapy to improve neuromuscular disability and impairment in adult patients with Chronic<br>Inflammatory Demyelinating Polyneuropathy (CIDP)<br>Limitations of Use:<br>- Safety and effectiveness of Gammagard Liquid has not been studied in immunoglobulin-naive patients<br>with CIDP.<br>- Gammagard Liquid maintenance therapy in CIDP has not been studied beyond 6 months.  | 840   | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• Pi 2 years and older<br>• MMN and CIDP: 18 years<br>and older   | 2/27/2024             |
| Drugs               | J1570         | Injection, ganciclovir sodium,<br>500 mg  | 500 mg                     | 1/1/2000                | Cytovene®-IV                   | ganciclovir sodium for<br>injection, for intravenous use  | Indicated for:<br>• Treatment of CMV retinitis in immunocompromised individuals, including patients with acquired<br>immunodeficiency syndrome (AIDS).<br>• Prevention of CMV disease in adult transplant recipients at risk for CMV disease.   | 104   | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 12/19/2022            |
| Immune<br>Globulins | J1571         | Injection, hepatitis B immune<br>globulin (Hepagam B),<br>intramuscular, 0.5 mL   | 0.5 mL                     | 1/1/2008                | Hepagam B®                     | hepatitis b immune globulin<br>intramuscular (human)  | Indicated for post exposure prophylaxis in the following settings:<br>• Acute Exposure to Blood Containing HBsAg<br>• Perinatal Exposure of Infants Bont to HBsAg-positive Mothers<br>• Sexual Exposure to HBsAg-positive Persons<br>+ Household Exposure to Persons with Acute HBV Infection   | 34  | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 9/12/2018             |
| Immune<br>Globulins | J1572         | Injection, immune globulin,<br>(Flebogamma/Flebogamma<br>DIF), intravenous, non-<br>lyophilized (e.g. liquid), 500<br>mg                  | 500 mg                     | 1/1/2008                | Flebogamma*                    | immune globulin intravenous<br>(human) for intravenous<br>administration, 10% liquid<br>preparation   | Indicated for the treatment of:<br>• Primary (inherited) Immunodeficiency (PI).<br>• Chronic Primary Immune Thrombocytopenia (ITP) in patients 2 years of age and older.  | 560   | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Primary (inherited)<br>Immunodeficiency (PI): None<br>• Chronic Primary Immune<br>Thrombocytopenia (TP): In<br>patients 2 years of age and<br>older.  | 7/3/2018              |
| Immune<br>Globulins | J1573         | Injection, hepatitis B immune<br>globulin (Hepagam B),<br>intravenous, 0.5 mL   | 0.5 mL                     | 1/1/2008                | HepaGam B <sup>®</sup>         | hepatitis b immune globulin<br>intravenous (human)  | Indicated for the prevention of hepatitis B virus recurrence after liver transplantation in HBsAg-positive<br>transplant patients (HepaGam B) – IV only.  | 1,290   | N/A   | N/A         | N/A                    | у               | Y                               |   | 7/3/2018              |

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| medicaid/medi       | caid-ncci-eo  | dit-files  |                            |                         |               |  |  |                                   |   |             |                        |                 |                                 |  |                       |
|---------------------|---------------|--|----------------------------|-------------------------|---------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category            | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name    | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs               | J1574         | Injection, ganciclovir sodium<br>(exela), not therapeutically<br>equivalent to j1570, 500 mg                   | 500 mg                     | 1/1/2023                | Ganzyk-RTU    | ganciclovir injection, for<br>intravenous use (Exela)  | Indicated for the:<br>• Treatment of CMV retinitis in immunocompromised adult patients, including patients with acquired<br>immunodeficiency syndrome (AIDS).<br>• Prevention of CMV disease in adult transplant recipients at risk for CMV disease.   | 104                               | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 12/6/2022             |
| Immune<br>Globulins | J1575         | Injection, immune<br>globulin/hyaluronidase,<br>(Hyqvia), 100 mg immune<br>globulin                            | 100 mg                     | 1/1/2016                | HyQvia®       | immune globulin infusion<br>10% (human) with<br>recombinant human<br>hyaluronidase solution for<br>subcutaneous administration | Indicated for treatment of primary immunodeficiency (PI) in patients two years of age and older.<br>Indicated for the treatment of maintenance therapy in adults with chronic inflammatory demyelinating<br>polyneuropathy (CIDP).   | 1,300                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication Specific Age<br>Restrictions:<br>PI: 2 years of age and older<br>CIDP: 18 years of age and<br>older   | 2/27/2024             |
| Immune<br>Globulins | J1576         | Injection, immune globulin<br>(panzyga), intravenous, non-<br>lyophilized (e.g., liquid), 500<br>mg            | 500 mg                     | 7/1/2023                | Panzyga*      | immune globulin intravenou:<br>human - ifas  | Chronic immune thrombocytopenia (ITP) in adults.     Chronic inflammatory demyelinating polyneuropathy (CIDP) in adults.   | 1,120                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Primary humoral<br>immunodeficiency (PI) - 2<br>years of age and older<br>• Chronic immune<br>thrombocytopenia (ITP) and<br>chronic inflammatory<br>demyelinating polyneuropath<br>(CIDP) - 18 years of age and<br>older | 6/22/2023<br>Y        |
| Drugs               | J1580         | Injection, garamycin,<br>gentamicin, up to 80 mg   | up to 80 mg                | 1/1/2000                | Garamycin®    | gentamicin sulfate injection<br>for intravenous infusion or<br>intramuscular injection   | administered as initial therapy in conjunction with a penicillin-type or cephalosporin-type drug before<br>obtaining results of susceptibility testing. If an aerobic organisma are suspected as etiologic agents,<br>consideration should be given to using other suitable antimicrobial therapy in conjunction with<br>gentamicin. Following identification of the organisma and its susceptibility, appropriate antibiotic therapy<br>should then be continued.<br>• Gentamicin sulfate has been used effectively in combination with carbenicillin for the treatment of life-<br>threatening infections caused by Pseudomonas areuginosa. It has also been found effective when used in<br>conjunction with a penicillin-type drug for the treatment of endocarditis caused by group D<br>streptococci.<br>• Gentamicin has also been shown to be effective in the treatment of serious staphylococcal infections.<br>While not the antibiotic of first choice, gentamicin may be considered when penicillins or other less<br>potentially toxic drugs are considered in mixed infections caused by susceptibility tests and clinical judgment indicate<br>its use. It may also be considered in mixed infections caused by susceptibile strains of staphylococci and<br>gram-negative canganisms. | 279                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/4/2019              |
| Biologicals         | J1602         | Injection, golimumab, 1 mg,<br>for intravenous use   | 1 mg                       | 1/1/2014                | Simponi Aria® | golimumab injection, for<br>intravenous use  | Indicated for treatment of a dult patients with:<br>Moderately osereely active Rheumatoid Arthritis (RA) in combination with methotrexate.<br>• Active Ankylosing Spondylitis (AS).<br>Indicated for treatment in patients 2 years of age and older with:<br>• Active Psoniatic Arthritis (PA).<br>• Active polyarticular Juvenile Idiopathic Arthritis (pJIA)   | 560                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | ¥               | Ŷ                               | Indication specific age<br>restrictions:<br>Rheumatoid Arthritis and<br>Ankylosing Spondylitis: 18<br>years of age and older<br>Polyarticular Juvenile<br>Idiopathic Arthritis: 2 years of<br>age and older  | 10/21/2020            |
| Drugs               | J1610         | Injection, glucagon<br>hydrochloride, per 1 mg   | 1 mg                       | 1/1/2000                | GlucaGen®     | glucagon for injection, for<br>subcutaneous, intramuscular<br>or intravenous use   | Indicated for:<br>• Treatment of severe hypoglycemia.<br>• Use as a diagnostic aid for use during radiologic examinations to temporarily inhibit movement of the<br>gastrointestinal tract.  | 10                                | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Treatment of severe<br>hypoglycemia: None<br>• Diagnostic aid: 18 years of<br>age and old  | 10/26/2018            |
| Drugs               | J1611         | Injection, glucagon<br>hydrochloride (fresenius kabi),<br>not therapeutically equivalent<br>to j1610, per 1 mg | 1 mg                       | 1/1/2023                | N/A           | glucagon for injection, for<br>subcutaneous, intramuscula<br>or intravenous use (Freseniu<br>Kabi)                             | Indicated:<br>• for the treatment of severe hypoglycemia in pediatric and adult patients with diabetes<br>• as a diagnostic aid for use during radiologic examinations to temporarily inhibit movement of the<br>gastrointestinal tract in adult patients  | 10                                | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>• Diagnostic aid during<br>radiologic examinations to<br>temporarily inhibit movemen<br>of the gastrointestinal tract:<br>18 years of age and older<br>• Treatment of severe<br>hypoglycemia: N/A                          | t 12/12/2022          |
| Drugs               | J1626         | Injection, granisetron<br>hydrochloride, 100 mcg   | 100 mcg                    | 1/1/2000                | N/A           | granisetron hydrochloride<br>injection, for intravenous us   | Indicated for:<br>• Prevention of nausea and/or vomiting associated with initial and repeat courses of emetogenic cancer<br>therapy including high-dose cisplatin.<br>• Prevention and treatment of postoperative nausea and vomiting in adults.   | 294                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific:<br>• Chemotherapy Induced<br>Nausea and Vomiting: 2 year:<br>of age and older<br>• Postoperative Nausea and<br>Vomiting: 18 years of age and<br>older   | 6/4/2019              |

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|---------------|---------------|--|----------------------------|-------------------------|----------------------|--|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name           | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
| Drugs         | J1627         | Injection, granisetron,<br>extended-release, 0.1 mg  | 0.1 mg                     | 1/1/2018                | Sustol®              | granisetron extended-release<br>injection, for subcutaneous<br>use           | Indicated in combination with other antiemetics in adults for the prevention of acute and delayed nausea<br>and vomiting associated with initial and repeat courses of moderately emetogenic chemotherapy (MEC) or<br>anthracycline and cyclophosphamide (AC) combination chemotherapy regimens  | 500                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 10/26/2018            |
| Drugs         | J1630         | Injection, haloperidol, up to 5<br>mg  | up to 5 mg                 | 1/1/2000                | Haldol®              | haloperidol lactate injection  | Indicated for use in the treatment of schizophrenia and for the control of tics and vocal utterances of<br>Tourette's Disorder.  | 124                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 10/26/2018            |
| Drugs         | J1631         | Injection, haloperidol<br>decanoate, per 50 mg   | per 50 mg                  | 1/1/2000                | Haldol®<br>Decanoate | haloperidol decanoate<br>injection, for intramuscular<br>use                 | Tourter's obsorver.<br>Indicated for the treatment of schizophrenic patients who require prolonged parenteral antipsychotic<br>therapy.  | 18                                | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/4/2019              |
| Drugs         | J1640         | Injection, hemin, 1 mg   | 1 mg                       | 1/1/2006                | Panhematin®          | hemin for injection  | Indicated for amelioration of recurrent attacks of acute intermittent porphyria temporally related to the<br>menstrual cycle in susceptible women, after initial carbohydrate therapy is known or suspected to be<br>inadequate.<br>Umitations of Use:<br>• Before administering Panhematin, consider an appropriate period of carbohydrate loading (i.e., 400 g<br>glucose/day for 1 to 2 days).<br>• Panhematin is not effective in repairing neuronal damage due to progression of porphyria attacks.   | 14,700                            | 16 years    | N/A         | Females Only           | Y               | Y                               |          | 11/30/2021            |
| Drugs         | J1643         | Injection, heparin sodium<br>(pfizer), not therapeutically<br>equivalent to j1644, per 1000<br>units | 1,000 units                | 1/1/2023                | N/A                  | heparin sodium injection, for<br>intravenous or subcutaneous<br>use (Pfizer) | Indicated for:   | 465                               | N/A         | N/A         | N/A                    | Ŷ               | Ŷ                               |          | 12/12/2022            |
| Drugs         | J1644         | Injection, heparin sodium, per<br>1,000 units  | per 1,000 units            | 1/1/2000                | N/A                  | heparin sodium injection, for<br>intravenous or subcutaneous<br>use          |  | 465                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/4/2019              |
| Drugs         | J1645         | Injection, dalteparin sodium,<br>per 2,500 IU  | per 2,500 IU               | 1/1/2000                | Fragmin®             | dalteparin sodium injection,<br>for subcutaneous use                         | Indicated for:<br>• Prophylaxis of ischemic complications of unstable angina and non-Q-wave myocardial infarction.<br>• Prophylaxis of deep vein thrombosis (DVT) in abdominal surgery, hip replacement surgery or medical<br>patients with severely restricted mobility during acute illness.<br>• Extended treatment of symptomatic venous thromboembolism (VTE) to reduce the recurrence in<br>patients with cancer. In these patients, the Fragmin therapy begins with the initial VTE treatment and<br>continues for six months.<br>• Treatment of symptomatic venous thromboembolism (VTE) to reduce the recurrence in pediatric<br>patients under some some some some some some some some | 372                               | 1 month     | N/A         | N/A                    | Y               | ¥                               |          | 6/4/2019              |
| Drugs         | J1650         | Injection, enoxaparin sodium,<br>10 mg   | 10 mg                      | 1/1/2000                | Lovenox®             | enoxaparin sodium injection,<br>for subcutaneous and<br>intravenous use      | Indicated for:<br>• Prophylaxis of deep vein thrombosis (DVT) in abdominal surgery, hip replacement surgery, knee<br>replacement surgery, or medical patients with severely restricted mobility during acute illness.<br>• Inpatient treatment of acute DVT without pulmonary embolism.<br>• Outpatient treatment of acute DVT without pulmonary embolism.<br>• Prophylaxis of ischemic complications of unstable angina and non-Q-wave myocardial infarction (MI).<br>• Treatment of acute ST-segment elevation myocardial infarction (STEMI) managed medically or with<br>subsequent percutaneous coronary intervention (PCI).   | 930                               | 18 years    | N/A         | N/A                    | Y               | Ŷ                               |          | 6/5/2019              |
| Drugs         | J1652         | Injection, fondaparinux<br>sodium, 0.5 mg  | 0.5 mg                     | 1/1/2003                | Arixtra®             | fondaparinux sodium<br>injection solution for<br>subcutaneous injection      | Indicated for:<br>• Prophylaxis of deep vein thrombosis (DVT) in patients undergoing hip fracture surgery (including<br>extended prophylaxis), hip replacement surgery, knee replacement surgery, or abdominal surgery.<br>• Treatment of DVT or acute pulmonary embolism (PE) when administered in conjunction with Coumadin.   | 520                               | 18 years    | N/A         | N/A                    | Ŷ               | Ŷ                               |          | 10/10/2018            |

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| Drugs         | J1720         | Injection, hydrocortisone<br>sodium succinate, up to 100<br>mg                   | up to 100 mg               | 1/1/2000                | Solu-Cortef* | hydrocortisone sodium<br>succinate for injection, for<br>intravenous or intramuscular<br>administration | When oral therapy is not reasible, and the strength, dosage rorm, and route or administration or the drug<br>reasonably lend the preparation to the treatment of the condition, the intravenous or intramuscular use of<br>Solu-Cortel is indicated as follows:<br>Allergic States: Control of severe or incapacitating allergic conditions intractable to adequate trials of<br>conventional treatment in asthma, atopic dermatitis, contact dermatitis, drug hypersensitivity reactions,<br>serum sichness; transfusion reactions.<br>Dermatologic Diseases: Bullous dermatitis herpetiformis, exfoliative erythroderma, mycosis fungoides,<br>pemphigus, severe erythema multiforme (Stevens-Johnson syndrome).<br>E hodorine Disordes: Primary or secondary admonocritical insufficiency (hydrocortisone or cortisone is<br>the drug of choice; synthetic analogs may be used in conjunction with mineralocorticolds where<br>applicable, in infanzy, mineralocritical supplementations is of particular importance), comgenital adrenal<br>hyperplasia, hypercalcemia associated with cancer, nonsuppurative thyroldits.<br>Gastrointestinal Diseases: Totik the patient orea critical period of the disease in regional enteritis<br>anemia (Diamond Blackfan anemia), idiopathic thrombcyctopenic purpura in adults (Intravenous<br>administration only, intramuscular administration is contraindicated), pure red cell aplasia, select cases of<br>secondary thrombcyctopenia.<br>• Miscellaneous: Trichnois with neurologic or myocardial involvement, tuberculous meningits with<br>subarachnoid black or impending block when used concurrently with appropriate antituberculous<br>chemotherapy.<br>• Neoplastic Diseases: For the palliative management of leukemias and lymphomas.<br>• Nervous System: Acute exacerbations of multiple sclerosis; creebral deema associated with primary or<br>metastatic brain tumo, or cranistomy.<br>• Ophthalmic Diseases: Torkhonicomy.<br>• Ophthalmic Diseases: Torkhonicomy. | 155                               | N/A         | N/A         | N/A  | ¥               | Y                               |          | 6/28/2021             |
| Drugs         | J1729         | Injection,<br>hydroxyprogesterone<br>caproate, Not Otherwise<br>Specified, 10 mg | 10 mg                      | 1/1/2018                | N/A          | hydroxyprogesterone<br>caproate injection   | Indicated in non-pregnant women:<br>• For the treatment of advanced adenocarcinoma of the uterine corpus (Stage III or IV)<br>• In the management of amenorhea (primary and secondary) and abnormal uterine bleeding due to<br>hormonal imbalance in the absence of organic pathology, such as submucous fibroids or uterine cancer<br>• As a test for endogenous estrogen production and for the production of secretory endometrium and<br>desquamation.   | 3,100                             | N/A         | N/A         | Indicated only for<br>non-pregnant<br>women. | Y               | Y                               |          | 6/4/2019              |
| Drugs         | J1738         | Injection, meloxicam, 1 mg   | 1 mg                       | 10/1/2020               | Anjeso™      | meloxicam injection, for<br>intravenous use   | Indicated for use in adults for the management of moderate-to-severe pain, alone or in combination with<br>non-NSAID analgesics.<br>Umitation of Use:<br>Because of delayed onset of analgesia, Anjeso alone is not recommended for use when rapid onset of<br>analgesia is required.<br>Indicated for the treatment of osteoporosis in postmenopausal women.  | 930                               | 18 years    | N/A         | N/A  | Y               | Y                               |          | 9/21/2020             |
| Drugs         | J1740         | Injection, ibandronate sodium<br>1 mg  | <sup>1,</sup> 1 mg         | 1/1/2007                | Boniva®      | ibandronate injection, for<br>intravenous use   | Limitations of Use:<br>Optimal duration of use has not been determined. For patients at low-risk form fracture, consider drug<br>discontinuation after 2 to 5 years of use.  | 3                                 | 40 years    | N/A         | Females Only                                 | Y               | Y                               |          | 10/18/2018            |
| Drugs         | J1742         | Injection, ibutilide fumarate,<br>mg   | 1 1 mg                     | 1/1/2000                | Corvert®     | ibutilide fumarate injection,<br>for intravenous infusion   | Indicated for the rapid conversion of atrial fibrillation or atrial flutter of recent onset to sinus rhythm.<br>Patients with atrial arrhythmias of longer duration are less likely to respond to ibutilide. The effectiveness<br>of ibutilide has not been determined in patients with arrhythmias of more than 90 days in duration.  | 10                                | 18 years    | N/A         | N/A  | Y               | Y                               |          | 10/18/2018            |
| Drugs         | J1743         | Injection, idursulfase, 1 mg   | 1 mg                       | 1/1/2008                | Elaprase®    | idursulfase injection, for<br>intravenous use   | Indicated for patients with Hunter syndrome (Mucopolysaccharidosis II, MPS II). Elaprase has been shown<br>to improve walking capacity in patients 5 years and older. In patients 16 months to 5 years of age, no data<br>are available to demonstrate improvement in disease-related symptoms or long term clinical outcome;<br>however, treatment with Elaprase has reduced spleen volume similarly to that of adults and children 5<br>years of age and older. The safety and efficacy of Elaprase have not been established in pediatric patients<br>less than 16 months of age.   | 360                               | 16 months   | N/A         | N/A  | Y               | Y                               |          | 6/4/2019              |
| Biologicals   | J1744         | Injection, icatibant, 1 mg   | 1 mg                       | 1/1/2013                | Firazyr®     | icatibant injection, for<br>subcutaneous use  | Indicated for the treatment of acute attacks of hereditary angioedema (HAE).   | 2700                              | 18 years    | N/A         | N/A  | Y               | Y                               |          | 6/4/2019              |

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|-------------|---------------|--|----------------------------|-------------------------|-------------------------|---|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|---|----------------------|
| Biologicals | J1745         | Injection, infliximab, excludes<br>biosimilar, 10 mg | 10 mg                      | 1/1/2017                | Remicade*               | infliximab lyophilized<br>concentrate for injection, for<br>intravenous use | Indicated for:<br>- Crohn's Disease: reducing signs and symptoms and inducing and maintaining clinical remission in adult<br>- Archn's Disease: reducing signs and symptoms and inducing and maintaining clinical remission in adult<br>therapy and reducing the number of draining enterocutaneous and rectovaginal fisulars and maintaining<br>fistula course in adult patients with fisularing disease who have had an inadequate response to conventional<br>in pediatric patients with moderately to severely active disease who have had an inadequate response to<br>conventional therapy.<br>- Ulcerative Collis: reducing signs and symptoms, inducing and maintaining clinical remission<br>in pediatric patients with moderately to severely active disease who have had an inadequate response to<br>conventional therapy.<br>- Ulcerative Collis: reducing signs and symptoms, inducing and maintaining clinical remission<br>in pediatric patients with moderately to severely active disease who have had an inadequate response to<br>- Nethorabe that an inadequate response to conventional therapy.<br>- Pediatric Ulcerative Collis: reducing signs and symptoms and inducing and maintaining clinical remission<br>in pediatric patients with moderately to severely active disease who have had an inadequate response to<br>conventional therapy.<br>- Pediatric Ulcerative Collis: reducing signs and symptoms and inducing and maintaining clinical remission<br>in pediatric patients with moderately to severely active disease.<br>- Analytosing Spondylitis: reducing signs and symptoms in patients with moderately to severely<br>active disease.<br>- Analytosing Spondylitis: reducing signs and symptoms in patients with active disease.<br>- Parkitis: Arthitis: reducing signs and symptoms in patients with active disease.<br>- Parkitis: charkitis: reducing signs and symptoms in patients with active disease.<br>- Parkitis: charkitis: reducing signs and symptoms in patients with active disease.<br>- Parkitis: charkitis: reducing signs and symptoms in patients with therapies are medically liess<br>appropriate.<br>- ***Recom | 300                               | 6 years     | N/A         | N/A                    | ¥               | Ŷ                               | 5/2024: NC Suggested Max<br>Monthly Units updated to align<br>with MUE values effective<br>5/6/2024.                      | 7/29/2024            |
| Biologicals | J1746         | Injection, ibalizumab-uiyk, 10<br>mg                 | 10 mg                      | 1/1/2019                | Trogarzo™               | ibalizumab-uiyk injection, for<br>intravenous use                           | <ul> <li>Treatment of hidradenitis suppurativa (HS), severe, refractory<br/>Indicated for use in combination with other antiretroviral(s), for the treatment of human<br/>immunodeficiency virus type 1 (HIV-1) infection in heavily treatment-experienced adults with multidrug<br/>resistant HIV-1 infection failing their current antiretroviral regimen.</li> </ul>  | 360                               | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 2/16/2024            |
| Drugs       | J1750         | Injection, iron dextran, 50 mg                       | 50 mg                      | 1/1/2009                | INFeD*                  | iron dextran injection  | Indicated for treatment of patients with documented iron deficiency in whom oral administration is<br>unsatisfactory or impossible.  | 62                                | 4 months    | N/A         | N/A                    | Y               | Y                               |   | 10/26/2018           |
| Drugs       | J1756         | Injection, iron sucrose, 1 mg                        | 1 mg                       | 1/1/2003                | Venofer*                | iron sucrose injection for<br>intravenous use                               | Indicated for the treatment of iron deficiency anemia in patients with chronic kidney disease (CKD).   | 2,000                             | 2 years     | N/A         | N/A                    | Y               | Ŷ                               |   | 7/29/2020            |
| Biologicals | J1786         | Injection, imiglucerase, 10<br>units                 | 10 units                   | 1/1/2011                | Cerezyme*               | imiglucerase for injection, for<br>intravenous use                          | Indicated for long-term enzyme replacement therapy for pediatric and adult patients with a confirmed<br>diagnosis of Type 1 Gaucher disease that results in one or more of the following conditions:   | 2,520                             | 2 years     | N/A         | N/A                    | Y               | Y                               |   | 6/22/2023            |
| Drugs       | J1790         | Injection, droperidol, up to 5 mg                    | up to 5 mg                 | 1/1/2000                | N/A                     | droperidol injection for<br>intravenous or intramuscular<br>use             | Indicated to reduce nausea and vomiting associated with surgical and diagnostic procedures.  | 5                                 | 2 years     | N/A         | N/A                    | Y               | Ŷ                               |   | 10/4/2018            |
| Drugs       | J1800         | Injection, propranolol HCl, up<br>to 1 mg            | up to 1 mg                 | 1/1/2000                | N/A                     | propranolol hydrochloride<br>injection, solution                            | Indicated for supraventricular arrhythmias, ventricular tachycardias, tachyarrhythmias of digitalis<br>intoxication and resistant tachyarrhythmias due to excessive catecholamine action during anesthesia.  | N/A                               | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 8/29/2018            |
| Biologicals | J1812         | Insulin (fiasp), per 5 units                         | 5 units                    | 7/1/2023                | Fiasp®                  | insulin aspart injection for<br>subcutaneous or intravenous<br>use          | Indicated to improve glycemic control in adult and pediatric patients with diabetes mellitus.  | N/A                               | 2 years     | N/A         | N/A                    | Y               | Y                               |   | 6/19/2023            |
| Biologicals | J1814         | Insulin (lyumjev), per 5 units                       | 5 units                    | 7/1/2023                | Lyumjev®                | insulin lispro-aabc injection,<br>for subcutaneous or<br>intravenous use    | Indicated to improve glycemic control in adult and pediatric patients with diabetes mellitus.  | N/A                               | 1 year      | N/A         | N/A                    | Y               | Ŷ                               |   | 6/19/2023            |
| Biologicals | J1815         | Injection, insulin, per 5 units                      | 5 units                    | 1/1/2003                | Various brand<br>names  | insulin, injectable suspension  | Indicated to improve glycemic control in adults and pediatric patients with diabetes mellitus.   | N/A                               | N/A         | N/A         | N/A                    | Ŷ               | Ŷ                               | 6/2024: NC Suggested Max<br>Monthly Units updated to align<br>with NCTracks, which has been<br>set to N/A since 1/1/2023. |                      |
| Biologicals | J1823         | Injection, inebilizumab-cdon, 1<br>mg                | 1 mg                       | 1/1/2021                | Uplizna™                | inebilizumab-cdon injection,<br>for intravenous use                         | Indicated for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult patients who are<br>anti-aquaporin-4 (AQP4) antibody positive.  | 600                               | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 12/28/2020           |
| Biologicals | J1830         | Injection, interferon beta-1B,<br>0.25 mg            | 0.25 mg                    | 1/1/2000                | Betaseron®,<br>Extavia® | interferon beta-1b for<br>injection, for subcutaneous<br>use                | Indicated for the treatment of relapsing forms of multiple sclerosis to reduce the frequency of clinical<br>exacerbations. Patients with multiple sclerosis in whom efficacy has been demonstrated include patients<br>who have experienced a first clinical episode and have MRI features consistent with multiple sclerosis.   | 16                                | 18 years    | N/A         | N/A                    | Y               | Ŷ                               |   | 6/4/2019             |
| Drugs       | J1833         | Injection, isavuconazonium<br>sulfate, 1 mg          | 1 mg                       | 1/1/2016                | Cresemba*               | isavuconazonium sulfate for<br>injection for intravenous<br>administration  | Indicated in adults and pediatric patients 1 year of age and older for the treatment of:   | 13,020                            | 1 year      | N/A         | N/A                    | Y               | Ŷ                               |   | 2/16/2024            |
| Drugs       | J1885         | Injection, ketorolac<br>tromethamine, per 15 mg      | 15 mg                      | 1/1/2000                | N/A                     | ketorolac tromethamine<br>injection for intravenous or<br>intramuscular use | Indicated for the short-term management (< 5 days) of moderately-severe acute pain requiring analgesia<br>at the opioid level in adults, usually in a postoperative setting.   | 40                                | 17 years    | N/A         | N/A                    | Y               | Y                               |   | 4/9/2019             |
| Drugs       | J1930         | Injection, lanreotide, 1 mg                          | 1 mg                       | 1/1/2009                | Somatuline®<br>Depot    | lanreotide injection, for<br>subcutaneous use                               | Indicated for the long-term treatment of acromegalic patients who have had an inadequate response to or<br>cannot be treated with surgery and/or radiotherapy.<br>indicated for the treatment of adult patients with unresectable, well- or moderately-differentiated, locally<br>advanced or metastatic gastoenteropanceratic neuroendocrine tumors (GEP-NETs) to improve progression<br>free survival.<br>Indicated for the treatment of adults with carcinoid syndrome; when used, it reduces the frequency of<br>short-acting somotistatin analogue rescue therapy.  | 240                               | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 10/26/2018           |

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| medicaid/medi | icaid-ncci-ed | dit-files  |                            |                         |   |  |   |                                   |   |             |  |                 |                                 |  |                      |
|---------------|---------------|--|----------------------------|-------------------------|---|--|---|-----------------------------------|---|-------------|--|-----------------|---------------------------------|--|----------------------|
| Category      | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                              | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions                                     | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modifie<br>Date |
| Biologicals   | J1931         | Injection, laronidase, 0.1 mg  | 0.1 mg                     | 1/1/2005                | Aldurazyme®                             | laronidase solution for<br>intravenous infusion only                                     | Indicated for patients with Hurler and Hurler-Scheie forms of Mucopolysaccharidosis I (MPS I) and for<br>patients with the Scheie form who have moderate to severe symptoms. The risks and benefits of treating<br>mildly affected patients with the Scheie form have not been established. Addurazyme has been shown to<br>improve pulmonary function and walking capacity. Addurazyme has not been evaluated for effects on the<br>central nervous system manifestations of the disorder.   | 4,060                             | 6 months  | N/A         | N/A  | Y               | Y                               |  | 4/10/2019            |
| Drugs         | J1932         | Injection, lanreotide, (cipla), 1<br>mg                                      | 1 mg                       | 10/1/2022               | N/A                                     | lanreotide injection, for<br>subcutaneous use (Cipla)                                    | Indicated for:<br>• The long-term treatment of acromegalic patients who have had an inadequate response to or cannot be<br>treated with surgery and/or radiotherapy.<br>• The treatment of adult patients with unresectable, well- or moderately differentiated, locally advanced<br>or metastatic gastroenteropancreatic neuroendocrine tumors (GEP-NETs) to improve progression-free<br>survival.   | 240                               | 18 years  | N/A         | N/A  | Y               | Y                               |  | 9/15/2022            |
| Drugs         | J1940         | Injection, furosemide, up to 20<br>mg  | up to 20 mg                | 1/1/2000                | Lasix®                                  | furosemide injection   | Indicated for the treatment of edema associated with congestive heart failure, cirrhosis of the liver, and<br>renal disease, including the nephrotic syndrome.<br>- Furosomide is particularly useful when an agent with greater diuretic potential is desired.<br>Indicated as an adjunct in the treatment of pulmonary edema.<br>- The intravenous administration of furosemide is indicated when a rapid onset of diuresis is desired.<br>If a strointestinal absorption is impaired or oral medication is not practical for any reason, furosemide is<br>indicated by the intravenous or intramuscular route. Parenteral use should be replaced with oral<br>furosemide as soon as practical. | 310                               | N/A   | N/A         | N/A  | Y               | ¥                               |  | 10/26/2018           |
| Drugs         | J1943         | Injection, aripiprazole lauroxil,<br>(aristada initio), 1 mg                 | 1 mg                       | 10/1/2019               | Aristada Initio™                        | aripiprazole lauroxil extended<br>release injectable suspension<br>for intramuscular use | Indicated for the initiation of Aristada when used for the treatment of schizophrenia in adults in  | 675                               | 18 years  | N/A         | N/A  | Y               | Ŷ                               | <ul> <li>Cervical Dystonia: Safety and<br/>effectiveness in pediatric<br/>patients have not been<br/>established.</li> </ul>   | 9/27/2019            |
| Drugs         | J1944         | Injection, aripiprazole lauroxil,<br>(aristada), 1 mg                        | 1 mg                       | 10/1/2019               | Aristada*                               | aripiprazole lauroxil extended<br>release injectable suspension<br>for intramuscular use | Indicated for the treatment of schizophrenia.   | 1,064                             | 18 years  | 65 years    | N/A  | Y               | Y                               |  | 9/27/2019            |
| Drugs         | J1950         | Injection, leuprolide acetate<br>(for depot suspension), per<br>3.75 mg      | per 3.75 mg                | 1/1/2000                | Lupron Depot*,<br>Lupron Depot-<br>PED* | leuprolide acetate for depot<br>suspension, for intramuscular<br>use                     |   | 12                                | Product Specific Age<br>Restrictions (see<br>comments)    | N/A         | Lupron Depot:<br>Females Only<br>Lupron Depot-<br>PED: N/A | ¥               | ¥                               | Product specific age<br>restrictions:<br>Lupron Depot:<br>Females of reproductive age<br>Lupron Depot-PED:<br>1 year of age and older  | 2/19/2024            |
| Drugs         | J1951         | Injection, leuprolide acetate<br>for depot suspension<br>(fensolvi), 0.25 mg | 0.25 mg                    | 7/1/2021                | Fensolvi®                               | leuprolide acetate for<br>injectable suspension, for<br>subcutaneous use                 | Indicated for the treatment of pediatric patients 2 years of age and older with central precocious puberty.   | 180                               | 2 years   | N/A         | N/A  | Y               | Y                               |  | 6/28/2021            |
| Drugs         | J1952         | Leuprolide injectable, camcevi,<br>1 mg                                      | 1 mg                       | 1/1/2022                | Camcevi™                                | leuprolide injectable<br>emulsion, for subcutaneous<br>use                               | Indicated for the treatment of adult patients with advanced prostate cancer.  | 42                                | 18 years  | N/A         | Males Only   | Y               | Ŷ                               |  | 5/16/2022            |
| Drugs         | J1953         | Injection, levetiracetam, 10<br>mg   | 10 mg                      | 1/1/2009                | Keppra®                                 | levetiracetam injection, for<br>intravenous use  | Indicated as an adjunctive therapy, as an alternative when oral administration is temporarily not feasible,<br>for the treatment of:<br>• Partial ones setures in patients 1 month of age and older with epilepsy<br>• Myoclonic setures in patients 12 years of age and older with juvenile myoclonic epilepsy<br>• Primary generalized tonic-clonic seizures in patients 6 years of age and older with idiopathic generalized<br>epilepsy   | 9,300                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A  | Ŷ               | Ŷ                               | Indication specific age<br>restrictions:<br>Partial Onset Seizures: 1<br>month of age and older<br>• Myoclonic Seizures in<br>Patients with Juvenile<br>Myoclonic Epilepsy: 12 years<br>of age and older<br>• Primary Generalized Tonic-<br>Clonic Seizures: 6 years of age<br>and older | 10/10/2018           |
| Drugs         | J1954         | Injection, leuprolide acetate<br>for depot suspension (lutrate),<br>7.5 mg   | 7.5 mg                     | 1/1/2023                | Lutrate Depot                           | leuprolide acetate for depot<br>suspension   | Indicated for treatment of advanced prostate cancer.  | 3                                 | 18 years  | N/A         | Males Only   | Y               | Y                               |  | 3/16/2023            |
| Drugs         | J1955         | Injection, levocarnitine, per 1 g  | 1 g                        | 1/1/2000                | Carnitor*                               | levocarnitine injection for<br>intravenous use   | Indicated for:<br>• the acute and chronic treatment of patients with an inborn error of metabolism which results in<br>secondary carnitine deficiency.<br>• the prevention and treatment of carnitine deficiency in patients with end stage renal disease who are<br>undergoing dialysis.   | 1,302                             | N/A   | N/A         | N/A  | Y               | Y                               |  | 4/10/2019            |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
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| ategory | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name            | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age               | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modifi<br>Date |
|---------|---------------|---|----------------------------|-------------------------|-----------------------|---|---|-----------------------------------|---|---------------------------|------------------------|-----------------|---------------------------------|---|---------------------|
| Drugs   | J1956         | Injection, levofloxacin, 250 mg   | 250 mg                     | 1/1/2000                | Levaquin <sup>®</sup> | levofloxacin injection for<br>intravenous use   | Indicated in adults (>=18 years of age) with infections caused by designated, susceptible bacteria:<br>• Pneumonia: Noscoomial and Community Acquired<br>• Skin and Skin Structure Infections: Complicated and Uncomplicated<br>• Chronic bacterial prostatitis<br>• Inhalational Anthrax, Post-Exposure<br>• Plague<br>• Urinary Tract Infections: Complicated and Uncomplicated<br>• Acute Pyelonephritis<br>• Acute Pyelonephritis<br>• Acute Bacterial Exacerbation of Chronic Bronchitis<br>• Acute Bacterial Sinusitis<br>Usage: To reduce the development of drug-resistant bacteria and maintain the effectiveness of Levaquin<br>and other antibacterial drugs, Levaquin should be used only to treat or prevent infections that are proven  | 62                                | Indication Specific<br>Age Restrictions<br>(see comments) | N/A                       | N/A                    | Y               | Y                               | Indication specific:<br>Inhalation Anthrax (Post-<br>Exposure): 6 months and<br>older.<br>Plague: 6 months and older.<br>All other indications: 18 years<br>of age and older. | 6/5/2019            |
| Drugs   | J1961         | Injection, lenacapavir, 1 mg  | 1 mg                       | 7/1/2023                | Sunlenca®             | lenacapavir injection, for subcutaneous use   | or strongly suspected to be caused by bacteria.<br>Indicated for the treatment of HIV-1 infection in heavily treatment-experienced adults with multidrug<br>resistant HIV-1 infection failing their current antiretroviral regimen due to resistance, intolerance, or<br>safety considerations.   | 927                               | 18 years  | N/A                       | N/A                    | Y               | Y                               |   | 6/22/202            |
| Drugs   | J2001         | Injection, lidocaine HCL for<br>intravenous infusion, 10 mg                           | 10 mg                      | 1/1/2004                | N/A                   | lidocaine hydrochloride<br>injection, solution  | Administered intravenously or intramuscularly, is specifically indicated in the acute management of<br>ventricular arrhythmias such as those occurring in relation to acute myocardial infarction, or during<br>cardiae manipulation, such as cardiae surgery.<br>I indicated for production of local or regional anesthesia by infiltration techniques such as percutaneous<br>injection and intravenous regional anesthesia by peripheral nerve block techniques such as brachial plexus<br>and intercostal and by central neural techniques such as lumbar and caudal epidural blocks, when the<br>accepted procedures for these techniques as described in standard textbooks are observed.   | 35                                | N/A   | N/A                       | N/A                    | Y               | Ŷ                               |   | 10/31/20:           |
| Drugs   | J2010         | Injection, lincomycin HCl, up<br>to 300 mg  | 300 mg                     | 1/1/2000                | Lincocin®             | lincomycin hydrochloride<br>injection, solution   | Indicated for the treatment of serious infections due to susceptible strains of streptococci, pneumococci,<br>and staphylococci. Its use should be reserved for penicillin-allergic patients or other patients for whom, in<br>the judgment of the physician, a penicillin is inappropriate.  | 837                               | 1 month   | N/A                       | N/A                    | Y               | Y                               |   | 10/26/201           |
| Drugs   | J2020         | Injection, linezolid, 200 mg  | 200 mg                     | 1/1/2002                | Ζγνοχ®                | linezolid injection, solution   | Indicated in adults and children for the treatment of the following infections caused by susceptible Gram-<br>positive bacteria: nosocomial pneumonia; community-acquired pneumonia, complicated skin and skin<br>structure infections, including diabetic foot infections, without concomitant osteomyelitis, uncomplicated<br>skin and skin structure infections, vancomycin-resistant Enterococcus faecium infections.<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Zyvox<br>formulations and other antibacterial drugs, Zyvos should be used only to treat infections that are proven  | 168                               | N/A   | N/A                       | N/A                    | Ŷ               | Y                               |   | 10/26/201           |
| Drugs   | J2021         | Injection, linezolid (hospira),<br>not therapeutically equivalent<br>to j2020, 200 mg | 200 mg                     | 1/1/2023                | N/A                   | linezolid injection, for<br>intravenous use (Hospira)   | Indicated in adults and children for the treatment of the following infections caused by susceptible Gram-<br>positive bacteria: Noscomial pneumonia; Community-acquired pneumonia; Complicated skin and skin<br>structure infections, including diabetic foot infections, without concomitant osteomyelitis; Vancomycin-<br>resistant Entrecoccus faecium infections.<br>Limitations of Use:   | 168                               | N/A   | N/A                       | N/A                    | Y               | Ŷ                               |   | 12/12/202           |
| Drugs   | J2060         | Injection, lorazepam, 2 mg  | 2 mg                       | 1/1/2000                | Ativan®               | lorazepam injection for<br>intravenous or intramuscular<br>use                                    | Indicated:<br>In adult patients for preanesthetic medication, producing sedation (sleepiness or drowsiness), relief of<br>anxiety and a decreased ability to recall events related to the day of surgery.<br>• For treatment of status epilepticus.   | 124                               | 18 years  | N/A                       | N/A                    | Ŷ               | Y                               |   | 4/10/201            |
| Drugs   | J2150         | Injection, mannitol, 25% in 50<br>mL  | 50 mL                      | 1/1/2000                | N/A                   | mannitol injection, for<br>intravenous use  | Indicated for the reduction of:<br>Indicated for the reduction of:<br>Indicated for the reduction of the re | 713                               | N/A   | N/A                       | N/A                    | Y               | ¥                               |   | 11/29/202           |
| Drugs   | J2175         | Injection, meperidine<br>hydrochloride, per 100 mg                                    | 100 mg                     | 1/1/2000                | Demerol™              | meperidine hydrochloride<br>injection, for subcutaneous,<br>intramuscular, and<br>intravenous use | Indicated for preoperative medication, support of anesthesia, obstetrical analgesia, and for the<br>management of pain severe enough to require an opioid analgesic and for which alternative treatments<br>are inadequate.<br>Limitations of Use:<br>Because of the risks of addiction, abuse, and misuse with opioids, even at recommended doses, reserve<br>for use in patients for whom alternative treatment options [e.g., non-opioid analgesics or opioid<br>combination products] have not been tolerated, or are not expected to be tolerated or have not provided<br>adequate analgesia, or are not expected to provide adequate analgesia.   | 124                               | N/A   | N/A                       | N/A                    | Ŷ               | Ŷ                               |   | 10/26/201           |
| Drugs   | J2186         | Injection, meropenem and<br>vaborbactam, 10mg/10mg<br>(20mg)                          | 1 vial                     | 1/1/2019                | Vabomere™             | meropenem and<br>vaborbactam for injection,<br>for intravenous use                                | Indicated for the treatment of patients 18 years and older with complicated urinary tract infections (CUTI)<br>including pyelonephritis caused by designated susceptible bacteria. To reduce the development of drug-<br>resistant bacteria and maintain the effectiveness of Vabomere and other antibacterial drugs. Vabomere<br>should be used only to treat or prevent infections that are proven or strongly suspected to be caused by<br>susceptible bacteria.   | 8,400                             | 18 years  | N/A                       | N/A                    | Y               | Y                               |   | 10/26/20:           |
| Drugs   | J2210         | Injection, methylergonovine<br>maleate, up to 0.2 mg                                  | up to 0.2 mg               | 1/1/2000                | Methergine®           | methylergonovine maleate<br>injection   | Indicated   | 5                                 | Women of childbearing age                                 | Women of childbearing age | Females Only           | Ŷ               | Y                               |   | 10/31/20:           |
| Drugs   | J2249         | Injection, remimazolam, 1 mg  | 1 mg                       | 7/1/2023                | Byfavo™               | remimazolam for injection,<br>for intravenous use   | Indicated for the induction and maintenance of procedural sedation in adults undergoing procedures<br>lasting 30 minutes or less.   | 200                               | 18 years  | N/A                       | N/A                    | Y               | Y                               |   | 6/22/202            |

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|-------------|---------------|---|----------------------------|-------------------------|--------------------------------------|---|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Drugs       | J2250         | injection, midazolam<br>hydrochloride, per 1 mg   | 1 mg                       | 1/1/2000                | N/A                                  | midazolam hydrochloride<br>injection for intravenous or<br>intramuscular use                    | Indicated:<br>• Intramuscularly or intravenously for preoperative sedation/anxiolysis/amnesia<br>• Intravenously as an agent for sedation/anxiolysis/amnesia prior to or during diagnostic, therapeutic or<br>endoscopic procedures, such as bronchoscopy, gastrocoy, coronary anglography, cardiac<br>catheterization, oncology procedures, radiologic procedures, suture of lacerations and other procedures<br>either alow or in combination with other CNS depressants;<br>• Intravenously for induction of general anesthesia, before administration of other anesthetic agents. With<br>the use of narcotic premedication, induction of anesthesia can be attained within a relatively narrow dose<br>range and in a short period of time. Intravenous mislocation can also be used as a component of<br>intravenous supplementation of introus oxide and oxygen (balanced anesthesia);<br>• Continuous intravenous infusion for sedation of intubated and mechanically ventilated patients as a<br>component of anesthesia or during treatment in a critical care setting.  | 25                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 10/31/2018            |
| Drugs       | J2251         | Injection, midazolam<br>hydrochloride (wg critical<br>care), not therapeutically<br>equivalent to j2250, per 1 mg | 1 mg                       | 1/1/2023                | N/A                                  | midazolam in sodium<br>chloride injection for<br>intravenous use (WG Critical<br>Care)          | Indicated for:<br>Continuous intravenous infusion for sedation of intubated and mechanically ventilated adult, pediatric,<br>and neonatal patients as a component of anesthesia or during treatment in a critical care setting.  | 500                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 12/12/2022            |
| Drugs       | J2260         | Injection, milrinone lactate,<br>per 5 mg   | per 5 mg                   | 1/1/2000                | N/A                                  | milrinone lactate injection   | Indicated for the short-term intravenous treatment of patients with acute decompensated heart failure.   | 64                                | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/6/2019              |
| Biologicals | J2267         | Injection, mirikizumab-mrkz, 1<br>mg  | 1 mg                       | 7/1/2024                | Omvoh™                               | mirikizumab-mrkz injection,<br>for intravenous or<br>subcutaneous use                           | Mirikizumab-mrkz injection is indicated for the treatment of moderately to severely active ulcerative<br>colitis in adults.  | 600                               | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |          | 6/24/2024             |
| Drugs       | J2270         | Injection, morphine sulfate, up<br>to 10 mg   | up to 10 mg                | 1/1/2000                | N/A                                  | morphine sulfate injection,<br>up to 10 mg  | Indicated for the management of pain severe enough to require an opioid analgesic and for which<br>alternative treatments are inadequate.<br>Limitations of Use: Because of the risks of addiction, abuse, and misuse with opioids, even at<br>recommended doses, reserve Morphine Sulfate Injection, for use in patients for whom alternative<br>treatment opioines (e.g., non-opioid cambination products):<br>Have not been tolerated, or are not expected to be tolerated,<br>Have not provided adequate analgesia, or are not expected to provide adequate analgesia<br>Prior: Indicated for:<br>the relief of severe acute and chronic pain<br>to relieve preoperative apprehension<br>to facilitizet ansthesia induction<br>= the treatment of dyspines associated with acute left ventricular failure and pulmonary edema<br>= analgesia during labor<br>= analgesia during labor<br>= anstersia .   | 527                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/7/2019              |
| Drugs       | J2272         | Injection, morphine sulfate<br>(fresenius kabi), not<br>therapeutically equivalent to<br>j2270, up to 10 mg       | 10 mg                      | 1/1/2023                | N/A                                  | morphine sulfate injection,<br>for intravenous or<br>intramuscular use, CII<br>(Fresenius Kabi) | Indicated for the management of pain severe enough to require an opioid analgesic and for which<br>alternative treatments are inadequate.<br>Umitations of Use<br>Because of the risks of addiction, abuse, and misuse with opioids, even at recommended doses, reserve<br>Morphine Suffate Injection, for use in patients for whom alternative treatment options [e.g., non-opioid<br>analgesics or opioid combination products]:<br>• Have not been tolerated, or are not expected to be tolerated,<br>• Have not provide adequate analgesia, or are not expected to provide adequate analgesia  | 527                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 12/12/2022            |
| Drugs       | J2274         | Injection, morphine sulfate,<br>preservative-free for epidural<br>or intrathecal use, 10 mg                       | 10 mg                      | 1/1/2015                | Duramorph*,<br>Infumorph*,<br>Mitigo | morphine sulfate injection<br>preservative-free   | • Mitigo: for use in continuous microinfusion devices and indicated only for intrathecal or epidural infusion<br>in the management of intractable chronic pain severe enough to require an opioid analgesic and for which<br>alternative treatments are inadequate. • Infumorph: for use in continuous microinfusion devices and indicated only for intrathecal or epidural<br>infusion in the management of intractable chronic pain severe enough to require an opioid analgesic and<br>for which alternative treatments are inadequate. • Duramorph: Indicated for: • Othe management of pain severe enough to require use of an opioid analgesic by intravenous<br>administration and for which alternative treatments are not expected to be adequate. • Our enough the intrathecal management of pain without attendant loss of motor, sensory, or<br>sympathetic function. • Umitation of Use: Duramorph is not for use in continuous microinfusion devices. Prior to 10/30/2018: Morphine sulfate (preservative-free sterifie solution) is a systemic narcotic analgesic.<br>for administred equivally or intrathecal, provides pain neited for extended periods without attendant loss<br>of motor, sensor, or sympathetic function. Infumorph <sup>+</sup> is indicated only for intrathecal or epidural infusion in the treatment of intractable druging on intrathecal, provides pain neited for extended periods without attendant loss<br>of motor, sensor, or sympathetic function. Infumorph <sup>+</sup> is indicated only for intrathecal or epidural infusion in the treatment of intractable chronic<br>pain. It is not recommended for single-dose intravenous, intramuscular, or subcutaneous administration<br>due to the large amount of morphine in the ampule and the associated risk of overdosage. | 100                               | 18 years    | N/A         | N/A                    | Y               | ¥                               |          | 4/9/2022              |
| Drugs       | J2277         | Injection, motixafortide, 0.25  | 0.25 mg                    | 4/1/2024                | Aphexda™                             | motixafortide for injection,<br>for subcutaneous use  | Indicated in combination with filgrastim (G-CSF) to mobilize hematopoietic stem cells to the peripheral<br>blood for collection and subsequent autologous transplantation in patients with multiple myeloma.   | 1,488                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 3/22/2024             |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
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| neulcalu/meul | icaid-ncci-ed | ht-mes  |                            |                         | 1                                     |   |  | 1                                 |             | 1           |                        |                 | Rebating            |   | 1                     |
|---------------|---------------|---|----------------------------|-------------------------|---------------------------------------|---|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------|---|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                            | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Labeler<br>Required | Comments  | Last Modified<br>Date |
| Drugs         | J2278         | Injection, ziconotide, 1<br>microgram   | 1 mcg                      | 1/1/2006                | Prialt®                               | ziconotide solution,<br>intrathecal infusion  | Indicated for the management of severe chronic pain in patients for whom intrathecal therapy is<br>warranted, and who ein intolerant of or refractory to other treatment, such as systemic analgesics,<br>adjunctive therapies, or intrathecal morphine.   | 620                               | 18 years    | N/A         | N/A                    | Y               | Ŷ                   |   | 9/21/2018             |
|               |               |   |                            |                         |                                       |   | Indicated for management of pain severe enough to require an opioid analgesic and for which alternative<br>treatments are inadequate. Also can be used as a supplement to balanced anesthesia, for pre/post<br>operative analgesia and obstetrical analgesia during labor and delivery.  |                                   |             |             |                        |                 |                     |   |                       |
| Drugs         | J2300         | Injection, nalbuphine<br>hydrochloride, per 10 mg   | 10 mg                      | 1/1/2000                | N/A                                   | nalbuphine hydrochloride<br>injection, solution                                     | Limitations of Use: Because of the risks of addiction, abuse, and misuse with opioids, even at<br>recommended does, reserve nalbuphine injection for use in patients for whom alternative treatment<br>options (e.g. non-opioid analgesics). The second sec | 248                               | 18 years    | N/A         | N/A                    | Y               | Y                   |   | 10/26/2018            |
| Drugs         | J2310         | Injection, naloxone<br>hydrochloride, per 1 mg  | 1 mg                       | 1/1/2000                | Narcan®                               | naloxone hydrochloride<br>injection   | Indicated for the complete or partial reversal of opioid depression, including respiratory depression,<br>induced by natural and synthetic opioids including: propoxyphene, methadone, nalbuphine, butorphanol<br>and pentazocine; It is also indicated for the diagnosis of suspected opioid tolerance or acute opioid<br>overdose.   | N/A                               | N/A         | N/A         | N/A                    | Y               | Y                   |   | 10/26/2018            |
| Drugs         | J2311         | Injection, naloxone<br>hydrochloride (zimhi), 1 mg  | 1 mg                       | 1/1/2023                | Zimhi™                                | naloxone hydrochloride<br>injection for intramuscular or<br>subcutaneous use        | Indicated in adult and pediatric patients for the emergency treatment of known or suspected opioid<br>overdose, as manifested by respiratory and/or central nervous system depression.   | 50                                | N/A         | N/A         | N/A                    | Y               | Ŷ                   |   | 12/6/2022             |
| Drugs         | J2315         | Injection, naltrexone, depot<br>form, 1 mg  | 1 mg                       | 1/1/2007                | Vivitrol®                             | naltrexone for extended-<br>release injectable suspension,<br>for intramuscular use | <ul> <li>Indicated for the treatment of alcohol dependence in patients who are able to abstain from alcohol in an outpatient setting prior to initiation of treatment with Vivitrol. Patients should not be actively drinking at the time of initial Vivitrol administration.</li> <li>Indicated for the prevention of relapse to opioid dependence, following opioid detoxification.</li> <li>Vivitrol should be part of a comprehensive management program that includes psychosocial support.</li> </ul>  | 760                               | 18 years    | N/A         | N/A                    | Y               | Y                   | 9/1/2023: Generic Name<br>updated to align with<br>Prescribing Information. | 9/13/2023             |
| Biologicals   | J2323         | injection, natalizumab, 1 mg  | 1 mg                       | 1/1/2008                | Tysabri*                              | natalizumab injection, for<br>intravenous use                                       | Indicated for treatment of:<br>Multiple Scienosis (MS)<br>• Tysabri is indicated as monotherapy for the treatment of patients with relapsing forms of multiple<br>scienosis. Tysabri increases the risk of PML. When<br>initiating and continuing treatment with Tysabri, physicians should consider whether the expected benefit<br>of Tysabri is sindicated for inducing and maintaining clinical response and remission in adult patients with<br>moderately to severely active Crohr's disease with evidence of inflammation who have had an inadequate<br>response to, or are unable to tolerate, conventional CD therapies and inhibitors of TNF-α.<br>Important Limitations:<br>• In CD, Tysabri should not be used in combination with immunosuppressants or inhibitors of TNF-α.  | 600                               | 18 years    | N/A         | N/A                    | Y               | Ŷ                   |   | 10/26/2018            |
| Drugs         | J2326         | Injection, nusinersen, 0.1 mg   | 0.1 mg                     | 1/1/2018                | Spinraza®                             | nusinersen injection, for<br>intrathecal use  | Indicated for the treatment of spinal muscular atrophy (SMA) in pediatric and adult patients.  | 360                               | N/A         | N/A         | N/A                    | Y               | Y                   |   | 5/6/2021              |
| Biologicals   | J2327         | Injection, risankizumab-rzaa,<br>intravenous, 1 mg  | 1 mg                       | 1/1/2023                | Skyrizi®                              | risankizumab-rzaa injection,<br>for intravenous use                                 | Indicated for the treatment of:<br>• moderately to severely active cohn's disease in adults.<br>• moderately to severely active ucerative colitis (UC) in adults.  | 2,400                             | 18 years    | N/A         | N/A                    | Y               | Y                   |   | 7/29/2024             |
| Biologicals   | J2329         | Injection, ublituximab-xiiy,<br>1mg   | 1 mg                       | 7/1/2023                | Briumvi™                              | ublituximab-xiiy injection, for<br>intravenous use                                  | Indicated for the treatment of relapsing forms of multiple sclerosis (MS), to include clinically isolated<br>syndrome, relapsing-remitting disease, and active secondary progressive disease, in adults.   | 600                               | 18 years    | N/A         | N/A                    | Y               | Y                   |   | 6/22/2023             |
| Drugs         | J2353         | Injection, octreotide, depot<br>form for intramuscular<br>injection, 1 mg                         | 1 mg                       | 1/1/2004                | Sandostatin <sup>®</sup> LAR<br>Depot | octreotide acetate for<br>injectable suspension                                     | Indicated for treatment in patients who have responded to and tolerated sandostatin injection<br>subcutaneous injection for:<br>• Acromegaly<br>• Severe diarrihea/flushing episodes associated with metastatic carcinoid tumors<br>• Profuse watery diarrihea associated with VIP-secreting tumors  | 40                                | 18 years    | N/A         | N/A                    | Y               | Y                   |   | 7/16/2018             |
| Drugs         | J2354         | Injection, octreotide, non-<br>depot form for subcutaneous<br>or intravenous injection, 25<br>mcg | 25 mcg                     | 1/1/2004                | Sandostatin®                          | octreotide acetate, injection   | Indicated:<br>• To reduce blood levels of growth hormone and IGF-I (somatomedin C) in acromegaly patients who have<br>had inadequate response to or cannot be treated with surgical resection, pituitary irradiation, and<br>bromocriptine meylate at maximally tolerated docse.<br>• For the symptomatic treatment of patients with metastatic carcinoid tumors where it suppresses or<br>inhibits the severe diarnhea and flushing episodes associated with VH-serceting tumors. Sandostatin<br>studies were not designed to show an effect on the size, rate of growth or development of metastases.  | 1,860                             | 18 years    | N/A         | N/A                    | Y               | Ŷ                   |   | 7/16/2018             |
| Drugs         | J2358         | Injection, olanzapine, long-<br>acting, 1 mg  | 1 mg                       | 1/1/2011                | Zyprexa®<br>Relprevv™                 | olanzapine pamoate for<br>extended release injectable<br>suspension                 | Indicated for the treatment of schizophrenia.  | 900                               | 18 years    | N/A         | N/A                    | Y               | Y                   |   | 9/21/2018             |
| Drugs         | J2359         | Injection, olanzapine, 0.5 mg   | 0.5 mg                     | 10/1/2023               | Zyprexa®<br>Intramuscular             | olanzapine injection, powder,<br>for solution                                       | Indicated for the treatment of acute agitation associated with schizophrenia and bipolar I mania.  | 1,860                             | 13 years    | N/A         | N/A                    | Y               | Y                   |   | 9/28/2023             |
| Drugs         | J2360         | Injection, orphenadrine<br>citrate, up to 60 mg   | up to 60 mg                | 1/1/2000                | Norflex*                              | orphenadrine citrate injection  | Indicated as an adjunct to rest, physical therapy, and other measures for the relief of discomfort<br>associated with acute painful musculoskeletal conditions.  | 20                                | 18 years    | N/A         | N/A                    | Y               | Y                   |   | 7/16/2018             |

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|----------|---------------|--|----------------------------|-------------------------|---|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
|          |               |  |                            |                         |   |  | Multidose vial with preservatives: Indicated for the production of local anesthesia by infiltration and<br>peripheral nerve block.   |                                   |   |             |                        |                 | roquiou                         |  |                       |
| Drugs    | J2401         | Injection, chloroprocaine<br>hydrochloride, per 1 mg   | 1 mg                       | 1/1/2023                | Nesacaine <sup>®</sup> ,<br>Nesacaine <sup>®</sup> -MPF | chloroprocaine HCl injection   | Single dose vial without preservatives and without EDTA: Indicated for the production of local anesthesia<br>by infiltration, peripheral, and central nerve block, including lumbar and caudal epidural blocks.  | 1,000                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 12/6/2022             |
|          |               |  |                            |                         |   |  | Nesacaine and Nesacaine-MPF Injections are not to be used for subarachnoid administration.   |                                   |   |             |                        |                 |                                 |  |                       |
| Drugs    | J2402         | Injection, chloroprocaine<br>hydrochloride (clorotekal), per<br>1 mg                                 | 1 mg                       | 1/1/2023                | Clorotekal®   | chloroprocaine hydrochloride<br>injection, for intrathecal use   | Indicated for intrathecal injection in adults for the production of subarachnoid block (spinal anesthesia).  | 50                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/6/2022             |
| Drugs    | J2403         | Chloroprocaine hcl<br>ophthalmic, 3% gel, 1 mg   | 1 mg                       | 4/1/2023                | lheezo™   | chloroprocaine hydrochloride<br>ophthalmic gel 3%, for topical<br>ophthalmic use                                 | Chloroprocaine hydrochloride ophthalmic gel is indicated for ocular surface anesthesia.  | 4,000                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/1/2023             |
| Drugs    | J2405         | Injection, ondansetron<br>hydrochloride, per 1 mg  | 1 mg                       | 1/1/2000                | Zofran®   | ondansetron hydrochloride<br>injection, for intravenous or<br>intramuscular use                                  | Indicated for the prevention of:<br>• Nausea and vomiting associated with initial and repeat courses of emetogenic cancer chemotherapy.<br>• Postoperative nausea and/or vomiting.   | 720                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• Prevention of nausea and<br>vomiting associated with<br>emetogenic chemotherapy; 6<br>months of age and older<br>• Prevention of postoperative<br>nausea and vomiting: 1 month<br>of age and older | 9/27/2018             |
| Drugs    | J2406         | Injection, oritavancin<br>(kimyrsa), 10 mg   | 10 mg                      | 10/1/2021               | Kimyrsa™  | oritavancin for injection, for<br>intravenous use  | Indicated for the treatment of adult patients with acute bacterial skin and skin structure infections<br>(ABSSI) caused by susceptible isolates of the following Gram-positive microorganisms: Staphylococcus<br>aureus (including methicillin-susceptible and methicillin-resistant isolates). Streptococcus<br>Streptococcus agalactiae, Streptococcus dygalactiae, Streptococcus anginosus group (includes S<br>anginosus, S. intermedius, and S. constellatus), and Enterococcus faecalis (vancomycin-susceptible isolates<br>only).<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Kimyrsa and other   | 120                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/29/2021             |
|          |               |  |                            |                         |   |  | antibacterial drugs, Kimyrsa should be used only to treat or prevent infections that are proven or strongly suspected to be caused by bacteria.  |                                   |   |             |                        |                 |                                 |  |                       |
| Drugs    | J2407         | Injection, oritavancin<br>(orbactiv), 10 mg  | 10 mg                      | 10/1/2021               | Orbactiv®   | oritavancin for injection, for intravenous use   | projection to be caused by backeting<br>Indicated for the treatment of adult patients with acute bacterial skin and skin structure infections caused<br>or suspected to be caused by susceptible isolates of designated Gram-positive microorganisms.  | 120                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/29/2021             |
| Drugs    | J2425         | injection, palifermin, 50<br>micrograms  | 50 mcg                     | 1/1/2006                | Kepivance®  | palifermin injection, for<br>intravenous use   | Indicated to decrease the incidence and duration of severe oral mucositis in patients with hematologic<br>malignancies receiving myelotoxic therapy in the setting of autologous hematopoietic stem cell support.<br>Kepivance is indicated as supportive care for preparative regimens predicted to result in 2 WHO Grade 3<br>mucositis in the majority of patients.<br>Limitations of Use:<br>The safety and efficacy of Kepivance have not been established in patients with non-hematologic<br>malignancies.<br>• Kepivance was not effective in decreasing the incidence of severe mucositis in patients with hematologic<br>malignancies receiving myelotoxic therapy in the setting of allogeneic hematopoietic stem cell support.<br>• Kepivance is not recommended for use with melphalan 200 mg/m <sup>2</sup> as a conditioning regimen. | 1,008                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 4/9/2019              |
| Drugs    | J2426         | Injection, paliperidone<br>palmitate extended release<br>(invega sustenna), 1 mg                     | 1 mg                       | 1/1/2011                | Invega Sustenna®  | paliperidone palmitate<br>extended-release injectable<br>suspension, for intramuscular<br>use                    | Indicated for:<br>• Treatment of schizophrenia in adults.<br>• Treatment of schizopffettive disorder in adults as monotherapy and as an adjunct to mood stabilizers or<br>antideoressants.   | 624                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 6/22/2023             |
| Drugs    | J2427         | Injection, paliperidone<br>palmitate extended release<br>(invega hafyera, or invega<br>trinza), 1 mg | 1 mg                       | 7/1/2023                | Invega Hafyera™,<br>Invega Trinza®                      | paliperidone palmitate<br>extended-release injectable<br>suspension, for <del>gluteal</del><br>intramuscular use | Invega Trinza:<br>Indicated for the treatment of schizophrenia in patients after they have been adequately treated with<br>Invega Sustenna* (1-month paliperidone palmitate extended-release injectable suspension) for at least<br>four months.<br>Invega Hafyera:<br>Indicated for the treatment of schizophrenia in adults after they have been adequately treated with:<br>+ A once-*-month paliperidone palmitate extended-release injectable suspension (e.g., Invega Sustenna)<br>for at least four months or<br>+ An every-three-month paliperidone palmitate extended-release injectable suspension (e.g., Invega<br>Trinza) for at least one three-month cycle   | 1,560                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 6/22/2023             |
| Drugs    | J2430         | Injection, pamidronate<br>disodium, per 30 mg  | 30 mg                      | 1/1/2000                | Aredia®   | pamidronate disodium for<br>injection for intravenous<br>infusion  | Indicated for:<br>• Hypercalzemia of malignancy<br>• Paget's disease<br>• Osteolytic bone metastases of breast cancer and osteolytic lesions of multiple myeloma   | 6                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/21/2018             |
| Drugs    | J2440         | Injection, papaverine HCl, up<br>to 60 mg  | up to 60 mg                | 1/1/2000                | N/A – various<br>generics                               | papaverine hydrochloride<br>injection, solution  | Indicated in various conditions accompanied by spasm of smooth muscle, such as vascular spasm<br>associated with acute myocardial infarction (coronary occlusion), angina pectoris, peripheral and<br>pulmonary embolism, peripheral vascular disease in which there is a vasoopastic element, or certain<br>cerebral angiopastic states; and visceral spasm, as in wreteral, billary, or gastrointestinal colic.  | 80                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 7/16/2018             |

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|-------------|---------------|---|----------------------------|-------------------------|--------------------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Drugs       | J2469         | Injection, palonosetron HCl, 25<br>mcg  | 25 mcg                     | 1/1/2005                | Aloxi®                         | palonosetron HCl injection<br>for intravenous use                     | Indicated in adults for:<br>• Moderately emetogenic cancer chemotherapy – prevention of acute and delayed nausea and vomiting<br>associated with Initial and repeat courses.<br>• Highly emetogenic cancer chemotherapy – prevention of acute nausea and vomiting associated with<br>Initial and repeat courses.<br>• Prevention of postoperative nausea and vomiting (PONV) for up to 24 hours following surgery. Efficacy<br>beyond 24 hours has not been demonstrated.<br>Indicated in pediatric patients aged 1 month to less than 17 years for:<br>• Prevention of acute nausea and vomiting associated with Initial and repeat courses of emetogenic<br>cancer chemotherapy, including highly emetogenic cancer chemotherapy. | 50                                | 1 month     | N/A         | N/A                    | Y               | Y                               |          | 7/16/2018             |
| Drugs       | J2501         | Injection, paricalcitol, 1 mcg  | 1 mcg                      | 1/1/2003                | Zemplar®                       | paricalcitol injection  | Indicated for the prevention and treatment of secondary hyperparathyroidism associated with stage 5<br>chronic kidney disease (CKD).  | 420                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 7/16/2018             |
| Drugs       | J2502         | Injection, pasireotide long<br>acting, 1 mg   | 1 mg                       | 1/1/2016                | Signifor® LAR                  | pasireotide for injectable<br>suspension, for intramuscular<br>use    | Indicated for the treatment of:<br>• Patients with acromeeals who have had an inadequate response to surgery and/or for whom surgery is   | 120                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 7/26/2018             |
| Biologicals | J2506         | Injection, pegfilgrastim,<br>excludes biosimilar, 0.5 mg  | 0.5 mg                     | 1/1/2022                | Neulasta®,<br>Neulasta® Onpro® | pegfilgrastim injection, for<br>subcutaneous use                      | Indicated to:<br>- To decrease the incidence of infection, as manifested by febrile neutropenia, in patients with nonmyeloic<br>malignancies receiving myelosuppressive anticancer drugs associated with a clinically significant incidence<br>of febrile neutropenia.<br>- Increase survival in patients acutely exposed to myelosuppressive doses of radiation (Hematopoietic<br>Subsyndrome of Acute Radiation Syndrome).<br>Limitations of Use:<br>- Neulasta is not indicated for the mobilization of peripheral blood progenitor cells for hematopoietic stem<br>cell transplantation.  | 36                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 12/14/2021            |
| Biologicals | J2507         | Injection, pegloticase, 1 mg  | 1 mg                       | 1/1/2012                | Krystexxa®                     | pegloticase injection, for<br>intravenous infusion                    | Indicated for the treatment of chronic gout in adult patients refractory to conventional therapy.   | 24                                | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/4/2019              |
| Biologicals | J2508         | Injection, pegunigalsidase alfa-<br>iwxj, 1 mg  | 1 mg                       | 1/1/2024                | Elfabrio®                      | pegunigalsidase alfa-iwxj<br>injection, for intravenous use           | Indicated for the treatment of adults with confirmed Fabry disease.   | 420                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 12/22/2023            |
| Drugs       | J2510         | Injection, penicillin G procaine,<br>aqueous, up to 600,000 units   | up to 600,000 units        | 1/1/2000                | N/A                            | penicillin G procaine<br>injectable suspension                        | Indicated in the treatment of moderately severe infections in both adults and pediatric patients due to<br>penicilin-G-susceptible microorganisms that are susceptible to the low and persistent serum levels<br>common to this particular dosage form. Therapy should be guided by bacteriological studies (including<br>susceptibility tests) and by clinical response. See package insert for list of infections and microorganisms.   | 52                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 8/24/2018             |
| Drugs       | J2515         | Injection, pentobarbital<br>sodium, per 50 mg   | 50 mg                      | 1/1/2000                | Nembutal®                      | pentobarbital sodium<br>injection, USP                                | Indicated for use as:<br>• Sedatives<br>• Hyportics, for the short-term treatment of insomnia, since they appear to lose their effectiveness for<br>sleep induction and sleep maintenance after 2 weeks<br>• Preamesthetics<br>• Anticonvulsant, in anesthetic doses, in the emergency control of certain acute convulsive episodes, e.g.,<br>those associated with status epilepticus, cholera, eclampsia, meningits, tetanus, and toxic reactions to<br>strychnine or local anesthetics   | 150                               | N/A         | N/A         | N/A                    | ¥               | Y                               |          | 8/24/2018             |
| Drugs       | J2540         | Injection, penicillin G<br>potassium, up to 600,000<br>units  | 600,000 units              | 1/1/2000                | Pfizerpen®                     | penicillin G potassium for<br>injection                               | Indicated in the therapy of severe infections caused by penicillin G-susceptible microorganisms when rapic<br>and high penicillin levels are required. Therapy should be guided by bacteriological studies (including<br>susceptibility tests) and by clinical response. See package insert for full list of microorganisms.  | i<br>1,240                        | N/A         | N/A         | N/A                    | Y               | Y                               |          | 8/24/2018             |
| Drugs       | J2543         | Injection, piperacillin<br>sodium/tazobactam sodium, 1<br>g/0.125 g (1.125 g)   | 1.125 g                    | 1/1/2000                | Zosyn®                         | piperacillin and tazobactam<br>for injection, for intravenous<br>use  | Indicated for treatment of:<br>• Intra-abdominal infections<br>• Sina and skin structure infections<br>• Female pelvic infections<br>• Community-acquired pneumonia<br>• Nosocomial pneumonia<br>• Usage<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Zosyn and other<br>antibacterial drugs, Zosyn should be used only to treat or prevent infections that are proven or strongly<br>suspected to be caused by bacteria.  | 224                               | 2 months    | N/A         | N/A                    | Y               | Y                               |          | 4/10/2019             |
| Drugs       | J2545         | Pentamidine isethionate,<br>inhalation solution, FDA-<br>approved final product, non-<br>compounded, administered<br>through DME, unit dose form,<br>per 300 mg | 300 mg                     | 1/1/2000                | NebuPent®                      | pentamidine isethionate<br>inhalant (DME) for oral<br>inhalation only | Indicated for the prevention of Pneumocystis Jiroveci pneumonia (PJP) in high-risk, HIV-infected patients<br>defined by one or both of the following criteria:<br>• a history of one or more episodes of PJP<br>• a peripheral CD4+ (T4 helper/inducer) lymphocyte count less than or equal to 200/mm3  | 2                                 | 16 years    | N/A         | N/A                    | Y               | Y                               |          | 8/24/2018             |
| Drugs       | J2547         | Injection, peramivir, 1 mg  | 1 mg                       | 1/1/2016                | Rapîvab <sup>®</sup>           | peramivir injection, for<br>intravenous use                           | Indicated for the treatment of acute uncomplicated influenza in patients 6 months and older who have<br>been symptomatic for no more than two days.<br>Limitations of Use:<br>• Efficacy based on clinical trials in which the predominant influenza virus type was influenza A; a limited<br>number of subjects infected with influenza 8 virus were enrolled.<br>• Consider available information on influenza drug susceptibility patterns and treatment effects when<br>deciding whether to use.<br>• Efficacy could not be established in patients with serious influenza requiring hospitalization.   | 600                               | 6 months    | N/A         | N/A                    | Y               | Y                               |          | 2/25/2021             |

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| Category | HCPCS<br>Code | HCPCS Description                                 | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|----------|---------------|---|----------------------------|-------------------------|------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Drugs    | J2550         | Injection, promethazine HCI,<br>up to 50 mg       | up to 50 mg                | 1/1/2000                | Phenergan  | promethazine hydrochioride<br>injection                                | Indicated for the following conditions:<br>• Amelioration of allergic reactions to blood or plasma.<br>• In anaphylaxis as an adjunct to epinephrine and other standard measures after the acute symptoms<br>have been controlled.<br>• For other uncomplicated allergic conditions of the immediate type when oral therapy is impossible or<br>contraindicated.<br>• For sedation and relief of apprehension and to produce light sleep from which the patient can be easily<br>aroused.<br>• Active treatment of motion sickness.<br>• Prevention and control of nausea and womiting associated with certain types of anesthesia and surgery.<br>• As an adjunct to analgesics for the control of postoperative pain.<br>• Preperative, postoperative, and obstetric (during lador) sedation.<br>• Intravenously in special situations, such as repeated bronchoscopy, ophthalmic surgery, and<br>poor-risk patients, with reduced amounts of meperidine or other narcotic analgesic as an adjunct to<br>anesthesia and analgesia.  | 93                                | 2 years   | N/A         | N/A                    | Y               | Y                               |  | 8/24/2018             |
| Drugs    | J2560         | Injection, phenobarbital<br>sodium, up to 120 mg  | up to 120 mg               | 1/1/2000                | N/A        | phenobarbital sodium<br>injection                                      | Indicated for use as:<br>Sedative. Sedation is obtainable within an hour, and in adequate dosage, the duration of action is more<br>than six hours. Included in the more common conditions in which the sedative action of this class of drugs<br>is desired are anxiety-tension states, hyperthyrolism, essential hypertension, nausea and vomiting of<br>functional origin, motion sichness, acute labyrinthits; pylorospasm in infants, chorea and cardiac failure.<br>Phenobarital is also a useful adjunct in treatment of hemorrhage from the respiratory or gastrointestinal<br>tract. Phenobarital controls anxiety, decreases muscular activity and lessens nervous exclibility in<br>hyperthyroid patients. However, thyrotoxic individuals occasionally react poorly to barbiturates.<br>+ Hynotic, for the short-term treatment of insomina, since it appears to lose its effectiveness for sleep<br>induction and sleep maintenance after 2 weeks.<br>• Praneshtetic:<br>• Long-term anticonvulsant, (phenobarbital, mephobarbital and metharbital) for the treatment of<br>generalized tonic-clonic and cortical focal secures. And, in the emergency control of certain acute<br>convulsive egiose, e.g., those associated with status egilepticus, chiefera, eclampia, cerebral<br>hemorrhage, meningitis, tetanus, and toxic reactions to strychnine or local anesthetics. Phenobarbital<br>sodium may be administered intravenously at an anticonvulsant for energency use.<br>When administered intravenously, it may require 15 or more minutes before reaching peak concentrations<br>in the harin. Herefore, injecting phenobarbital out with the convulsions stop may cause the harin<br>level to exceed that required to control the convulsions and lead to severe barbiturate-induced<br>depression.<br>• Phenobarbital is indicated in pediatric patients as an anticonvulsant and as a sedative, including its<br>preoperative and postoperative use. | N/A                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 8/29/2018             |
| Drugs    | J2562         | Injection, plerixafor, 1 mg                       | 1 mg                       | 1/1/2010                | Mozobil®   | plerixafor injection, solution<br>for subcutaneous use                 | Indicated in combination with granulocyte-colony stimulating factor (G-CSF) to mobilize hematopoietic<br>stem cells (HSCs) to the peripheral blood for collection and subsequent autologous transplantation in<br>patients with non-Hodgkin's lymphoma and multiple myeloma.  | 160                               | 18 years  | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 6/6/2019              |
| Drugs    | J2590         | Injection, oxytocin, up to 10<br>units            | up to 10 units             | 1/1/2000                | Pitocin®   | oxytocin injection, USP<br>synthetic                                   | Indicated for:<br>• Antepartum<br>- The initiation or improvement of uterine contractions, where there is desirable and considered suitable<br>for reasons of fetal or maternal concern, in order to achieve vaginal delivery.<br>- Induction of labor in patients with a medical indication for the initiation of labor.<br>- Stimulation or reinforcement of labor, as in selected cases of uterine inertia.<br>- Adjunctive therapy in the management of incomplete or inevitable abortion.<br>• Postpartum<br>- Produce uterine contractions during the third stage of labor and to control postpartum bleeding or<br>hemorrhage.   | 12                                | N/A   | N/A         | Females Only           | ¥               | Ŷ                               |  | 7/16/2018             |
| Drugs    | J2597         | Injection, desmopressin<br>acetate, per 1 mcg     | 1 mcg                      | 1/1/2000                | DDAVP®     | desmopressin acetate<br>injection                                      | Indicated for patients with hemophilia A with factor VIII coagulant activity levels greater than 5%, patients<br>with mild to moderate classic von Willebrand's disease (Type 1) with factor VIII levels greater than 5%, as<br>an antidiuretic replacement therapy in the management of central (crania) (alabetes insipidus and for the<br>management of the temporary polynia and polydigisa following head truma or surgery in the pituitary<br>region. DDAVP is ineffective for the treatment of nephrogenic diabetes insipidus.   | 660                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication age specific:<br>Hemophilia A and von<br>Willebrand's Disease: 3<br>months of age and older<br>Diabetes Insipidus: 12 years of<br>age and older | 7/2/2018              |
| Drugs    | J2675         | Injection, progesterone, per 50<br>mg             | per 50 mg                  | 1/1/2003                | N/A        | progesterone injection, in<br>sesame oil for intramuscular<br>use only | Indicated in amenorrhea and abnormal uterine bleeding caused by hormonal imbalance in the absence of<br>organic pathology, such as submucous fibroids or uterine cancer.  | 2                                 | 18 years  | N/A         | Females Only           | Y               | Y                               |  | 6/6/2019              |
| Drugs    | J2679         | Injection, fluphenazine hcl,<br>1.25 mg           | 1.25 mg                    | 1/1/2024                | N/A        | fluphenazine hydrochloride<br>injection, solution                      | <ul> <li>Fluphenazine hydrochloride has not been shown effective in the management of behavioral<br/>complications in patients with mental retardation.</li> </ul>  | 248                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/22/2023            |
| Drugs    | J2680         | Injection, fluphenazine<br>decanoate, up to 25 mg | up to 25 mg                | 1/1/2000                | N/A        | fluphenazine decanoate<br>injection                                    | Intended for use in the management of patients requiring prolonged parenteral neuroleptic therapy (e.g.<br>chronic schitopriencis). Fluphenazine decanoate has not been shown effective in the management of<br>behavioral complications in patients with mental retardation.   | 8                                 | 12 years  | N/A         | N/A                    | Y               | Y                               |  | 6/4/2019              |
| Drugs    | J2690         | Injection, procainamide HCl,<br>up to 1 g         | up to 1 g                  | 1/1/2000                | N/A        | procainamide hydrochloride<br>injection, solution                      | Indicated for the treatment of documented ventricular arrhythmias, such as sustained ventricular<br>tachycardia, that, in the judgement of the physician, are life-threatening. Because of the proarrhythmic<br>effects of procainamide, its use with lesser arrhythmias is generally not recommended. Treatment of<br>patients with samptionatic ventricular premature contractions should be avoided.   | 7                                 | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 6/6/2019              |

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| nedicaid/medi | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name             | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modifi<br>Date |
|---------------|---------------|--|----------------------------|-------------------------|------------------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|---------------------|
| Drugs         | J2700         | Injection, oxacillin sodium, up<br>to 250 mg                             | up to 250 mg               | 1/1/2000                | N/A, various generics  | oxacillin sodium injection,<br>powder, for solution for<br>intramuscular or intravenous<br>use | Indicated for the treatment of infections caused by penicillinase-producing staphylococci which have<br>demonstrated susceptibility to the drug. Cultures and susceptibility tests should be performed initially to<br>determine the causative organism and their susceptibility to the drug.  | 744                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 9/21/2018           |
| Drugs         | J2710         | Injection, neostigmine<br>methylsulfate, up to 0.5 mg                    | up to 0.5 mg               | 1/1/2000                | Bloxiverz <sup>®</sup> | neostigmine methylsulfate<br>injection, for intravenous use                                    | Indicated for the reversal of the effects of nondepolarizing neuromuscular blocking agents (NMBAs) after<br>surgery.   | 50                                | N/A   | N/A         | N/A                    | Y               | Y                               |   | 4/10/2019           |
| Drugs         | J2720         | Injection, protamine sulfate,<br>per 10 mg                               | 10 mg                      | 1/1/2000                | N/A                    | protamine sulfate injection, solution for intravenous use                                      | Indicated for the treatment of heparin overdosage.   | 5                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 8/29/2018           |
| Biologicals   | J2724         | Injection, protein C<br>concentrate, intravenous,<br>human, 10 IU        | 10 IU                      | 1/1/2008                | Ceprotin               | protein c concentrate<br>(human) lyophilized power<br>for solution for injection               | Indicated for pediatric and adult patients with severe congenital Protein C deficiency for the prevention<br>and treatment of venous thrombosis and purpura fulminans.   | 105,840                           | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 6/4/2019            |
| Drugs         | J2730         | Injection, pralidoxime chloride,<br>up to 1 g                            | up to 1 g                  | 1/1/2000                | Protopam®              | pralidoxime chloride for<br>injection  | Indicated as an antidote:<br>• In the treatment of poisoning caused by those pesticides and chemicals of the organophosphate class<br>which have anticholinesterase activity.<br>• In the control of overdosage by anticholinesterase drugs used in the treatment of myasthenia gravis.  | 20                                | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 8/24/2018           |
| Drugs         | J2760         | Injection, phentolamine<br>mesylate, up to 5 mg                          | up to 5 mg                 | 1/1/2000                | Regitine®              | phentolamine mesylate<br>injection, powder,<br>lyophilized, for suspension                     | Indicated for:<br>• The prevention or control of hypertensive episodes that may occur in a patient with pheochromocytoma<br>sa result of stress or manipulation during preoperative preparation and surgical excision.<br>• The prevention or treatment of dermal necrosis and sloughing following intravenous administration or<br>extravasation of nergeinpehnic.<br>The diagnosis of pheochromocytoma by the phentolamine mesylate for injection blocking test.   | 372                               | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 8/24/2018           |
| Drugs         | J2765         | Injection, metoclopramide<br>HCl, up to 10 mg                            | up to 10 mg                | 1/1/2000                | N/A                    | metoclopramide<br>hydrochloride injection  | Indicated for:<br>• The relief of symptoms associated with acute and recurrent diabetic gastric stasis<br>• The prophylaxis of vomiting associated with emetogenic cancer chemotherapy<br>• The prophylaxis of postoperative nausea and vomiting in those circumstances where nasogastric suction<br>is undesirable<br>• Faoittaing small bowel intubation in adults and pediatric patients in whom the tube does not pass the<br>pylons with conventions maneuvers<br>• Stimulating gastric emptying and intestinal transit of barium in cases where delayed emptying interferes<br>with radiological examination of the stomach and/or small intesting | 560                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific:<br>• Facilitating Small Bowel<br>Intubation: 18 years of age and<br>older<br>• All other indications: None | 6/6/2019            |
| Biologicals   | J2777         | Injection, faricimab-svoa, 0.1<br>mg                                     | 0.1 mg                     | 10/1/2022               | Vabysmo®               | faricimab-svoa injection, for<br>intravitreal use  | Indicated for the treatment of patients with:<br>• Neovascular (Wet) Age-Related Macular Degeneration (nAMD)<br>• Diabetic Macular Edema (DME)<br>• Macular Edema Following Retinal Vein Occlusion (RVO)   | 240                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 12/1/202            |
| Biologicals   | J2778         | Injection, ranibizumab, 0.1 mg   | 0.1 mg                     | 1/1/2008                | Lucentis®              | ranibizumab injection for<br>intravitreal injection  | indicated for the treatment of patients with:<br>• Neovascular (Wet) Age-Related Macular Degeneration (AMD)<br>• Macular Edema Following Retinal Vein Occlusion (RVO)<br>• Diabetic Macular Edema (DME)<br>• Diabetic Retinopathy (DR)<br>• Wyopic Charoidal Neovascularization (mCNV)   | 20                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 10/31/201           |
| Biologicals   | J2779         | Injection, ranibizumab, via<br>intravitreal implant (susvimo),<br>0.1 mg | 0.1 mg                     | 7/1/2022                | Susvimo™               | ranibizumab injection for<br>intravitreal use via ocular<br>implant                            | Indicated for the treatment of patients with Neovascular (wet) Age-related Macular Degeneration (AMD)<br>who have previously responded to at least two intravitreal injections of a VEGF inhibitor.  | 100                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 6/6/2022            |
| Drugs         | J2781         | Injection, pegcetacoplan,<br>intravitreal, 1 mg                          | 1 mg                       | 10/1/2023               | Syfovre™               | pegcetacoplan injection, for<br>intravitreal use   | Indicated for the treatment of geographic atrophy (GA) secondary to age-related macular degeneration<br>(AMD).   | 60                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 9/28/2023           |
| Drugs         | J2782         | Injection, avacincaptad pegol,<br>0.1 mg                                 | 0.1 mg                     | 4/1/2024                | lzervay™               | avacincaptad pegol<br>intravitreal solution  | Indicated for the treatment of geographic atrophy (GA) secondary to age-related macular degeneration<br>(AMD).   | 80                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 4/12/202            |
| Biologicals   | J2783         | Injection, rasburicase, 0.5 mg   | 0.5 mg                     | 1/1/2004                | Elitek®                | rasburicase for injection, for intravenous use   | Indicated for the initial management of plasma uric acid levels in pediatric and adult patients with<br>leukemia, lymphoma, and solid tumor malignancies who are receiving anti-cancer therapy expected to<br>result in tumor lysis and subsequent elevation of plasma uric acid.  | 280                               | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 6/4/201             |
| Drugs         | J2785         | Injection, regadenoson, 0.1 mg   | 0.1 mg                     | 1/1/2009                | Lexiscan®              | regadenoson injection for<br>intravenous use   | Limitation of Use: Elitek is indicated for a single course of treatment.<br>Indicated for radionuclide myocardial perfusion imaging (MPI) in patients unable to undergo adequate<br>exercise stress.   | 4                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 6/4/2021            |
| Biologicals   | J2786         | Injection, reslizumab, 1 mg  | 1 mg                       | 1/1/2017                | Cinqair®               | reslizumab injection, for<br>intravenous use   | Indicated for add-on maintenance treatment of patients with severe asthma aged 18 years and older, and<br>with an eosinophilic phenotype.<br>Limitations of Use: Cinqair is not indicated for:<br>• Treatment of other eosinophilic conditions.<br>• Relief of acute bronchospasm or status asthmaticus.   | 840                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 7/2/2018            |

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|---------------------|---------------|--|----------------------------|-------------------------|---|--|--|-----------------------------------|---|-------------|---------------------------|-----------------|---------------------------------|---|----------------------|
| Immune<br>Globulins | J2788         | Injection, Rho d immune<br>globulin, human, minidose, 50<br>micrograms (250 IU)                      | 50 mcg                     | 1/1/2003                | HyperRHO <sup>®</sup> S/D<br>Mini Dose,<br>MICRhoGAM <sup>®</sup> , | rho(D) immune globulin<br>(human), mini dose   | HyperRHO 5//D Mini Dose: recommended to prevent the isoimmunization of Rho(D) negative women at<br>the time of spontaneous or induced abortion of up to 12 weeks' gestation provided the following criteria<br>are met:<br>1. The mother must be Rho(D) negative and must not already be sensitized to the Rho(D) antigen.<br>2. The father is not known to be Rho(D) negative.<br>3. Gestation is not more than 12 weeks at termination.<br>**See package insert for full usage criteria.**<br>MICRhoGAM: For use in preventing Rh immunization.<br>* Pregnancy and other obstetrical conditions in Rh-negative bother of hab are<br>conclusively Rh-negative, e.g. delivery of an Rh-negative bother of hab are<br>and baby, any netapartum fetah-maternal henorrhange (suspected or proven), actual or threatened<br>pregnancy loss at any stage of gestation and ectopic pregnancy.<br>* Prevention of Rh immunization in any Rh-negative person after incompatible transfusion of Rh-positive<br>blood or blood products.   | 1                                 | N/A   | N/A         | HyperRHO:<br>Females Only | Y               | Ŷ                               |   | 7/3/2018             |
| Immune<br>Globulins | J2790         | Injection, Rho d immune<br>globulin, human, full dose, 300<br>micrograms (1500 IU)                   | 300 mcg (1500 IU)          | 1/1/2003                | HyperRho® S/D<br>Full Dose,<br>RhoGAM®                              | rho(d) immune globulin<br>(human), full dose   | Indicated for use in preventing Rh immunization:<br>• In pregnancy and other obstetrical conditions (see full prescribing information).<br>• In any Rh-negative person after incompatible transfusion of Rh-positive blood or blood products.  | 3                                 | N/A   | N/A         | N/A                       | Y               | Y                               |   | 4/9/2022             |
| Immune<br>Globulins | J2791         | Injection, Rho(D) immune<br>globulin (human), (Rhophylac)<br>intramuscular or intravenous,<br>100 IU | . 100 IU                   | 1/1/2008                | Rhophylac®  | rho(d) immune globulin<br>intravenous (human) 1500 IU<br>(300 mcg) solution for<br>intravenous (IV) or<br>Intramuscular (IM) injection | Indicated for:<br>Suppression of Rhesus (Rh) Isoimmunization in:<br>Pregnancy and obstetric conditions in non-sensitized, Rho (D)-negative women with an Rh-incompatible<br>pregnancy, including:<br>- Rhoutine antegratum and postpartum Rh prophylaxis<br>- Rho prophylaxis in obstetric complications or invasive procedures<br>- Incompatible transfusions in Rho (D)-negative individuals transfused with blood components containing<br>Rho (D)-positive red blood cells (RdCs).<br>Immune Thrombocytopenic Purpura (ITP)<br>- Raising platelet counts in Rho (D)-positive, non-splenectomized adults with chronic ITP.  | 350                               | N/A   | N/A         | N/A                       | Y               | Ŷ                               | 12/2023: Age restrictions<br>updated to align with other<br>rho(D) immune globulin<br>products effective<br>12/20/2023. | 1/26/2024            |
| Immune<br>Globulins | J2792         | Injection, rho D immune<br>globulin, intravenous, human,<br>solvent detergent, 100 IU                | 100 IU                     | 1/1/2000                | WinRho SDF*   | rho(D) immune globulin<br>intravenous (human) solution<br>for intravenous or<br>intramuscular injection                                | Indicated for:<br>Immune Thrombocytopenic Purpura (ITP)<br>Raising platelet counts in Rho(D) positive, non-splenectomized:<br>• Children with chronic or acute ITP,<br>• Aduts with chronic ITP and<br>• Children and aduts with ITP secondary to HIV Infection<br>Suppression of Rhiseus (Rh) Isoimmunization<br>• Pregnancy and other obstetric conditions in non-sensitized, Rho(D)-negative women with an Rh-<br>incompatible pregnancy including:<br>O Routine antepartum and postpartum Rh prophylaxis<br>O Rh prophylaxis in obstetric complications or invasive procedures<br>• Incompatible transfusions in Rho(D)-negative individuals transfused with blood components containing<br>Rho(D)-positive ed blood cells (ReCs).   | 1,500                             | N/A   | N/A         | N/A                       | Y               | Ŷ                               |   | 9/12/2018            |
| Biologicals         | J2793         | Injection, rilonacept, 1 mg  | 1 mg                       | 1/1/2010                | Arcalyst <sup>®</sup>   | rilonacept injection for subcutaneous use  | Indicated for:<br>Indicate for:<br>'the treatment of patients with Cryopyrin-Associated Periodic Syndromes (CAPS), including Familial Cold<br>Autoinflammatory Syndrome (FACS) and Muckie-Wells Syndrome (MWS) in adults and children 12 years of<br>age and older.<br>Zmaintenance of remission of Deficiency of Interleukin-1 Receptor Antagonist (DIRA) in adults and<br>pediatric patients weighing at least 10 kg.<br>'the treatment of recurrent pericarditis (RP) and reduction in risk of recurrence in adults and children 12<br>years and older.   | 1,600                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                       | Y               | Y                               | Indication specific age<br>restrictions:<br>CAPS and RP: 12 years of age<br>and older<br>DIRA: N/A                      | 4/26/2021            |
| Drugs               | J2794         | Injection, risperidone<br>(risperdal consta), 0.5 mg   | 0.5 mg                     | 1/1/2005                | Risperdal Consta®   | risperidone long-acting<br>injection   | Indicated:<br>• for the treatment of schizophrenia.<br>• as monotherapy or as adjunctive therapy to lithium or valproate for the maintenance treatment of<br>Biodair I Disorder.   | 300                               | N/A   | N/A         | N/A                       | Y               | Y                               |   | 10/3/2019            |
| Drugs               | J2795         | Injection, ropivacaine<br>hydrochloride, 1 mg  | 1 mg                       | 1/1/2001                | Naropin®  | ropivacaine HCl injection  | Indicated for the production of local or regional anesthesia for surgery and for acute pain management.<br>Surgical Anesthesia: epidural block for surgery including cesarean section; major nerve block; local<br>infiltration.<br>Acute pain management: epidural continuous infusion or intermittent bolus, eg. postoperative or labor;<br>local infiltration.  | 2,166                             | 18 years  | N/A         | N/A                       | Y               | Y                               |   | 8/29/2018            |
| Drugs               | J2796         | Injection, romiplostim, 10<br>micrograms   | 10 mcg                     | 1/1/2010                | Nplate*   | romiplostim for injection, for<br>subcutaneous use   | Indicated for the treatment of thrombocytopenia in:<br>• Adult patients with immune thrombocytopenia (ITP) who have had an insufficient response to<br>corticosteroids, immunglobulins, or splenectomy.<br>• Pediatric patients 1 year of age and older with ITP for at least 6 months who have had an insufficient<br>response to corticosteroids, immunglobulins, or splenectomy.<br>Nplate is indicated to increase survival in adults and in pediatric patients (including term neonates) acutely<br>exposed to myelosuppressive doses of radiation (Hematopoietic Syndrome of Acute Radiation Syndrome<br>(HSARS)).<br>Limitations of Use:<br>• Nplate is not indicated for the treatment of thrombocytopenia due to myelodysplastic syndrome (MDS)<br>or any cause of thrombocytopenia other than ITP.<br>• Nplate should be used only in patients with ITP whose degree of thrombocytopenia and clinical condition<br>increases the risk for bleeding.<br>• Nplate should be used only in patients with ITP whose degree of thrombocytopenia and clinical condition<br>increases the risk for bleeding. | 700                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                       | Ŷ               | Y                               | Indication Specific Age<br>Restrictions:<br>ITP: 1 year of age and older<br>HS-ARS: None                                | 2/25/2021            |
| Drugs               | J2798         | Injection, risperidone,<br>(perseris), 0.5 mg  | 0.5 mg                     | 10/1/2019               | Perseris™   | risperidone for extended-<br>release injectable suspension,<br>for subcutaneous use  | Indicated for the treatment of schizophrenia in adults.  | 480                               | 18 years  | N/A         | N/A                       | Ŷ               | Ŷ                               |   | 10/3/2019            |

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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age                           | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modifie<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|------------|--|---|-----------------------------------|---|---------------------------------------|------------------------|-----------------|---------------------------------|---|----------------------|
| Drugs       | J2799         | Injection, risperidone (uzedy),<br>1 mg  | 1 mg                       | 1/1/2024                | Uzedy™     | risperidone extended-release<br>injectable suspension, for<br>subcutaneous use       | Indicated for the treatment of schizophrenia in adults.   | 250                               | 18 years  | N/A                                   | N/A                    | Y               | Y                               |   | 12/22/2023           |
| Drugs       | J2800         | Injection, methocarbamol, up<br>to 10 mL                                       | up to 10 mL                | 1/1/2000                | Robaxin®   | methocarbamol injection for<br>intravenous or intramuscular<br>use                   | Indicated as an adjunct to rest, physical therapy, and other measures for the relief of discomfort<br>associated with acute, painful, musculoskeletal conditions; supportive therapy in tetanus.  | 54                                | Indication Specific<br>Age Restrictions<br>(see comments) | N/A                                   | N/A                    | Y               | Y                               | Indication specific.<br>Relief of discomfort associated<br>with acute, painful,<br>musculoskeletal conditions: 18<br>years of age and older.<br>Tetanus: None   | 6/8/2010             |
| Drugs       | J2801         | Injection, risperidone<br>(rykindo), 0.5 mg                                    | 0.5 mg                     | 4/1/2024                | Rykindo®   | risperidone for extended-<br>release injectable suspension,<br>for intramuscular use | Risperidone for extended-release injectable suspension is indicated:<br>• for the treatment of schizophrenia in adults.<br>• as monotherapy or as adjunctive therapy to lithium or valproate for the maintenance treatment of<br>bippolar I disorder in adults.   | 300                               | 18 years  | N/A                                   | N/A                    | Y               | Ŷ                               |   | 4/12/2024            |
| Biologicals | J2820         | Injection, sargramostim (GM-<br>CSF), 50 mcg                                   | 50 mcg                     | 1/1/2000                | Leukino*   |  | Indicated:<br>• To shorten time to neutrophil recovery and to reduce the incidence of severe and life-threatening<br>induction chemotherapy in adult patients 55 years and older with acute myeloid leukemia (AML).<br>• For the mobilization of hematopoietic progenitor cells into peripheral blood for collection by<br>leukapheresis and autologous transplantation in adults.<br>• For the acceleration of myeloid reconstitution following autologous bone marrow or peripheral blood<br>progenitor cell transplantation in adults.<br>• For the acceleration of myeloid reconstitution following autologous bone marrow or peripheral blood<br>progenitor cell transplantation in adults.<br>• For the acceleration of myeloid reconstitution following allogeneic bone marrow transplantation in adult<br>and pediatric patients 2 years of age and<br>older.<br>• For theratent of delayed neutrophil recovery or graft failure after autologous or allogeneic bone marrow<br>transplantation in adult and pediatric patients 2 years of age and older.<br>• To increase survial in adult and pediatric patients 2 years of age and older.<br>• To increase survial in adult and pediatric patients 2 years of age and older.<br>• To increase survial in adult and pediatric patients 2 years of age and older.<br>• To increase survial in adult and pediatric patients 2 years of age acutely exposed to<br>myelosuppressive doses of radiation (Hematopoietic Syndrome of Acute Radiation Syndrome [H-ARS]). | 620                               | Indication Specific<br>Age Restrictions<br>(see comments) | Indication Specific<br>(see comments) | N/A                    | ¥               | ¥                               | Indication specific age<br>restrictions:<br>• To shorten time to<br>neutrophil recovery and to<br>reduce the incidence of severe<br>and life-threatening infections<br>and infections resulting in<br>death following induction<br>chemotherapy in adult<br>patients 55 years and older<br>with acute myeloid leukemia<br>(AML).<br>• For the mobilization of<br>hematopoietic progenitor cells<br>into peripheral blood for<br>collection by leukapheresis<br>and autologous<br>transplantation in adults.<br>• For the acceleration of<br>myeloid reconstitution<br>following autologous bone<br>marrow or peripheral blood<br>progenitor cell transplantation<br>in adult and pediatric patients<br>2 years of age and older.<br>• For the acceleration of<br>myeloid reconstitution<br>following autopage based based based<br>progenitor cell transplantation<br>in adult and pediatric patients | 8/29/2018            |
| Biologicals | J2840         | Injection, sebelipase alfa, 1 mg   | 1 mg                       | 1/1/2017                | Kanuma®    | sebelipase alfa injection, for<br>intravenous use                                    | Indicated for the treatment of patients with a diagnosis of Lysosomal Acid Lipase (LAL) deficiency.   | 1,260                             | 1 month   | N/A                                   | N/A                    | Y               | Y                               |   | 12/16/2021           |
| Biologicals | J2860         | Injection, siltuximab, 10 mg   | 10 mg                      | 1/1/2016                | Sylvant®   | siltuximab for injection, for<br>intravenous use                                     | Indicated for treatment of patients with multicentric Castleman's disease (MCD) who are human<br>immunodeficiency virus (HIV) negative and human herpesvirus-8 (HHV-8) negative.<br>Limitations of Use: Sylvant was not studied in patients with MCD who are HIV positive or HHV-8 positive<br>because Sylvant did not bind to viruly produced LG-in a non-clinical study.  | 400                               | 18 years  | N/A                                   | N/A                    | Ŷ               | Ŷ                               |   | 6/7/2019             |
| Drugs       | J2916         | Injection, sodium ferric<br>gluconate complex in sucrose<br>injection, 12.5 mg | 12.5 mg                    | 1/1/2003                | Ferrlecit® | sodium ferric gluconate<br>complex in sucrose injection,<br>for intravenous (IV) use | Indicated for the treatment of iron deficiency anemia in patients 6 years of age and older with chronic<br>kidney disease receiving hemodialysis who are receiving supplemental epoetin therapy.  | 80                                | 6 years   | N/A                                   | N/A                    | Y               | Y                               |   | 9/21/2018            |

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|             | HCPCS<br>Code | HCPCS Description                                       | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                          | Generic Name  | FDA Approved Indications   | NC Suggested Max | Minimum Age | Maximum Age | Gender       | NDC      | Rebating<br>Labeler | Comments   | Last Modified |
|-------------|---------------|---|----------------------------|-------------------------|-------------------------------------|---|--|------------------|-------------|-------------|--------------|----------|---------------------|--|---------------|
|             |               |   |                            |                         |                                     |   | (See Package Insert for full FDA approved indication descriptions)   | Monthly Units    | Minimum Age | maximum Age | Restrictions | Required | Required            | Commente   | Date          |
| Drugs       | J2919         | Injection, methylprednisolone<br>sodium succinate, 5 mg | 5 mg                       | 4/1/2024                | Solu-Medrol*                        | methylprednisolone sodium<br>succinate for injection  | When oral therapy is not reasible, and the storeging, obsage form, and route of administration of the drug<br>reasonably lend the preparation to the treatment of the condition, the intravenous or intranuscular use of<br>Solu-Medrol is indicated as follows:<br>Solu-Medrol is indicated as follows:<br>Allergic states: Control of severe or incapacitating allergic conditions intractable to adequate trials of<br>conventional treatment in asthma, atopic dermatitis, contact dermatitis, drug hypersensitivity reactions,<br>serum sickness, transfusion reactions.<br>I endocrine disorders: Primary or secondary adenocortical insufficiency (hydrocortisone or cortisone is<br>the drug of choice; synthetic analogs may be used in conjunction with mineralocorticoids where<br>applicable; in financy, mineralocortical supplementation is of particular importance). Congenital adrenal<br>hyperplasia, hypercalcemia associated with cancer, nonsuppurative thyroidits.<br>• Gastorinettraind diseases: To tilde quotinum hemohytic anemia, congenital (exprimed) particular<br>(sastorinettraind diseases): To tilde quotimuum hemohytic anemia, congenital (exprimed) particular<br>elemanologi diseases: To tilde quotimumup hemohytic anemia, congenital (exprimed) hypeplastic<br>anemia (Diamond-Blackfan anemia), idiopathic thrombocytopenic purpura in adults (intravenous<br>administration ony; intramuscular administration is contraindicated), pure red el aplasia, selected cases<br>of secondary thromocytopenia.<br>• Miscellaneous: Trichinosis with neurologic or myocardial involvement, tuberculous meningitis with<br>subarachinod block or impeding block when used concurrently with appropriate antituberculous<br>chemotherapy.<br>• Neoplastic diseases: For the palliative management of leukemias and lymphomaa.<br>• Nevenita diseases: For the palliative management of leukemias and lymphomas.<br>• Nevenita diseases: For the palliative diversions of proteinuria in idiopathic nephrotic syndrome or that<br>even bareneous evenices or emission of proteinuria in idiopathic nephrotic syndrome or that<br>even barenexervent. | 4,500            | N/A         | N/A         | N/A          | Y        | ¥                   |  | 3/22/2024     |
| Biologicals | J2993         | Injection, reteplase, 18.1 mg                           | 18.1 mg                    | 1/1/2002                | Retavase®                           | reteplase for injection, for<br>intravenous use   | Indicated for treatment of acute ST-elevation myocardial infarction (STEMI) to reduce the risk of death<br>and heart failure.<br>Limitation of Use: The risk of stroke may outweigh the benefit produced by thrombolytic therapy in<br>patients whoos STEMI puts them at low risk for death or heart failure.  | 2                | 18 years    | N/A         | N/A          | Y        | Ŷ                   |  | 10/31/2018    |
| Biologicals | J2997         | Injection, alteplase recombinant, 1 mg                  | 1 mg                       | 1/1/2001                | Activase®,<br>Cathflo®<br>Activase® | alteplase for injection, for intravenous use  | Cathflo Activase: Indicated for the restoration of function to central venous access devices as assessed by<br>the ability to withdraw blood.<br>Activase: Indicated for the treatment of:<br>• Acute Lischemic Stroke (AIS)<br>• Acute Myocardial Infarction (AMI) to reduce mortality and incidence of heart failure. Limitation of use<br>in AMI: The risk of stroke may be greater than the benefit in patients at low risk of death from cardiac<br>causes.<br>• Acute Maxieve Pulmonary Embolism (PE) for tysis.   | 3,100            | 18 years    | N/A         | N/A          | Y        | Y                   | 1/2024: Category corrected<br>from Drugs to Biologicals. | 1/26/2024     |
| Biologicals | J2998         | Injection, plasminogen, human<br>tvmh, 1 mg             | 1 mg                       | 1/1/2002                | Ryplazim®                           | plasminogen, human-tvmh<br>lyophilized powder for<br>reconstitution, for<br>intravenous use | Indicated for the treatment of patients with plasminogen deficiency type 1 (hypoplasminogenemia).  | 15,411.2         | 11 months   | N/A         | N/A          | Y        | Y                   |  | 6/6/2022      |
| Drugs       | J3000         | Injection, streptomycin, up to<br>1 gram                | up to 1 g                  | 1/1/2000                | N/A                                 |   | Indicated for the treatment of individuals with moderate to severe infections caused by susceptible strains<br>of microorganisms in the specific conditions of Mycobacterium tuberculosis and Non-tuberculosis<br>infections: Mycobacterium tuberculosis, and other sensitive non tuberculosis pathogens including<br>Pasteurella pestis (plague); Francisella tularensis (tularemia); Brucella; Calymmatobacterium granulomatis<br>(donovanosis; granuloma inguinale); H. ducrey (charcold); H. influenze (in respiratory, endocardia), and<br>meningeal infections, concomitantly with another antibacterial agent); K. pneumoniae pneumonia, and<br>Enterococcus faecalis in urinary tract infections; Streptococcus viridans; Enterococcus faecalis (in<br>endocardia) infectiona, concomitantly with pencillin); Gram-negative bacillary bacteremia (concomitantly<br>with another antibacterial agent); C. Streptoceccus viridans; Enterococcus faecalis (in<br>endocardia) infectiona, concomitantly with pencillin); Gram-negative bacillary bacteremia (concomitantly<br>with another antibacterial agent); C. Streptoceccus viridans; Enterococcus faecalis (in<br>endocardia) infectiona, concomitantly with pencillin); Gram-negative bacillary bacteremia (concomitantly<br>with another antibacterial agent); C. Streptoceccus viridans; Enterococcus faecalis (in<br>endocardia) infectiona, concomitantly with pencillin); Gram-negative bacillary bacteremia (concomitanty)   | 62               | N/A         | N/A         | N/A          | Y        | ¥                   |  | 6/7/2019      |
| Drugs       | J3010         | Injection, fentanyl citrate, 0.1<br>mg                  | 0.1 mg                     | 1/1/2000                | N/A                                 | fentanyl citrate injection, for<br>intravenous or intramuscular<br>use                      | Indicated for:<br>• analgesic action of short duration during the anesthetic periods, premedication, induction and<br>maintenance, and in the immediate postoperative period (recovery room) as the need arises.<br>• use as an opioid analgesic supplement in general or regional anesthesia.<br>• administration with a neuroleptic as an anesthetic premedication, for the induction of anesthesia and as<br>an adjunct in the maintenance of general and regional anesthesia.<br>• use as an anesthetic agent with oxygen in selected high risk patients, such as those undergoing open<br>heart surgery or certain complicated neurological or orthopedic procedures.   | 210              | 2 years     | N/A         | N/A          | Y        | Y                   |  | 6/4/2019      |
| Drugs       | J3030         | Injection, sumatriptan,<br>succinate, 6 mg              | 6 mg                       | 1/1/2000                | Imitrex®                            | sumatriptan succinate<br>injection, for subcutaneous<br>use                                 | Indicated for:<br>• Acute treatment of migraine with or without aura in adults<br>• Acute treatment of cluster headache in adults<br>timitations of Use:<br>Use only if a clear diagnosis of migraine or cluster headache has been established. Not indicated for the<br>prophylacit: therapy of migraine or cluster headache attacks.   | 8                | 18 years    | N/A         | N/A          | Y        | Ŷ                   |  | 9/21/2018     |
| Biologicals | J3055         | Injection, talquetamab-tgvs,<br>0.25 mg                 | 0.25 mg                    | 4/1/2024                | Talvey™                             | talquetamab-tgvs injection,<br>for subcutaneous use   | Indicated for the treatment of adult patients with relapsed or refractory multiple myeloma who have<br>received at least four prior lines of therapy, including a proteasome inhibitor, an immunomodulatory<br>agent and an anti-CD38 monoclonal antibody.   | 1,808            | 18 years    | N/A         | N/A          | Y        | Y                   |  | 4/12/2024     |
| Biologicals | J3060         | Injection, taliglucerase alfa, 10<br>units              | 10 units                   | 1/1/2014                | Elelyso®                            | taliglucerase alfa for<br>injection, for intravenous use                                    | Indicated for the treatment of patients with a confirmed diagnosis of Type 1 Gaucher disease.  | 2,520            | 4 years     | N/A         | N/A          | Y        | Y                   |  | 6/4/2019      |

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|----------------|---------------|--|----------------------------|-------------------------|------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Category       | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age                               | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
| Drugs          | 13090         | Injection, tedizolid phosphate,<br>1 mg                          | 1 mg                       | 1/1/2016                | Sivextro®  | tedizolid phosphate for<br>injection, for intravenous use      | Indicated in adults and pediatric patients 12 years of age and older for the treatment of acute bacterial<br>skin and skin structure infections (ABSSSI) caused by designated susceptible bacteria.   | 1,200                             | 12 years                                  | N/A         | N/A                    | Y               | Y                               |          | 7/28/2020             |
| Drugs          | 13095         | Injection, telavancin, 10 mg                                     | 10 mg                      | 1/1/2011                | Vibativ®   | telavancin for injection, for<br>intravenous use               | Indicated for the treatment of the following infections in adult patients caused by designated susceptible<br>bacteria:<br>• Complicated skin and skin structure infections (cSSSI)<br>• Hospital-acquired and ventilator-associated bacterial pneumonia (HABP/VABP) caused by susceptible<br>isolates of Staphylococcus aureus. Vibativ should be reserved for use when alternative treatments are not<br>suitable.  | 3,150                             | 18 years                                  | N/A         | N/A                    | Y               | Y                               |          | 6/8/2019              |
| Drugs          | J3105         | Injection, terbutaline sulfate,<br>up to 1 mg                    | up to 1 mg                 | 1/1/2000                | N/A        | terbutaline sulfate injection,<br>solution                     | Indicated for the prevention and reversal of bronchospasm in patients 12 years of age and older with<br>asthma and reversible bronchospasm associated with bronchitis and emphysema.  | 45                                | 12 years                                  | N/A         | N/A                    | Y               | Y                               |          | 9/12/2018             |
| Biologicals    | J3111         | Injection, romosozumab-aqqg,<br>1 mg                             | 1 mg                       | 10/1/2019               | Evenity™   | romosozumab-aqqg injection<br>for subcutaneous use             | Indicated for the treatment of osteoporosis in postmenopausal women at high risk for fracture, defined as<br>a history of osteoporotic fracture, or multiple risk factors for fracture; or patients who have failed or are<br>intolerant to other available osteoporosis therapy.<br>Limitations of Use: Limit duration of use to 12 monthly doses. If osteoporosis therapy remains warranted,<br>continued therapy with an anti-resorptive agent should be considered  | 420                               | Not for use in<br>premenopausal<br>women. | N/A         | Females Only           | Y               | Y                               |          | 10/3/2019             |
| Drugs          | J3121         | Injection, testosterone<br>enanthate, 1 mg                       | 1 mg                       | 1/1/2015                | N/A        | testosterone enanthate injection, solution                     | Indicated for replacement therapy in conditions associated with a deficiency or absence of endogenous<br>testosterone including primary hypogonadism (congenital or acquired), hypogonadotropic hypogonadism<br>(congenital or acquired), and delayed puberty. Testosterone frannthates injection may be used secondarily<br>in women with advancing inoperable metastatic (skeletal) mammary cancer who are 1 – 5 years<br>postmenopausal.   | 1,200                             | N/A                                       | N/A         | N/A                    | Y               | ¥                               |          | 9/12/2018             |
| Drugs          | J3145         | Injection, testosterone<br>undecanoate, 1mg                      | 1 mg                       | 1/1/2015                | Aveed®     | testosterone undecanoate<br>injection for intramuscular<br>use | Indicated for testosterom replacement therapy in adult males for conditions associated with a deficiency<br>or absence of endogenous testosterome:<br>primary hypogenadism (congenital or acquired) or hypogenadotropic hypogenadism (congenital or<br>acquired).<br>Limitations of Use:<br>• Safety and efficacy of Aveed in men with "age-related hypogenadism" have not been established.<br>• Safety and efficacy of Aveed in males less than 18 years old have not been established.   | 1,500                             | 18 years                                  | N/A         | Males Only             | Y               | ¥                               |          | 9/21/2018             |
| Drugs          | J3230         | Injection, chlorpromazine HCl,<br>up to 50 mg                    | 50 mg                      | 1/1/2000                | N/A        | chlorpromazine<br>hydrochloride injection                      | Indicated for the treatment of schizophrenia; to control nausea and vomiting; for relief of restlessness and<br>apprehension before surgery; for acute intermittent porphyria; as an adjunct in the treatment of tetanus;<br>to control the manifestations of the manic type of manic-depressive illness; for relief of intractable<br>hiccups; for the treatment of severe behavioral problems in children (1 to 12 years of age) marked by<br>combativeness and/or explosive hyperexcitable behavior (out of proportion to immediate provocations),<br>and in the short-term treatment of hyperactive children who show excessive motor activity with<br>accompanying conduct disorders consisting of some or all of the following symptoms: implicitly,<br>difficulty sustaining attention, aggressivity, mood lability, and poor frustration tolerance.   | 248                               | 6 months                                  | N/A         | N/A                    | Y               | ¥                               |          | 9/27/2018             |
| Biologicals    | J3240         | Injection, thyrotropin alpha,<br>0.9 mg, provided in 1.1 mg vial | 0.9 mg                     | 1/1/2003                | Thyrogen®  | thyrotropin alfa for injection<br>for intramuscular use        | Indicated for:<br>Diagnostic: Use as an adjunctive diagnostic tool for serum thyroglobulin (Tg) testing with or without<br>radiolodine imaging in the follow-up of patients with well-differentiated thyroid cancer who have<br>previously undergone thyroidectomy.<br>• Ablation: Use as an adjunctive treatment for radiolodine ablation of thyroid tissue remnants in patients<br>who have undergone a near-total or total thyroidectomy for well-differentiated thyroid cancer and who do<br>not have evidence of distant metastatic thyroid cancer.<br>Limitations of Use:<br>• Diagnostic:<br>• Thyrogen-stimulated Tg levels are generally lower than, and do not correlate with Tg levels after thyroid<br>hormone withdrawal.<br>- Serven when Thyrogen-Tg testing is performed in combination with radiolodine imaging, there remains a<br>risk of missing a diagnosis of thyroid cancer or underestimating the extent of the disease.<br>- Anti-Tg Antibodies may confound the Tg assay and render Tg levels uninterpretable.<br>• Ablation:<br>- The effect of Thyrogen on long term thyroid cancer outcomes has not been determined. | 2                                 | 18 years                                  | N/A         | N/A                    | Y               | ¥                               |          | 6/19/2023             |
| Biologicals    | J3241         | Injection, teprotumumab-<br>trbw, 10 mg                          | 10 mg                      | 10/1/2020               | Tepezza®   | teprotumumab-trbw for injection, for intravenous use           | Indicated for the treatment of Thyroid Eye Disease regardless of Thyroid Eye Disease activity or duration.  | 600                               | 18 years                                  | N/A         | N/A                    | Y               | Y                               |          | 5/25/2023             |
| Drugs          | J3243         | Injection, tigecycline, 1 mg                                     | 1 mg                       | 1/1/2007                | Tygacil®   | tigecycline for injection, for intravenous use                 | Indicated in patients 18 years of age and older for:<br>• Complicated sin and skin structure infections<br>• Complicated intra-abdominal infections<br>• Community-acquired bacterial pneumonia<br>Limitations of Use: Tygacil is not indicated for treatment of diabetic foot infection or hospital-acquired<br>pneumonia, including ventilator-acsociated pneumonia.  | 1,450                             | 18 years                                  | N/A         | N/A                    | Y               | Ŷ                               |          | 9/21/2018             |

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| medicaid/medi | caid-ncci-eo  | dit-files   |                            |                         |            |   |  |                                   |   |             |                        |                 |                                 |   |                       |
|---------------|---------------|---|----------------------------|-------------------------|------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|               |               | Injection, tigecycline (accord)                                   |                            |                         |            |   | Indicated in patients 18 years of age and older for:<br>• Complicated skin and skin structure infections<br>• Complicated intra-abdominal infections<br>• Community-acquired bacterial pneumonia   |                                   |   |             |                        |                 |                                 |   |                       |
| Drugs         | J3244         | not therapeutically equivalent<br>to j3243, 1 mg                  |                            | 1/1/2023                | N/A        | tigecycline for injection, for<br>intravenous use (Accord)                  | Limitations of Use: Tigecycline for injection is not indicated for treatment of diabetic foot infection or<br>hospital-acquired pneumonia, including ventilator-associated pneumonia.  | 1,450                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 12/12/2022            |
|               |               |   |                            |                         |            |   | To reduce the development of drug-resistant bacteria and maintain the effectiveness of tigecycline for<br>injection and other antibacterial drugs, Tigecycline for injection should be used only to treat infections that<br>are proven or strongly suspected to be caused by bacteria.  |                                   |   |             |                        |                 |                                 |   |                       |
| Biologicals   | J3247         | Injection, secukinumab,<br>intravenous, 1 mg                      | 1 mg                       | 7/1/2024                | Cosentyx®  | secukinumab injection, for<br>intravenous use                               | Secukinumab intravenous injection is indicated for the treatment of:<br>- Adults with active parvial enthritis (PA)<br>- Adults with active ankylosing spondylitis (AS).<br>- Adults with active non-radiographic axial spondyloarthritis (n-axSpA) with objective signs of<br>inflammation.   | 1,125                             | 18 years  | N/A         | N/A                    | Y               | Y                               | 3/2024: Removal of<br>subcutaneous formulations<br>from PADP effective<br>3/31/2024 per DHB request<br>3/20/2024.   | 6/24/2024             |
| Drugs         | J3250         | Injection, trimethobenzamide<br>HCl, up to 200 mg                 | up to 200 mg               | 1/1/2000                | Tigan®     | trimethobenzamide<br>hydrochloride  | Indicated for the treatment of postoperative nausea and vomiting and for nausea associated with<br>gastroenteritis.  | 124                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 9/12/2018             |
| Drugs         | J3260         | Injection, tobramycin sulfate,<br>up to 80 mg                     | up to 80 mg                | 1/1/2000                | N/A        | tobramycin sulfate injection  | Indicated for the treatment of serious bacterial infections caused by susceptible strains of the designated<br>microorganisms in the diseases listed below:<br>• Septicemia in the neonate, child, and adult caused by P. aeruginosa, E. coli, and Klebsiella ap.<br>• Lower reginitarcy tract infections caused by P. aeruginosa, Kebsiella ap. Enterobacter sp. Serratia sp, E.<br>coli, and S. aureus (penkillinase and non-penkillinase-producing strains)<br>• Serious central nervous system infections (menaignits) caused by susceptible organisms<br>• Intra-badominal infections, including peritoritis, caused by F. coli, Klebsiella sp, and Enterobacter sp.<br>• Sini, bone, and skin-structure infections caused by P. aeruginosa, Proteus sp, E. coli, Klebsiella sp, Enterobacter sp.<br>• Enterobacter sp, and S. aureus | 558                               | N/A   | N/A         | N/A                    | Y               | Y                               |   | 9/12/2018             |
| Biologicals   | J3262         | Injection, tocilizumab, 1 mg                                      | 1 mg                       | 1/1/2011                | Actemra®   | tocilizumab injection, for<br>intravenous use                               | Indicated for the treatment of:<br>- Adult patients with moderately to severely active rheumatold arthritis (RA) who have had an inadequate<br>response to one or more Disease-Modifying Anti-Rheumatic Drugs (DMARDS).<br>- Active systemic juvenie idiopatici arthritis in patients two years of age and older.<br>- Active polyarticular juvenie idiopathic arthritis in patients two years of age and older.<br>- Adult and pediatric patients 2 years of age and older with chimeric antigen receptor (CAR) T cell-induced<br>severe or life-threatening cytokine release syndrome.<br>- Adult patients with giant cell arteritis.  | 3,200                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• 2 years of age and older:<br>systemic juvenile idiopathic<br>arthritis, polyarticular juvenile<br>idiopathic arthritis, CAR T cell-<br>induced CRS<br>• 18 years of age and older:<br>rheumatoid arthritis, giant cell<br>arteritis | 3/17/2022             |
| Biologicals   | J3263         | Injection, toripalimab-tpzi, 1<br>mg                              | 1 mg                       | 7/1/2024                | Loqtorzi™  | toripalimab-tpzi injection, for<br>intravenous use                          | recurrent locally advanced nasopharyngeal carcinoma (NPC).<br>• as a single agent for the treatment of adults with recurrent unresectable or metastatic NPC with disease<br>progression on or after a platinum-containing chemotherapy.  | 1,440                             | 18 years  | N/A         | N/A                    | Ŷ               | Ŷ                               |   | 6/24/2024             |
| Drugs         | J3285         | Injection, treprostinil, 1 mg                                     | 1 mg                       | 1/1/2006                | Remodulin* | treprostinil injection, for<br>subcutaneous or intravenous<br>use           | Indicated for treatment of pulmonary arterial hypertension (PAH) (WHO Group 1) to diminish symptoms<br>associated with exercise and to reduce the rate of clinical deterioration in patients requiring transition<br>from epoprostenol.  | 1,813                             | 17 years  | N/A         | N/A                    | Y               | Y                               |   | 5/14/2019             |
| Drugs         | J3299         | Injection, triamcinolone acetonide (xipere), 1 mg                 | 1 mg                       | 1/1/2000                | Xipere™    | triamcinolone acetonide<br>injectable suspension, for<br>suprachoroidal use | Indicated for the treatment of macular edema associated with uveitis.  | 80                                | 18 years  | N/A         | N/A                    | Ŷ               | Ŷ                               |   | 6/6/2022              |
| Drugs         | J3300         | Injection, triamcinolone<br>acetonide, preservative free, 1<br>mg | 1 1 mg                     | 1/1/2009                | Triesence® | triamcinolone acetonide<br>injectable suspension                            | Indicated for:<br>• Treatment of the following ophthalmic diseases: sympathetic ophthalmia, temporal arteritis, uveitis,<br>and ocular inflammatory conditions unresponsive to topical corticosteroids.<br>• Visualization during vitrectomy   | 8                                 | N/A   | N/A         | N/A                    | Y               | Y                               |   | 6/7/2019              |

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| medicaid/med | icaid-ncci-ec | dit-files  | 1                          |                         | 1                                | 1  |   |                                   |   | 1           |                        |                 | Rebating                        |   | т — т                 |
|--------------|---------------|--|----------------------------|-------------------------|----------------------------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                       | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Drugs        | J3301         | Injection, triamcinolone<br>acetonide, Not Otherwise<br>Specified, per 10 mg                                       | 10 mg                      | 1/1/2000                | Kenalog-10*,<br>Kenalog-40*      | triamcinolone acetonide<br>ingicetable suspension, for<br>intra-articular or intralesional<br>use only | Idenational and a set of the set | 150                               | N/A   | N/A         | N/A                    | ¥               | ¥                               |   | 9/12/2018             |
| Drugs        | J3304         | Injection, triamcinolone<br>acetonide, preservative-free,<br>extended-release,<br>microsphere formulation, 1<br>mg | 1 mg                       | 1/1/2019                | Zilretta™                        | triamcinolone acetonide<br>extended-release injectable<br>suspension, for intra-articular<br>use       | Indicated as an intra-articular injection for the management of osteoarthritis pain of the knee.<br>Limitation of Use: Zilretta is not intended for repeat administration.  | 64                                | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 9/12/2018             |
| Drugs        | J3315         | Injection, triptorelin pamoate,<br>3.75 mg   | 3.75 mg                    | 1/1/2003                | Trelstar®                        | triptorelin pamoate for<br>injectable suspension   | Indicated for the palliative treatment of advanced prostate cancer.   | 6                                 | 18 years  | N/A         | Males Only             | Y               | Y                               |   | 2/19/2024             |
| Drugs        | J3316         | Injection, triptorelin, extended release, 3.75 mg  | 3.75 mg                    | 1/1/2019                | Triptodur™                       | triptorelin for extended-<br>release injectable suspension,<br>for intramuscular use                   | Indicated for the treatment of pediatric patients 2 years and older with central precocious puberty.  | 6                                 | 2 years   | N/A         | N/A                    | Y               | Ŷ                               |   | 2/19/2024             |
| Biologicals  | J3357         | Ustekinumab, for<br>subcutaneous injection, 1 mg   | 1 mg                       | 1/1/2011                | Stelara® for<br>subcutaneous use | ustekinumab injection, for<br>subcutaneous use   | Indicated for the treatment of:<br>Adult patients with:<br>• Moderate to severe plaque psoriasis (Ps) who are candidates for phototherapy or systemic therapy<br>• Active psoriatic arthritis (PsA)<br>• Moderately to severely active crohn's disease (CD)<br>• Moderately to severely active uclerative collisis<br>Pediatric patients 6 to 17 years of age with:<br>• Moderate to severe plaque psoriasis, who are candidates for phototherapy or systemic therapy<br>• Active psoriatic arthritis (PsA)   | 180                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions.<br>• 6 years of age and older:<br>plaque psoriasis (Ps), psoriatic<br>arthritis (PsA)<br>• 18 years of age and older:<br>Crohn's disease (CD),<br>ulcerative colitis | 8/16/2022             |
| Biologicals  | J3358         | Ustekinumab, for intravenous injection, 1 mg   | 1 mg                       | 1/1/2018                | Stelara® for<br>intravenous use  | ustekinumab injection, for intravenous use   | Indicated for the treatment of adult patients with:<br>• Moderately to severely active Crohn's disease (CD)<br>• Moderately to severely active ulcerative collisis  | 520                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 12/3/2019             |
| Drugs        | J3360         | Injection, diazepam, up to 5<br>mg   | up to 5 mg                 | 1/1/2000                | N/A                              | diazepam injection   | Indicated:<br>• For the management of anxiety disorders or for the short-term relief of the symptoms of anxiety.<br>Anxiety or tension associated with the stress of everyday life usually does not require treatment with an<br>anxiety or.<br>In acute alcohol withdrawal, diazepam may be useful in the symptomatic relief of acute agitation,<br>temor, impending or acute delirum tremens and hallucinosis.<br>• As an adjunct prior to endoscopic procedures if apprehension, anxiety or acute stress reactions are<br>present, and to diminish the patient's recall of the procedures.<br>• As a useful adjunct for the melled of skeletal muscle spasm due to reflex spasm to local pathology (such as<br>filammation of the muscles or joints, or secondary to trauma); spasitity caused by upper motor neuron<br>disorders (such as cerebral palsy and paraplegia); athetosis; suff-man syndrome; and tetanus.<br>• As a useful adjunct in status eglifeticus and severe recurrent convulves esizures.<br>• As a useful adjunct in status eglifeticus and severe recurrent for values esizures<br>• As a useful adjunct in status eglifeticus and severe recurrent convulves esizures.<br>• As a useful adjunct in status eglifeticus and severe recurrent convulves esizures.<br>• As a useful adjunct in status eglifeticus and severe recurrent convulves esizures.<br>• As a useful adjunct in status eglifeticus and severe recurrent convulves esizures.  | 250                               | 31 days   | N/A         | N/A                    | ¥               | Ÿ                               |   | 10/10/2018            |

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| nedicaid/medi | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit    | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|---------------|---------------|--|-------------------------------|-------------------------|------------|--|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Drugs         | J3370         | Injection, vancomycin HCI, 500<br>mg   | 500 mg                        | 1/1/2000                | N/A        | vancomycin hydrochloride for<br>injection, USP for intravenous<br>use            | Indicated for the treatment of serious or severe infections caused by susceptible strains of methicillin-<br>resistant (B-lactam-resistant) staphylococci. It is indicated for penicillin-allergic patients, for patients who<br>cannot receive or who have failed to respond to other drugs, including the penicillins or cephalosporins,<br>and for infections caused by vancomycin-susceptible organisms that are resistant to tother antimicrobial<br>drugs. Vancomycin hydrochloride for injection is indicated for initial therapy when methicillin-resistant<br>staphylococci are suspected, but after susceptibility data are available, therapy should be adjusted<br>accordingly.<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of vancomycin<br>hydrochloride for injection USP and other antibiacterial drugs, vancomycin hydrochloride for injection<br>should be used only to treat or prevent infections that are proven or strongly suspected to be caused by<br>susceptible bacteria. When culture and susceptibility information are available, they should be considered<br>in selecting or modifying antibacterial therapy. In the absence of such data, local epidemiology and<br>susceptibility patterns may contribute to the empiric selection of therapy.<br>See package insert for list of infections. | 124                               | N/A         | N/A         | N/A                    | Y               | Ŷ                               |   | 6/8/2019              |
| Drugs         | J3371         | Injection, vancomycin hcl<br>(mylan), not therapeutically<br>equivalent to j3370, 500 mg   | 500 mg                        | 1/1/2023                | N/A        | vancomycin hydrochloride for<br>injection, for intravenous use<br>(Mylan)        | Indicated in adult and pediatric patients (neonates and older) for the treatment of:<br>• Septictemia<br>• Infective Fondocarditis<br>• Skin and Skin Structure Infections<br>• Bone Infections<br>• Lower Respiratory Tract Infections<br>To reduce the development of drug-resistant bacteria and maintain the effectiveness of Vancomycin<br>Hydrochloride for Injection and other antibacterial drugs, Vancomycin Hydrochloride for Injection should<br>be used only to treat or prevent Infections that are proven or strongly suspected to be caused by<br>susceptible bacteria.  | 124                               | N/A         | N/A         | N/A                    | Y               | Ŷ                               |   | 12/6/2022             |
| Drugs         | J3372         | Injection, vancomycin hcl<br>(xellia), not therapeutically<br>equivalent to j3370, 500 mg  | 500 mg                        | 1/1/2023                | N/A        | vancomycin injection, for<br>intravenous use (Xellia)                            | Indicated in adult and pediatric patients less than 18 years of age as follows:  • Vancomycin injection administered intravenously is indicated for the treatment of: • Septicemia • Infective Endocardlits • Skin and Skin Structure Infections • Bone Infections • Lower Respiratory Tract Infections • Lower Respiratory Tract Infections • Lower Respiratory Tract Infections • Unimitations of Use: • Vancomycin injection administered intravenously is not approved for the treatment of C. difficile- associated diarrhea and enterocolitis caused by susceptible isolates of Staphylococcus aureus because it is not effective. • Vancomycin injection administered orally is not approved for the treatment of septicemia, infective endocardlits, skin and skin structure infections, bone infections and lower respiratory tract Infections because it is not effective. To reduce the development of drug-resistant bacteria and maintain the effectiveness of Vancomycin infections and other antibacterial drugy, suspected to be caused by usceptible bacteria.   | 124                               | N/A         | N/A         | N/A                    | Y               | v                               |   | 12/6/2022             |
| Biologicals   | J3380         | Injection, vedolizumab,<br>intravenous, 1 mg   | 1 mg                          | 1/1/2016                | Entyvio®   | vedolizumab for injection, for<br>intravenous use                                | Indicated in adults for the treatment of:<br>• moderately to severely active ulcerative colitis (UC).<br>• moderately to severely active Crohn's disease (CD).  | 600                               | 18 years    | N/A         | N/A                    | Y               | Y                               | 4/2024: Subcutaneous<br>formulation removed from<br>coverage effective 3/31/2024<br>due to HCPCS code description<br>change effective 4/1/2024. | 3/22/2024             |
| Biologicals   | J3385         | Injection, velaglucerase alfa,<br>100 units  | 100 units                     | 1/1/2011                | VPRIV®     | velaglucerase alfa for<br>injection, for intravenous use                         | Indicated for long-term enzyme replacement therapy (ERT) for patients with type 1 Gaucher disease.  | 252                               | 4 years     | N/A         | N/A                    | Y               | Y                               |   | 6/8/2019              |
| Drugs         | J3396         | Injection, verteporfin, 0.1 mg   | 0.1 mg                        | 1/1/2005                | Visudyne®  | verteporfin for injection, for intravenous use                                   | Indicated for the treatment of patients with predominantly classic subfoveal choroidal neovascularization<br>due to age-related macular degeneration, pathologic myopia or presumed ocular histoplasmosis.  | 150                               | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 9/12/2018             |
| Biologicals   | J3397         | Injection, vestronidase alfa-<br>vjbk, 1 mg  | 1 mg                          | 1/1/2019                | Mepsevii™  | vestronidase alfa-vjbk<br>injection, for intravenous use                         | Indicated in pediatric and adult patients for the treatment of Mucopolysaccharidosis VII (MPS VII, Sly<br>syndrome).<br>Limitations of Use:<br>The effect of Mepsevii on the central nervous system manifestations of MPS VII has not been determined.  | 1,680                             | N/A         | N/A         | N/A                    | Y               | Y                               |   | 8/5/2021              |
| Biologicals   | 13398         | Injection, voretigene<br>neparvovec-rzyl, 1 billion<br>vector genomes  | 1 billion vector genomes (vg) | 1/1/2019                | Luxturna™  | voretigene neparvovec-rzyl<br>intraocular suspension for<br>subretinal injection | Indicated for the treatment of patients with confirmed biallelic RPE65 mutation-associated retinal<br>dystrophy. Patients must have viable retinal cells as determined by the treating physician(s).  | 300                               | 1 year      | N/A         | N/A                    | Y               | Y                               |   | 9/17/2021             |
| Biologicals   | J3401         | Beremagene geperpavec-svdt<br>for topical administration,<br>containing nominal 5 x 10^9<br>pfu/ml vector genomes, per<br>0.1 ml | 0.1 mL                        | 1/1/2024                | Vyjuvek™   |  | Indicated for the treatment of wounds in patients 6 months of age and older with dystrophic<br>epidermolysis bullosa with mutation(s) in the collagen type VII alpha 1 chain (COL7A1) gene.   | 125                               | 6 months    | N/A         | N/A                    | Y               | Y                               |   | 12/22/2023            |

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| Category    | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name              | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modifie<br>Date |
|-------------|---------------|---|----------------------------|-------------------------|-------------------------|--|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|----------------------|
| Drugs       | J3410         | Injection, hydroxyzine HCl, up<br>to 25 mg                    | up to 25 mg                | 1/1/2000                | Vistaril*               | hydroxyzine hydrochloride<br>injection for intramuscular<br>use  | <ul> <li>The total management of anxiety, tension, and psychomotor agitation in conditions of emotional stress requires in most instances a combined approach of psychotherapy and chemotherapy. Hydroxyine has been found to be particularly useful for this later phase of therapy in las ability to render the disturbed patient more amenable to psychotherapy in long term treatment of the psychoneurotic and psychotic, although is should not be used as the sole treatment of psychosis or of clearly demonstrated cases of depression.</li> <li>Also useful in alleviating the manifestations of anxiety and tension as in the preparation for dental procedures and in acute emotional problems. It has also been recommended for the management of anxiety associated with organic disturbances and a subtrack emotional problems. This as been recommended for the management of anxiety associated with organic disturbances and as adjunctive therapy in alcoholism and altergic conditions with storing emotional or locating. Such as an subma, chronic uritaria, and pruntus.</li> <li>Hydroxyzine hydrochloride intramuscular solution is useful in treating the following types of patients when intramuscular administration is indicated:</li> <li>The acute of chronic alcoholic with maviety withdrawal symptoms or delirium tremens.</li> <li>-As pre-and postoperative and pre- and postpartum adjunctive medication to permit reduction in narcotic dosage, allay anxiety and control enesis.</li> <li>Hydroxyzine hydrochloride has also demonstrated effectiveness in controlling nausea and vomiting, excluding nauses and vomiting of regenany.</li> <li>Hydroxyzine benefits the cardiac patient by its sbility to allay the associated anxiety and apprehension attendent to critari types of hera disease. Hydroxyzine is not known to interfere with the action of digitalis in any way and may be used concurrently with this agent.</li> </ul> | 240                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 10/26/2018           |
| Drugs       | J3420         | Injection, vitamin B-12<br>cyanocobalamin, up to 1,000<br>mcg | up to 1,000 mcg            | 1/1/2000                | N/A                     | cyanocobalamin injection,<br>USP (vitamin B-12)  | Indicated for vitamin B12 deficiencies due to malabsorption which may be associated with the following<br>conditions:<br>Addisonian (pernicious) anemia<br>• Gastrointestinal pathology, dysfunction, or surgery, including gluten enteropathy or sprue, small bowel<br>bacteria overgrowth, total or partial gastrectomy<br>• Fish tapeworm infestation<br>• Malignarcy of pancreas or bowel<br>• Folic acid deficiency  | 10                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 9/27/2018            |
| Drugs       | J3430         | Injection, phytonadione<br>(vitamin K) per 1 mg               | 1 mg                       | 1/1/2000                | Mephyton®               | phytonadione injectable<br>emulsion, USP   | Cyanocobalamin injection is also suitable for the vitamin B12 absorption test (Schilling test).<br>Indicated in the following coagulation disorders which are due to faulty formation of factors II, VII, IX and<br>X when caused by Vitamin X deficiency or interference with vitamin K activity:<br>• anticoagulant-induced prothrombin deficiency caused by coumarin or indanedione derivatives;<br>• prophylaxis and therapy of hemorrhagic disease of the newborn;<br>• hypoprothrombinemia due to antibacterial therapy;<br>• hypoprothrombinemia secondary to factors limiting absorption or synthesis of vitamin K, e.g.,<br>obstructive jaundice, billiany fistus, pure, ujecarative collis, celac disease, intestinal resection, cystic<br>fibrosis of the pancreas, and regional enterits;<br>• other drug-induced hypoprothrombinemia where it is definitely shown that the result is due to<br>interference with vitamin K metabolism, e.g., salicylates.  | 50                                | N/A         | N/A         | N/A                    | Y               | Y                               | 1        | 6/5/2019             |
| Biologicals | J3470         | Injection, hyaluronidase, up to<br>150 units                  | up to 150 units            | 1/1/2000                | Amphadase®              | hyaluronidase injection  | Indicated as an adjuvant:<br>• In subcutaneous fluid administration for achieving hydration.<br>• To increase absorption and dispersion of other injected drugs.<br>• In subcutaneous urography for improving resorption of radiopaque agents.  | 93                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/19/2023            |
| Biologicals | J3473         | Injection, hyaluronidase,<br>recombinant, 1 USP unit          | 1 USP unit                 | 1/1/2007                | Hylenex®<br>Recombinant | hyaluronidase human<br>injection, for infiltration use,<br>for interstitial use, for<br>intraocular use, for<br>metrobulbar use, for<br>retrobulbar use, for soft<br>tissue use, and for<br>subcutaneous use | Indicated as an:<br>• Adjuvant to increase the dispersion and absorption of other injected drugs.<br>• In subcutaneous fluid administration for achieving hydration.<br>• In subcutaneous urography for improving resorption of radiopaque agents.  | 2,250                             | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/19/2023            |
| Drugs       | J3475         | Injection, magnesium sulfate,<br>per 500 mg                   | 500 mg                     | 1/1/2000                | N/A                     | magnesium sulfate injection  | Indicated for replacement therapy in magnesium deficiency, especially in acute hypomagnesemia<br>accompanied by signs of tetany similar to those observed in hypocalcemia. In such cases, the serum<br>magnesium level is usually below the lower limit of normal (1.5 to 2.5 mEq/L) and the serum calcium level<br>is normal (4.3 to 5.3 mEq/L) or elevated. Magnesium sulfate injection is also indicated for the prevention<br>and control of seizures in pre-eclampsia and eclampsia, respectively and for use in hyperalimentation.  | 560                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/5/2019             |
| Drugs       | J3480         | Injection, potassium chloride,<br>per 2 mEq                   | 2 mEq                      | 1/1/2000                | N/A                     | potassium chloride injection   | Indicated for the treatment or prevention of hypokalemia when oral treatment is not feasible.   | 1,240                             | N/A         | N/A         | N/A                    | Y               | Y                               |          | 8/24/2018            |
| Drugs       | J3486         | Injection, ziprasidone<br>mesylate, 10 mg                     | 10 mg                      | 1/1/2004                | Geodon®                 | ziprasidone mesylate for<br>injection, for intramuscular<br>use  | Indicated for the acute treatment of agitation in schizophrenic patients.   | 124                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 3/17/2022            |

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| medicaid/med | HCPCS<br>Code | HCPCS Description                   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                                     | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|--------------|---------------|-------------------------------------|----------------------------|-------------------------|--|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Drugs        | J3489         | Injection, zoledronic acid, 1<br>mg | 1 mg                       | 1/1/2014                | Reclast*;<br>Zometa*                           | zoledronic acid injection, for<br>intravenous use             | Reclast is indicated for:<br>• Treatment in Circase bone mass in men with osteoporosis<br>• Treatment in Circase bone mass in men with osteoporosis<br>• Treatment on Circase bone in mean and women<br>Limitations of Use: Optimal duration of use has not been determined. For patients at low-risk for fracture,<br>consider drug discontinuation after 3 to 5 years of use.<br>Zometa is indicated for the treatment of:<br>• Hypercalcemic of malignancy.<br>• Patients with multiple myeloma and patients with documented bone metastases from solid turnors, in<br>conjunction with standard antineoplasic therapy. Prostate cancer should have progressed after treatment<br>with at least one hormonal therapy.<br>Limitations of Use: The safety and efficacy of Zometa has not been established for use in<br>hypergrachtropidism or on-turno-related hypercalcemia.   | 20                                | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 9/21/2018             |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | Aponvie™                                       | aprepitant injectable emulsion, for intravenous use           | Indicated for the prevention of postoperative nausea and vomiting (PONV) in adults.<br>Uminations of Use: Aponvie has not been studied for treatment of established nausea and vomiting.  | 160                               | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 3/16/2023             |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | Baxdela**                                      | delafloxacin for injection, for<br>intravenous use            | Indicated in adults for the treatment of acute bacterial skin and skin structure infections (ABSSSI) caused<br>by suceptible isolates of the following:<br>- Gram-positive organisms: Staphylococcus aureus (including methicillin-resistant [MRSA] and methicillin-<br>susceptible [MSA] loidate]. Staphylococcus aureus (including Streptococcus lugdunensis, Streptococcus<br>agalacitae, Streptococcus anginosus Group (including Streptococcus anginosus, Streptococcus<br>agalacitae, Streptococcus osciellatus). Streptococcus pagenese, and Enterococcus<br>finetremedius, and Streptococcus consolitatus, Staphylococcus and Enterococcus<br>findicated in adults for the treatment of community-acquired bacterial pneumoniae, (CABP) caused by the<br>following susceptible incircorganisms: Streptococcus pneumoniae, Staphylococcus aureus (methicillin-<br>susceptible [MSSA] isolates only), Klebsiella pneumoniae, Escherichia coli, Pseudomonas aeruginosa,<br>Haemophilus influenzae, Haemophilus parainfluenzae, Chlamydia pneumoniae, Legionella pneumophila,<br>and Mycoplasma pneumoniae. | 8,400                             | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |   | 12/3/2019             |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | Bludigo™                                       | indigotindisulfonate sodium<br>injection, for intravenous use |   | 40                                | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 10/20/2022            |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | Bridion®                                       | sugammadex injection, for<br>intravenous use                  | Indicated for the reversal of neuromuscular blockade induced by rocuronium bromide and vecuronium<br>bromide in adults undergoing surgery.  | 12,500                            | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 11/14/2019            |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | Cleviprex®                                     | clevidipine injectable<br>emulsion, for intravenous use       | Indicated for the reduction of blood pressure when oral therapy is not feasible or not desirable.   | 1,500                             | 18 years    | N/A         | N/A                    | Y               | Y                               |   | 10/4/2018             |
| Drugs        | J3490         | Unclassified drugs                  | 1 mL                       | 1/1/2000                | Defitelio®                                     | defibrotide sodium injection,<br>for intravenous use          | Indicated for the treatment of adult and pediatric patients with hepatic veno-occlusive disease (VOD), also<br>known as sinusoidal obstruction syndrome (SOS), with renal or pulmonary dysfunction following<br>hematopoletic stem-cell transplantation (HSCT).   | 1,395                             | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |   | 6/10/2019             |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | Depacon®                                       | valproate sodium, for<br>intravenous injection                | Indicated as an intravenous alternative in patients in whom oral administration of valproate products is<br>temporarily not feasible in the following conditions:<br>Monotherapy and adjunctive therapy of complex partial seizures and simple and complex absence<br>seizures; adjunctive therapy in patients with multiple seizure types that include absence seizures.   | 119,000                           | 2 years     | N/A         | N/A                    | Y               | Ŷ                               |   | 5/30/2019             |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg lidocaine USP<br>base | 1/1/2000                | Lidocaine (various<br>topical<br>formulations) | lidocaine (various topical formulations)                      | Indicated for production of anesthesia of accessible mucous membranes of the oropharynx. It is also<br>useful as an anesthetic lubricant for intubation and for the temporary relief of pain associated with minor<br>burns, including sunburn, abrasions of the skin, and insect bites.  | 31,000                            | N/A         | N/A         | N/A                    | Y               | Y                               |   | 10/26/2018            |
| Drugs        | J3490         | Unclassified drugs                  | 1 mg                       | 1/1/2000                | N/A  | nalmefene hydrochloride<br>injection                          | Indicated:<br>- for the complete or partial reversal of opioid drug effects, including respiratory depression, induced by<br>either natural or synthetic opioids<br>- in the management of known or suspected opioid overdose   | 20                                | 18 years    | N/A         | N/A                    | Ŷ               | Ŷ                               | 12/2023: Due to NDA product<br>Revex no longer being<br>marketed, recommended<br>dosing updated to align with<br>ANDA product Prescribing<br>Information and brand name<br>Revex updated to N/A<br>effective 6/22/2022. | 1/26/2024             |

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| Category    | HCPCS<br>Code | HCPCS Description      | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name            | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler | Comments  | Last Modified<br>Date |
|-------------|---------------|------------------------|----------------------------|-------------------------|-----------------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------|---|-----------------------|
| Drugs       | 13490         | Unclassified drugs     | 50 mL                      | 1/1/2000                | N/A                   | solution  | (see Package insert for hor FDK approved indication descriptions)<br>indicated in:<br>- The treatment of metabolic acidosis which may occur in severe renal disease, uncontrolled diabetes,<br>circulatory insufficiency due to shock or severe dehydration, extracorporeal circulation of blood, cardiac<br>arrest and severe primary lacits acidosis.<br>- The treatment of certain drug intoxications, including barbiturates (where dissociation of the barbiturate<br>protein complex is desired), in poisoning by<br>salkylates on methyl alcohol and in hemolytic reactions requiring alkalinization of the urine to diminish<br>nephrotoxicity of blood pigments.<br>- Severe diarnhea which is often accompanied by a significant loss of bicarbonate.<br>- Treatment of metabolic acidosis should, if possible, be superimposed on messures designed to control<br>the basic cause of the acidosis — e.g., insulin in uncomplicated diabetes, blood volume restoration in<br>shock. But since an appreciable time interval may elapse before all of the ancillary effects are brought<br>about, bicarbonate therapy is required in any form of metabolic acidosis where a rapid increase in<br>plasma total Content is circular. |                                   | N/A   | N/A         | N/A                    | Y               | <u>Required</u>     |   | 10/31/2018            |
| Drugs       | J3490         | Unclassified drugs     | 1 mg                       | 1/1/2000                | Noxafil®              | posaconazole injection, for<br>intravenous use                                      | dehydration, and in severe primary lactic acidosis or severe diabetic acidosis.<br>Indicated for the prophylaxis of invasive Aspergillus and Candida infections in patients who are at high risk<br>of developing these infections due to being severely immunocompromised, such as HSCT recipients with<br>GVHD or those with hematologic malignancies with prolonged neutroping from chemotherapy.<br>Indicated for the treatment of invasive aspergillosis in adults and pediatric patients 13 years of age and<br>older.   | 9,600                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                   | Indication specific age<br>restrictions:<br>Prophylaxis of invasive<br>Aspergillus and Candida<br>infections: 2 years of age and<br>older<br>Treatment of invasive<br>aspergillosis: 13 years of age<br>and older | 7/27/2021             |
| Drugs       | J3490         | Unclassified drugs     | 1 mg                       | 1/1/2000                | Opvee <sup>®</sup>    | nalmefene nasal spray   | Indicated for the emergency treatment of known or suspected overdose induced by natural or synthetic<br>opioids in adults and pediatric patients aged 12 years and older, as manifested by respiratory and/or<br>central nervous system depression.  | 27                                | 12 years  | N/A         | N/A                    | Y               | Y                   |   | 10/26/2023            |
| Drugs       | J3490         | Unclassified drugs     | 1 vial                     | 1/1/2000                | Prevymis <sup>®</sup> | letermovir injection, for<br>intravenous use  | Indicated for:<br>- orophylaxis of cytomegalowirus (CMV) infection and disease in adult CMV-seropositive recipients [R+] of<br>an allogeneic hematopoietic stem cell transplant (HSCT).<br>- prophylaxis of CMV disease in adult kidney transplant recipients at high risk (Donor CMV<br>seropositive/Recipient CMV seroegative [D+/R-].   | 31                                | 18 years  | N/A         | N/A                    | Y               | Ŷ                   |   | 7/26/2023             |
| Drugs       | J3490         | Unclassified drugs     | 1 mL                       | 1/1/2000                | Provayblue®           | methylene blue injection, for<br>intravenous use                                    | Indicated for the treatment of pediatric and adult patients with acquired methemoglobinemia. This<br>indication is approved under accelerated approval. Continued approval for this indication may be<br>contingent upon verification of clinical benefit in subsequent trials.  | 60                                | N/A   | N/A         | N/A                    | Y               | Y                   |   | 3/17/2022             |
| Drugs       | J3490         | Unclassified drugs     | 10 mg                      | 1/1/2000                | Revatio*              | sildenafil injection, for<br>intravenous use  | Indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group I) in adults to improve<br>exercise ability and delay clinical worsening. Studies establishing effectiveness were short-term (12 to 16<br>weeks), and included predominately patients with NYHA Functional Class II-III symptoms. Etiologies were<br>idiopathic (71%) or associated with connective tissue disease (25%).<br>Limitation of Use: Adding sildenafil to bosentan therapy does not result in any beneficial effect on exercise<br>capacity.  | 93                                | 3 years   | N/A         | N/A                    | Y               | Ŷ                   |   | 3/17/2022             |
| Drugs       | J3490         | Unclassified drugs     | 1 mg                       | 1/1/2000                | Rezipres*             | ephedrine hydrochloride<br>injection, for intravenous use                           | Indicated for the treatment of clinically important hypotension occurring in the setting of anesthesia.  | 1,457                             | 18 years  | N/A         | N/A                    | Y               | Y                   |   | 4/17/2022             |
| Drugs       | J3490         | Unclassified drugs     | 1 mcg                      | 1/1/2000                | Uptravi®              | selexipag for injection, for intravenous use  | indicated for the treatment of pulmonary arterial hypertension (PAH, WHO Group I) to delay disease<br>progression and reduce the risk of hospitalization for PAH.<br>Note: Use Uptravi for injection in patients who are temporarily unable to take oral therapy.  | 111,600                           | 18 years  | N/A         | N/A                    | Y               | Ŷ                   |   | 9/28/2021             |
| Drugs       | J3490         | Unclassified drugs     | 10 mg                      | 1/1/2000                | Vimpat®               | lacosamide injection, for<br>intravenous use  | Vimpat is indicated for:<br>• Treatment of partial-onset seizures in patients 1 month of age and older.<br>• Adjunctive therapy in the treatment of primary generalized tonic-clonic seizures in patients 4 years of<br>age and older.   | 1,240                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                   | Indication specific age<br>restrictions:<br>Partial-onset seizures: 1 month<br>of age and older<br>Primary generalized tonic-<br>clonic seizures: 4 years of age<br>and older                                     | 11/17/2021            |
| Drugs       | J3490         | Unclassified drugs     | 0.6 mg                     | 1/1/2000                | Zegalogue®            | dasiglucagon injection, for<br>subcutaneous use                                     | Indicated for the treatment of severe hypoglycemia in pediatric and adult patients with diabetes aged 6<br>years and above.  | 10                                | 6 years   | N/A         | N/A                    | Y               | Y                   |   | 7/27/2021             |
| Drugs       | J3490         | Unclassified drugs     | 1 mg                       | 1/1/2000                | Wainua™               | eplontersen injection, for<br>subcutaneous use                                      | years and adove:<br>Eplontersen injection is indicated for the treatment of the polyneuropathy of hereditary transthyretin-<br>mediated amyloidosis in adults.   | 45                                | 18 years  | N/A         | N/A                    | Y               | Y                   |   | 3/25/2024             |
| Biologicals | J3590         | Unclassified biologics | 11 mg (1 kit)              | 1/1/2002                | Cablivi®              | caplacizumab-yhdp for   | Inclused or the reactions of adult patients with acquired thrombotic thrombocytopenic purpura (aTTP),<br>in combination with plasma exchange and immunosuppressive therapy.  | 32                                | 18 years  | N/A         | N/A                    | Y               | Y                   |   | 3/26/2019             |
| Biologicals | J3590         | Unclassified biologics | per daily dose             | 1/1/2002                | Palforzia™            | peanut (Arachis hypogaea)<br>allergen powder-dnfp powder<br>for oral administration | Indicated for the mitigation of allergic reactions, including anaphylaxis, that may occur with accidental<br>exposure to peanut.<br>Limitation of Use: Not indicated for the emergency treatment of allergic reactions, including anaphylaxis.   | 31                                | 4 years   | N/A         | N/A                    | Y               | Ŷ                   | Initial Dose Escalation may be<br>administered to patients aged<br>4 through 17 years. Up-Dosing<br>and Maintenance may be<br>continued in patients 4 years<br>of age and older.                                  | 4/29/2020             |
| Biologicals | J3590         | Unclassified biologics | 0.5 mL                     | 1/1/2002                | Plegridy™             | peginterferon beta-1a<br>injection, for subcutaneous or<br>intramuscular use        | Indicated for the treatment of patients with relapsing forms of multiple sclerosis.  | 3                                 | 18 years  | N/A         | N/A                    | Y               | Y                   |   | 2/25/2021             |

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| nedicaid/medi | icaid-ncci-e  | dit-files  |                            |                         |            |  |   |                                   |   |             |                        |                 |                                 |  |                       |
|---------------|---------------|--|----------------------------|-------------------------|------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Biologicals   | J3590         | Unclassified biologics   | 50 mL                      | 1/1/2002                | Praxbind®  | idarucizumab injection, for<br>intravenous use   | Indicated in patients treated with Pradaxa when reversal of the anticoagulant effects of dabigatran is<br>needed:<br>• For emergency surgency/urgent procedures<br>• In life-threatening or uncontrolled bleeding   | 4                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 7/16/2018             |
| Biologicals   | J3590         | Unclassified biologics   | 1 IU                       | 1/1/2002                | Recothrom® | thrombin topical<br>(recombinant) lyophilized<br>powder for solution - for<br>topical use only                               | Indicated to aid hemostasis whenever oozing blood and minor bleeding from capillaries and small venules<br>is accessible and control of bleeding by standard surgical techniques is ineffective or impractical in adults<br>and pediatric populations greater than or equal to one month of age.  | 80,000                            | 1 month   | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Biologicals   | J3590         | Unclassified biologics   | 1 mg                       | 1/1/2002                | Revcovi™   | elapegademase-lvlr injection,<br>for intramuscular use   | Indicated for the treatment of adenosine deaminase severe combined immune deficiency (ADA-SCID) in<br>pediatric and adult patients.   | 288                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 12/28/2018            |
| Biologicals   | J3590         | Unclassified biologics   | 1 mg                       | 1/1/2002                | Strensiq®  | asfotase alfa injection, for<br>subcutaneous use   | Treatment of patients with perinatal/infantile-onset and juvenile-onset hypophosphatasia (HPP).   | 5,460                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Biologicals   | J3590         | Unclassified biologics   | 1 mg                       | 1/1/2002                | Tyenne®    | tocilizumab-aazg injection,<br>for intravenous use   | Tocilizumab-aarg injection is indicated for treatment of:<br>- Rheumatoid Arthritis (RA)<br>- Adult patients with moderately to severely active rheumatoid arthritis who have had an inadequate<br>response to one or more Disease-Modifying Anti-Rheumatic Drugs (DMARDs).<br>- Giant Cell Arteritis (CCA)<br>- Adult patients with giant cell arteritis.<br>- Polyarticular Juvenile Idiopathic Arthritis (PIA)<br>- Patients 2 years of age and older with active polyarticular juvenile Idiopathic arthritis.<br>- Systemic Juvenile Idiopathic Arthritis (SIA)<br>- Patients 2 years of age and older with active polyarticular juvenile Idiopathic arthritis.<br>- Systemic Juvenile Idiopathic Arthritis (SIA) | 1,600                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | ¥               | ¥                               | Indication specific:<br>RA, GCA: 18 years of age and<br>older<br>PJIA, SJIA: 2 years of age and<br>older | 5/3/2024              |
| Drugs         | J7030         | Infusion, normal saline<br>solution, 1,000 cc  | 1,000 cc                   | 1/1/2000                | N/A        | normal saline solution 1,000<br>cc (sodium chloride injection)   | Indicated as a source of water and electrolytes. Also indicated for use as a priming solution in  | N/A                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/26/2018            |
| Drugs         | J7040         | Infusion, normal saline solution, sterile  | 500 mL                     | 1/1/2000                | N/A        | normal saline solution 500 cc<br>(sodium chloride injection)   | Indicated as a source of water and electrolytes. Also indicated for use as a priming solution in<br>hemodialysis procedures.  | 186                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/7/2019              |
| Drugs         | J7042         | 5% Dextrose/normal saline<br>(500 mL = 1 unit)   | 500 mL                     | 1/1/2000                | N/A        | dextrose 5% / normal saline  | Indicated for use in adults and pediatric patients as sources of calories and water for hydration.  | 200                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/10/2018            |
| Drugs         | J7050         | Infusion, normal saline<br>solution, 250 cc  | 250 cc                     | 1/1/2000                | N/A        | normal saline solution 250 cc<br>(sodium chloride injection)   | Indicated as a source of water and electrolytes. Also indicated for use as a priming solution in<br>hemodialysis procedures.  | 186                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/7/2019              |
| Drugs         | J7060         | 5% Dextrose/water (500 mL =<br>1 unit)   | 500 mL                     | 1/1/2000                | N/A        | dextrose 5% / water  | Indicated for use in adults and pediatric patients as sources of calories and water for hydration.  | 200                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/10/2018            |
| Drugs         | J7070         | Infusion, D5W, 1,000 cc  | 1,000 cc                   | 1/1/2000                | N/A        | D5W (dextrose injection)   | Indicated for parenteral replenishment of fluid and minimal carbohydrate calories as required by clinical<br>condition of the patient.  | 124                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/4/2018             |
| Drugs         | J7120         | Ringer's lactate infusion, up to<br>1,000 cc   | up to 1,000 cc             | 1/1/2000                | N/A        | lactated ringer's infusion   | Indicated as a source of water and electrolytes or as an alkalinizing agent.  | 124                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 8/29/2018             |
| Drugs         | J7121         | 5% dextrose in lactated ringers<br>infusion, up to 1,000 cc                                      | up to 1,000 cc             | 1/1/2016                | N/A        | D5LR (5% dextrose in<br>lactated ringer's injection)   | Indicated for parenteral replacement of extracellular losses of fluid and electrolytes, with or without<br>minimal carbohydrate calories, as required by the clinical condition of the patient.   | 124                               | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/4/2018             |
| Biologicals   | J7165         | Injection, prothrombin<br>complex concentrate, human-<br>lans, per i.u. of factor ix<br>activity | 1 IU                       | 4/1/2024                | Balfaxar®  | prothrombin complex<br>concentrate, human-lans<br>lyophilized powder for<br>solution, for intravenous use                    | Prothrombin complex concentrate, human-lans is a blood coagulation factor replacement product<br>indicated for the urgent reversal of acquired coagulation factor deficiency induced by Vitamin K antagonist<br>(VKA, e.g., warfarin) therapy in adult patients with need for an urgent surgery/invasive procedure.   | 5,000                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 3/22/2024             |
| Biologicals   | J7168         | Prothrombin complex<br>concentrate (human), kcentra,<br>per i.u. of factor ix activity           | 1 IU                       | 7/1/2021                | Kcentra®   | prothrombin complex<br>concentrate (human) for<br>intravenous use, lyophilized<br>powder for reconstitution                  | Indicated for the urgent reversal of acquired coagulation factor deficiency induced by Vitamin K antagonist<br>(VKA, e.g., warfarin) therapy in adult patients with acute major bleeding or need for an urgent<br>surgery/invasive procedure.   | 5,000                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 6/28/2021             |
| Biologicals   | J7169         | Injection, coagulation factor xa<br>(recombinant), inactivated-<br>zhzo (andexxa), 10 mg         | 10 mg                      | 7/1/2020                | Andexxa®   | coagulation factor Xa<br>(recombinant), inactivated-<br>zhzo lyophilized powder for<br>solution for intravenous<br>injection | Indicated for patients treated with rivaroxaban and apixaban, when reversal of anticoagulation is needed<br>due to life-threatening or uncontrolled bleeding.   | 180                               | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 6/17/2020             |
| Biologicals   | J7170         | Injection, emicizumab-kxwh,<br>0.5 mg  | 0.5 mg                     | 1/1/2019                | Hemlibra*  | emicizumab-kxwh injection,<br>for subcutaneous use   | Indicated for routine prophylaxis to prevent or reduce the frequency of bleeding episodes in adult and<br>pediatric patients ages newborn and older with hemophilia A (congenital factor VIII deficiency) with or<br>without factor VIII inhibitors.  | 5,040                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 7/2/2018              |
| Biologicals   | J7171         | Injection, adamts13,<br>recombinant-krhn, 10 iu  | 10 IU                      | 7/1/2024                | Adzynma    | ADAMTS13, recombinant-<br>krhn lyophilized powder for<br>injection, for intravenous use                                      |   | 3,000                             | 2 years   | N/A         | N/A                    | Y               | Y                               |  | 6/24/2024             |
| Biologicals   | J7175         | Injection, factor X, (human), 1<br>IU  | 1 IU                       | 1/1/2017                | Coagadex*  | coagulation factor X (human)<br>lyophilized powder for<br>solution for intravenous<br>injection                              | Indicated in adults and children with hereditary Factor X deficiency for:<br>• On-demand treatment and control of bleeding episodes<br>• Perioperative management of bleeding in patients with mild, moderate and severe hereditary Factor X<br>deficiency<br>• Routine prophylaxis to reduce the frequency of bleeding episodes  | 84,000                            | N/A   | N/A         | N/A                    | Y               | Y                               |  | 5/25/2023             |
| Biologicals   | J7177         | Injection, human fibrinogen<br>concentrate (fibryga), 1 mg                                       | 1 mg                       | 1/1/2019                | Fibryga®   | fibrinogen (human)<br>lyophilized powder for<br>reconstitution, for<br>intravenous use                                       | Indicated for the treatment of acute bleeding episodes in adults and children with congenital fibrinogen<br>deficiency, including afibrinogenemia and hypofibrinogenemia. Fibryga is not indicated for<br>dysfibrinogenemia.  | 9,800                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 11/29/2021            |
| Biologicals   | J7178         | Injection, human fibrinogen<br>concentrate, not otherwise<br>specified, 1 mg                     | 1 mg                       | 1/1/2013                | RiaSTAP®   | fibrinogen concentrate<br>(human) for intravenous use,<br>lyophilized powder for<br>reconstitution                           | Indicated for the treatment of acute bleeding episodes in patients with congenital fibrinogen deficiency,<br>including afibrinogenemia and hypofibrinogenemia.  | 9,800                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/8/2019              |

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| medicaid/med | icaid-ncci-ed | dit-files  |                            |                         |                              |   |  |                                   |   |             |                        |                 |                                 |  |                       |
|--------------|---------------|--|----------------------------|-------------------------|------------------------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                   | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Biologicals  | J7179         | Injection, Von Willebrand<br>factor (recombinant),<br>(Vonvendi), 1IU VWF:RCo                            | 1 IU                       | 1/1/2017                | Vonvendi®                    | von Willebrand factor<br>(recombinant) lyophilized<br>powder for solution, for<br>intravenous injection                               | Indicated for use in adults (age 18 and older) diagnosed with von Willebrand disease (VWD) for:<br>• On-demand treatment and control of bleeding episodes.<br>• Perioperative management of bleeding.<br>• Routine prophylaxis to reduce the frequency of bleeding episodes in patients with severe Type 3 von<br>Willebrand disease receiving on-demand therapy.  | 254,800                           | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 2/11/2022             |
| Biologicals  | J7180         | Injection, factor XIII<br>(antihemophilic factor,<br>human), 1 IU  | 1 IU                       | 1/1/2012                | Corifact                     | factor XIII concentrate<br>(human) injection for<br>intravenous use   | Indicated for adult and pediatric patients with congenital Factor XIII deficiency for:<br>• Routine prophylactic treatment<br>• Peri-operative management of surgical bleeding.  | 10,000                            | N/A   | N/A         | N/A                    | Y               | Y                               |  | 10/10/2018            |
| Biologicals  | J7181         | Injection, factor XIII A-subunit,<br>(recombinant), per IU   | per IU                     | 1/1/2015                | Tretten®                     | coagulation factor XIII a-<br>subunit (recombinant)   | Indicated for routine prophylaxis of bleeding in patients with congenital factor XIII A-subunit deficiency.<br>Not for use in patients with congenital factor XIII B-subunit deficiency.   | 9,800                             | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/8/2019              |
| Biologicals  | J7182         | Injection, factor VIII,<br>(antihemophilic factor,<br>recombinant), (Novoeight),<br>per IU               | 1 IU                       | 1/1/2015                | Novoeight®                   | antihemophilic factor<br>(recombinant) for<br>intravenous injection<br>lyophilized powder for<br>solution                             | Adults and children with hemophilia A for: Control and prevention of bleeding; Perioperative management<br>Routine prophylaxis to prevent or reduce the frequency of bleeding episodes.  | 168,000                           | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/6/2019              |
| Biologicals  | J7183         | Injection, Von Willebrand<br>factor complex (human),<br>Wilate, 1 IU VWF:RCO                             | 1 IU VWF:RCO               | 1/1/2012                | Wilate®                      | von willebrand<br>factor/coagulation factor VIII<br>complex (human) lyophilized<br>powder for solution for<br>intravenous injection   | Von Wilebrand disease:<br>Indicated in children and adults with von Willebrand disease for:<br>• On-demand treatment and control of bleeding episodes.<br>• Perioperative management of bleeding.<br>• Nuither prohydrasis to reduce the frequency of bleeding episodes.<br>- Willebrand disease.<br>Willebrand disease.<br>Hemophilia A:<br>Indicated in adolescents and adults with hemophilia A for:<br>• Routine prophylasis to reduce the frequency of bleeding episodes.<br>• Route prophylasis to reduce the frequency of bleeding episodes.                    | 90,000                            | N/A   | N/A         | N/A                    | Y               | Y                               |  | 2/16/2024             |
| Biologicals  | J7185         | Injection, factor VIII<br>(antihemophilic factor,<br>recombinant) (Xyntha), per IU                       | 1 IU                       | 1/1/2010                | Xyntha®                      | factor VIII (antihemophilic<br>factor, recombinant) for<br>intravenous injection  | <ul> <li>Indicated in adults and children with hemophilia A for control and prevention of bleeding episodes and<br/>for perioperative management.</li> <li>Indicated in adults and children with hemophilia A for routine prophylaxis to reduce the frequency of<br/>bleeding episodes.</li> <li>Wyntha is not indicated in patients with von Willebrand's disease.</li> </ul>   | 58,800                            | N/A   | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 9/21/2020             |
| Biologicals  | J7186         | Injection, antihemophilic<br>factor VIII/Von Willebrand<br>factor complex (human), per<br>factor VIII IU | 1 IU                       | 1/1/2009                | Alphanate®                   | antihemophilic factor/von<br>Willebrand factor complex<br>(human) lyophilized powder<br>for solution for intravenous<br>injection     | Indicated for:<br>• Control and prevention of bleeding in adult and pediatric patients with hemophilia A.<br>• Surgical and/or imasive procedures in adult and pediatric patients with von Willebrand Disease in whom<br>desmopressin (DDAVP) is either ineffective or contraindicated. It is not indicated for patients with severe<br>VWD (Type 3) undergoing major surgery.   | 133,250                           | N/A   | N/A         | N/A                    | Y               | Y                               | Max Units: Although the<br>monthly dose can exceed this<br>amount, use of higher doses<br>administered by a provider<br>must be supported with<br>adequate documentation<br>supplied to DMA and<br>established in the medical<br>record.   | 9/21/2018             |
| Biologicals  | J7187         | Injection, Von Willebrand<br>factor complex (Humate-P),<br>per IU, VWF:RCO                               | 110                        | 1/1/2007                | Humate-P*                    | antihemophilic factor/von<br>Wilebrand factor complex<br>(human), lyophitzed powder<br>for reconstitution for<br>intravenous use only | Indicated for:<br>• Hemophilia A – Treatment and prevention of bleeding in adults.<br>• Von Willebrand disease (VWD) – in adults and pediatric patients in the<br>(1) Treatment of spontaneous and trauma-induced bleeding episodes, and<br>(2) Prevention of vecssive bleeding during and after surgery.<br>This applies to patients with severe VWD as well as patients with mild to moderate VWD where the use of<br>desmopressin is known or suspected to be inadequate. Humate-P is not indicated for the prophylaxis of<br>spontaneous bleeding episodes in VWD. | 136,250                           | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | Indication specific age<br>restrictions:<br>eternophila A: 13 years of<br>age and older<br>Von Wilebrand disease<br>(VWD): None<br>Max Units: Although the daily<br>dose can exceed this amount,<br>use of higher doses<br>administered by a provider<br>must be supported with<br>adequate documentation<br>supplied to DM and<br>established in the medical<br>record. | 9/21/2018             |
| Biologicals  | J7188         | Injection, factor VIII<br>(antihemophilic factor,<br>recombinant), (Obizur), per IU                      | 1 IU                       | 1/1/2016                | Obizur®                      | antihemophilic factor<br>(recombinant), porcine<br>sequence lyophilized powder<br>for solution for intravenous<br>injection           | Treatment of bleeding episodes in adults with acquired hemophilia A.   | 630,000                           | 18 years  | N/A         | N/A                    | Y               | Y                               | Tecolo.  | 4/10/2019             |
| Biologicals  | J7189         | Factor viia (antihemophilic<br>factor, recombinant),<br>(novoseven rt), 1 microgram                      | 1 mcg                      | 1/1/2006                | NovoSeven®,<br>NovoSeven® RT | coagulation factor VIIa<br>(recombinant) for<br>intravenous use   | Indicated for:<br>• Treatment of bleeding episodes and peri-operative management in adults and children with hemophilia<br>A or 8 with inhibitors, congenital Factor VII (FVII) deficiency, and Gianzmann's thrombasthenia with<br>refractorines to platelet transfictions, with or without antibidies to platelets.<br>• Treatment of bleeding episodes and peri-operative management in adults with acquired hemophilia.   | 96,000                            | N/A   | N/A         | N/A                    | Y               | Ŷ                               |  | 12/28/2020            |

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| Category    | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name   | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
|-------------|---------------|---|----------------------------|-------------------------|--|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Biologicals | J7190         | Factor VIII (antihemophilic<br>factor [human]) per IU   | 110                        | 1/1/2000                | Hemofil® M,<br>Koate®-DVI,<br>Monoclate-P®   | factor VIII (antihemophilic<br>factor, human) for<br>intravenous injection                      | Koate: indicated for the control and prevention of bleeding episodes or in order to perform emergency and<br>elective surgery in patients with hemophila A thereditary Factor VIII deficiency).<br>Limitation of Use: Koate is not indicated for the treatment of von Willebrand disease.<br>Monoclate-P: Indicated for treatment of classical hemophilia (Hemophilia A). Affected individuals<br>frequently require therapy following minor accidents. Surgery, when required in such individuals, must be<br>preceded by temporary corrections of the clotting abnormality. Surgical prophysics in severe AHF<br>deficiency can be accomplished with an appropriate/odsed pre-surgery and bolos of Monoclate-P followed<br>by intermittent maintenance doses. Monoclate P is not effective in controlling the bleeding of patients<br>with von Willebrand disease.<br>Hemofil M: Indicated in hemophilia A (classical hemophilia) for the prevention and control of hemorrhagic<br>episodes. Hemofil M is not indicated in von Willebrand disease.  | 24,000                            | N/A         | N/A         | N/A                    | Ŷ               | ¥                               |          | 10/10/2018            |
| Biologicals | J7192         | Factor VIII (antihemophilic<br>factor, recombinant) per IU,<br>not otherwise specified            | 110                        | 1/1/2000                | Advate*,<br>Bioclate*,<br>Helixate*,<br>Kogenate* FS,<br>Recombinate*'',<br>ReFacto* | factor VII (antihemophilic<br>factor, recombinant) for<br>intravenous use                       | Kogenate: Indicated for:<br>• On-demand treatment and control of bleeding episodes in adults and children with hemophilia A.<br>• Perioperative management of bleeding in adults and children with hemophilia A.<br>• Routine prophylaxis to reduce the frequency of bleeding episodes in children with hemophilia A and to<br>reduce the risk of joint damage in children without pre-existing joint damage.<br>• Routine prophylaxis to reduce the frequency of bleeding episodes in adults with hemophilia A.<br>Kogenate is not indicated for the treatment of von Willebrand disease.<br>Advate: Indicated for use in children and adults with hemophilia A for:<br>• Control and prevention of bleeding episodes.<br>• Perioperative management.<br>• Routine prophylaxis to prevent or reduce the frequency of bleeding episodes.<br>Advate: Indicated for the treatment of von Willebrand disease.<br>Recombinate: Indicated for the treatment of von Willebrand disease.<br>• Perioperative management.<br>• For the prevention and control of hemorrhagic episodes.<br>• Perioperative management.<br>• Recombinate: Indicated in von Willebrand's disease. | 54,000                            | N/A         | N/A         | N/A                    | Y               | Y                               |          | 10/10/2018            |
| Biologicals | J7193         | Factor IX (antihemophilic<br>factor, purified, non-<br>recombinant) per IU                        | 1 IU                       | 1/1/2002                | AlphaNine® SD,<br>Mononine®  | coagulation factor IX (human)   | Indicated for the prevention and control of bleeding episodes in patients with Factor IX deficiency<br>(hemophilia B, Christmas disease).   | 42,000                            | N/A         | N/A         | N/A                    | Y               | Y                               |          | 10/10/2018            |
| Biologicals | J7194         | Factor IX, complex, per IU  | per IU                     | 1/1/2000                | Bebulin® VH,<br>Profilnine® SD,<br>Profilnine®                                       | factor IX complex for intravenous administration  | Bebulin: Indicated for the prevention and control of bleeding episodes in adult patients with hemophilia B<br>(congenital Factor IX deficiency or Christmas disease). Bebulin is not indicated for use in the treatment of<br>Factor VII deficiency. No clinical studies have been conducted to show benefit from this product for<br>treating deficiencies other than Factor IX deficiency.<br>Profinine: Indicated for the prevention and control of bleeding in patients with factor IX deficiency<br>(hemophila B). Profilinine contains non-therapeutic levels of factor VII and is not indicated for use in the<br>treatment of factor VII deficiency.  | 59,500                            | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 10/26/2018            |
| Biologicals | J7195         | Injection factor IX<br>(antihemophilic factor,<br>recombinant) per IU, not<br>otherwise specified | 1 IU                       | 1/1/2002                | BeneFIX®   | coagulation factor IX<br>(recombinant) for<br>intravenous use                                   | Indicated for:<br>• Control and prevention of bleeding episodes in adult and pediatric patients with hemophilia B.<br>• Peri-operative management in adult and pediatric patients with hemophilia B.<br>Limitations of Use: Benefix is not indicated for the treatment of other factor deficiencies (e.g. factors II,<br>VI, VIII, adX, VI, menophilia A patients with inhibitors to factor VIII, reversal of coumarin-induced<br>anticoagulation, and bleeding due to low levels of liver-dependent coagulation factors.   | 42,000                            | N/A         | N/A         | N/A                    | Y               | Ŷ                               |          | 10/10/2018            |
| Biologicals | J7196         | Injection, antithrombin<br>recombinant, 50 IU   | 50 IU                      | 1/1/2011                | ATryn®   | antithrombin (recombinant)<br>lyophilized powder for<br>reconstitution                          | Indicated for the prevention of peri-operative and peri-partum thromboembolic events in hereditary<br>antithrombin deficient patients.  | 1,100                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/25/2018             |
| Biologicals | J7197         | Antithrombin III (human), per<br>IU   | 1 IU                       | 1/1/2000                | Thrombate III®   | antithrombin III (human)<br>lyophilized powder for<br>solution for intravenous<br>injection     | Indicated in patients with hereditary antithrombin deficiency for:<br>• Treatment and prevention of thromboembolism<br>• Prevention of peri-operative and peri-partum thromboembolism   | 40,000                            | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/25/2018             |
| Biologicals | J7198         | Anti-inhibitor, per IU  | per IU                     | 1/1/2000                | Feiba  | anti-inhibitor coagulant<br>complex, for intravenous use,<br>lyophilized powder for<br>solution | Indicated for use in hemophila A and B patients with inhibitors for:<br>• Control and prevention of bleeding episodes<br>• Perioperative management<br>• Routine prophylaxis to prevent or reduce the frequency of bleeding episodes.<br>Feiba is not indicated for the treatment of bleeding episodes resulting from coagulation factor deficiencies<br>in the absence of inhibitors to factor VIII or factor IX.  | 560,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 9/21/2018             |
| Biologicals | J7200         | Injection, factor IX,<br>(antihemophilic factor,<br>recombinant), Rixubis, per IU                 | 1 IU                       | 1/1/2015                | Rixubis®   | coagulation factor IX<br>(recombinant) for<br>intravenous injection                             | Indicated in adults and children with hemophilia B for control and prevention of bleeding episodes,<br>perioperative management, and routine prophylaxis. Rixubis is not indicated for induction of immune<br>tolerance in patients with Hemophilia B.  | 60,300                            | N/A         | N/A         | N/A                    | Y               | Y                               |          | 10/10/2018            |

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|---------------|---------------|---|----------------------------|-------------------------|------------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name             | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
| Biologicals   | J7201         | Injection, factor IX, Fc fusion<br>protein, (recombinant),<br>Alprolix, 1 IU                                  | 1 IU                       | 1/1/2017                | Alprolix®              | coagulation factor IX<br>(recombinant), Fc fusion<br>protein, lyophilized powder<br>for solution for intravenous<br>injection | Indicated for adults and children with hemophilia 8 for:<br>• On-demand treatment and control of bleeding episodes.<br>• Perioperative management of bleeding.<br>• Routine prophylaxis to reduce the frequency of bleeding episodes.<br>Limitations of Use: Alprolix is not indicated for induction of immune tolerance in patients with hemophilia<br>R   | 72,000                            | N/A         | N/A         | N/A                    | Y               | Y                               |          | 4/10/2019             |
| Biologicals   | J7202         | Injection, factor IX, albumin<br>fusion protein, (recombinant),<br>Idelvion, 1 IU                             | 1 IU                       | 1/1/2017                | Idelvion®              | coagulation factor IX<br>(recombinant), albumin<br>fusion protein lyophilized<br>powder for solution for<br>intravenous use   | or.<br>Indicated in children and adults with hemophilia B (congenital Factor IX deficiency) for:<br>• On-demand treatment and control and prevention of bleeding episodes<br>> Perioparative management of bleeding<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>Limitations of Use: Idelvion is not indicated for immune tolerance induction in patients with Hemophilia B.  | 96,921                            | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/6/2019              |
| Biologicals   | J7203         | Injection factor ix,<br>(antihemophilic factor,<br>recombinant), glycopegylated,<br>(rebinyn), 1 iu           | 1 IU                       | 1/1/2019                | Rebinyn®               | coagulation factor IX<br>(recombinant),<br>glycoPEGylated, lyophilized<br>powder for solution for<br>intravenous injection    | Indicated for use in adults and children with hemophilia B for:<br>• On-demand treatment and control of bleeding episodes<br>• Perioperative management of bleeding<br>Limitations of Use: Rebinyn is not indicated for routine prophylaxis in the treatment of patients with<br>hemophilia B or or immune tolerance induction in patients with hemophilia B.   | 67,200                            | N/A         | N/A         | N/A                    | Ŷ               | ¥                               |          | 7/2/2018              |
| Biologicals   | J7204         | Injection, factor viii,<br>antihemophilic factor<br>(recombinant), (esperoct),<br>glycopegylated-exei, per iu | 110                        | 7/1/2020                | Esperoct®              | antihemophilic factor<br>(recombinant),<br>glycopegylated-exei<br>lyophilized powder for<br>solution, for intravenous use     | Indicated for use in adults and children with hemophilia A for:<br>• On-demand treatment and control of bleeding episodes<br>> Perioperative management of bleeding<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>Limitation of Use: Esperoct is not indicated for the treatment of von Willebrand disease.  | 133,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/17/2020             |
| Biologicals   | J7205         | Injection, factor VIII Fc fusion<br>protein (recombinant), per IU   | 1 IU                       | 1/1/2016                | Eloctate®              | antihemophilic factor<br>(recombinant) Fc fusion<br>protein lyophilized powder<br>for solution for intravenous<br>injection   | Indicated in adults and chloren with Hemophila A (congenital Factor VIII deficiency) for:<br>• On-demand treatment and control of bleeding episodes.<br>• Perioperative management of bleeding.<br>• Routine prophylaxis to reduce the frequency of bleeding episodes.  | 140,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 7/2/2018              |
| Biologicals   | J7207         | Injection, factor VIII,<br>(antihemophilic factor,<br>recombinant), pegylated, 1 IU                           | 1 IU                       | 1/1/2017                | Adynovate <sup>®</sup> | antihemophilic factor<br>(recombinant), PEGylated<br>lyophilized powder for<br>solution for intravenous<br>injection          | Limitation of Use: Eloctate is not indicated for the treatment of yon Willebrand disease.<br>Indicated in children and adult patients with hemophilia A (congenital factor VIII deficiency) for:<br>• On-demand treatment and control of bleeding episodes<br>• Perioperative management<br>• Routine prophylasis to reduce the frequency of bleeding episodes<br>Adynovate is on indicated for the treatment of yon Willebrand disease.  | 210,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 9/25/2018             |
| Biologicals   | J7208         | Injection, factor viii,<br>(antihemophilic factor,<br>recombinant), pegylated-aucl,<br>(jivi), 1 i.u.         | 110                        | 7/1/2019                | Jivi®                  | antihemophilic factor<br>(recombinant) PEGylated-<br>aucl, for intravenous use  | Indicated for use in previously treated adults and adolescents (12 years of age and older) with hemophilia<br>A (congenital Factor VIII deficiency) for:<br>0 - Ondemand treatment and control of bleeding episodes<br>• Perioperative management of bleeding<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>Limitations of use:<br>- JiW is not indicated for use in children < 12 years of age due to a greater risk for hypersensitivity<br>reactions.<br>- JiW is not indicated for use in previously untreated patients (PUPs).<br>- JiW is not indicated for the restment of yow Wilebrand disease. | 180,000                           | 12 years    | N/A         | N/A                    | Y               | ¥                               |          | 9/25/2018             |
| Biologicals   | J7209         | Injection, factor VIII,<br>(antihemophilic factor,<br>recombinant), (Nuwiq), 1 IU                             | 1 IU                       | 1/1/2017                | Nuwiq®                 | antihemophilic factor<br>(recombinant), lyophilized<br>powder for solution for<br>intravenous injection                       | Indicated in adults and children with Hemophilia A for:<br>• On-demand treatment and control of bleeding episodes<br>> Perioperative management of bleeding<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>Nuwig is not indicated for the treatment of von Willebrand Disease.  | 210,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 4/10/2019             |
| Biologicals   | J7210         | Injection, factor VIII,<br>(antihemophilic factor,<br>recombinant), (Afstyla), 1 IU                           | 1 IU                       | 1/1/2018                | Afstyla®               | antihemophilic factor<br>(recombinant), single chain<br>for intravenous injection,<br>lyophilized powder for<br>solution      | Indicated in adults and children with hemophilia A (congenital Factor VIII deficiency) for:<br>• On-demand treatment and control of bleeding episodes.<br>• Soutine prophysias to reduce the frequency of bleeding episodes.<br>• Perioperative management of bleeding.<br>Limitation of Use:<br>Afstyla is not indicated for the treatment of von Wilebrand disease.   | 210,000                           | N/A         | N/A         | N/A                    | Y               | ¥                               |          | 4/10/2019             |
| Biologicals   | J7211         | Injection, factor VIII,<br>(antihemophilic factor,<br>recombinant), (Kovaltry), 1 IU                          | 1 IU                       | 1/1/2018                | Kovaltry®              | factor VIII (antihemophilic<br>factor, recombinant) for<br>intravenous injection  | Indicated for use in adults and children with hemophilia A (congenital Factor VIII deficiency) for:<br>• On-demand treatment and control of bleeding episodes<br>• Perioperative management of bleeding<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>Kovaltry is not indicated for the treatment of von Willebrand disease.   | 210,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 10/10/2018            |
| Biologicals   | J7212         | Factor viia (antihemophilic<br>factor, recombinant)-jncw<br>(sevenfact), 1 microgram                          | 1 mcg                      | 1/1/2021                | Sevenfact®             | [coagulation factor VIIa<br>(recombinant)-jncw]<br>lyophilized powder for<br>solution, for intravenous use                    | Indicated for the treatment and control of bleeding episodes occurring in adults and adolescents (12 years<br>of age and older) with hemophilia A or B with inhibitors.<br>Limitation of Use: Sevenfact is not indicated for treatment of congenital factor VII deficiency.   | 1,260,000                         | 12 years    | N/A         | N/A                    | Y               | Y                               |          | 12/28/2020            |
| Biologicals   | J7213         | Injection, coagulation factor ix<br>(recombinant), ixinity, 1 i.u.  | 110                        | 7/1/2023                | lxinity®               | coagulation factor IX<br>(recombinant) lyophilized<br>powder for solution for<br>intravenous injection                        | Indicated in adults and children (< 12 years of age) with hemophilia B for:<br>• On-demand treatment and control of bleeding episodes<br>• Perioperative management<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>binity is not indicated for induction of immune tolerance in patients with hemophilia B.   | 322,000                           | N/A         | N/A         | N/A                    | Y               | Y                               |          | 5/3/2024              |

### Physician Administered Drug Program Catalog

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
 The HCPCS Code effective date represents the date the HCPCS code was established

• Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

| medicaid/med        | icaid-ncci-eo | dit-files  |                            |                         |                        |  |   |                                   |                |             |                        |                 |                                 |          |                       |
|---------------------|---------------|--|----------------------------|-------------------------|------------------------|--|---|-----------------------------------|----------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Category            | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name             | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age    | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
| Biologicals         | J7214         | Injection, factor viii/von<br>willebrand factor complex,<br>recombinant (altuviiio), per<br>factor viii i.u. | 1 IU                       | 10/1/2023               | Altuviiio™             | antihemophilic factor<br>(recombinant), Fc-VWF-XTEN<br>fusion protein-ehtl,<br>lyophilized powder for<br>solution, for intravenous use | Indicated for use in adults and children with hemophilia A (congenital factor VIII deficiency) for:<br>• Routine prophylaxis to reduce the frequency of bleeding episodes<br>• On-demand treatment & control of bleeding episodes<br>• Perioperative management of bleeding<br>Limitation of Use:<br>Altuvinio is not indicated for the treatment of von Willebrand disease.  | 112,000                           | N/A            | N/A         | N/A                    | Y               | Y                               |          | 9/28/2023             |
| Drugs               | J7296         | Levonorgestrel-releasing<br>intrauterine contraceptive<br>system, (Kyleena), 19.5 mg                         | 19.5 mg                    | 1/1/2018                | Kyleena®               | levonorgestrel-releasing<br>intrauterine system  | Indicated for prevention of pregnancy for up to 5 years.  | 1                                 | After menarche | N/A         | Females Only           | Y               | Y                               |          | 10/26/2018            |
| Drugs               | J7297         | Levonorgestrel-releasing<br>intrauterine contraceptive<br>system (Liletta), 52mg                             | 52 mg                      | 1/1/2017                | Liletta®               | levonorgestrel-releasing<br>intrauterine system  | Indicated for the prevention of pregnancy for up to 8 years.<br>Indicated for treatment of heavy menstrual bleeding for up to 5 years in patients who choose intrauterine<br>contraception as their method of contraception.  | 1                                 | After menarche | N/A         | Females Only           | Y               | Y                               |          | 7/26/2023             |
| Drugs               | J7298         | Levonorgestrel-releasing<br>intrauterine contraceptive<br>system (Mirena), 52 mg                             | 52 mg                      | 1/1/2017                | Mirena®                | levonorgestrel-releasing<br>intrauterine system  | Indicated for:<br>• Pregnancy prevention for up to 8 years.<br>• Treatment of heavy menstrual bleeding in women who choose to use intrauterine contraception as their<br>method of contraception for up to 5 years.   | 1                                 | After menarche | N/A         | Females Only           | Y               | Y                               |          | 9/15/2022             |
| Miscellaneous       | J7300         | Intrauterine copper<br>contraceptive   | 1 intrauterine device      | 1/1/2000                | Paragard®              | intrauterine copper<br>contraceptive   | Indicated for intrauterine contraception for up to 10 years.  | 1                                 | 16 years       | N/A         | Females Only           | Y               | Y                               |          | 7/16/2018             |
| Drugs               | J7301         | Levonorgestrel-releasing<br>intrauterine contraceptive<br>system (Skyla), 13.5 mg                            | 13.5 mg                    | 1/1/2017                | Skyla®                 | levonorgestrel-releasing<br>intrauterine system  | Indicated for the prevention of pregnancy for up to 3 years.  | 1                                 | After menarche | N/A         | Females Only           | Y               | Y                               |          | 10/26/2018            |
| Drugs               | J7307         | Etonogestrel (contraceptive)<br>implant system, including<br>implant and supplies                            | 1 implant                  | 1/1/2008                | Nexplanon®             | etonogestrel implant for<br>subdermal use  | Indicated for use by women to prevent pregnancy.  | 1                                 | After menarche | N/A         | Females Only           | Y               | Y                               |          | 10/10/2018            |
| Drugs               | J7308         | Aminolevulinic acid HCl for<br>topical administration, 20%,<br>single unit dosage form (354<br>mg)           | 354 mg                     | 1/1/2004                | Levulan®<br>Kerastick® | aminolevulinic acid HCl for<br>topical solution, 20%   | Indicated for photodynamic therapy (treatment) of minimally to moderately thick actinic keratoses of the<br>face or scalp, or actinic keratoses of the upper extremities. FDA approval of upper extremity treatment<br>approved 3/6/2018.   | 1                                 | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 9/25/2018             |
| Drugs               | J7311         | Injection, fluocinolone<br>acetonide, intravitreal implant<br>(retisert), 0.01 mg                            | 0.01 mg                    | 1/1/2007                | Retisert <sup>®</sup>  | fluocinolone acetonide<br>intravitreal implant   | Indicated for the treatment of chronic noninfectious uveitis affecting the posterior segment of the eye.  | 118                               | 12 years       | N/A         | N/A                    | Y               | Y                               |          | 10/10/2018            |
| Drugs               | J7312         | Injection, dexamethasone,<br>intravitreal implant, 0.1 mg  | 0.1 mg                     | 1/1/2011                | Ozurdex <sup>®</sup>   | dexamethasone intravitreal<br>implant  | Indicated for the treatment of macular edema following branch retinal vein occlusion (BRVO) or central<br>retinal vein occlusion (CRVO), non-infectious uveitis affecting the posterior segment of the eye and<br>diabetic macular edema.   | 14                                | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 6/6/2019              |
| Drugs               | J7313         | Injection, fluocinolone<br>acetonide, intravitreal implant<br>(Iluvien), 0.01 mg                             | 0.01 mg                    | 1/1/2016                | lluvien®               | fluocinolone acetonide<br>intravitreal implant   | Indicated for the treatment of diabetic macular edema in patients who have been previously treated with<br>a course of corticosteroids and did not have a clinically significant rise in intraocular pressure.  | 38                                | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 10/16/2019            |
| Drugs               | J7314         | Injection, fluocinolone<br>acetonide, intravitreal implant<br>(Yutiq), 0.01 mg                               | 0.01 mg                    | 10/1/2019               | Yutiq™                 | fluocinolone acetonide<br>intravitreal implant 0.18 mg,<br>for intravitreal injection  | Indicated for the treatment of non-infectious uveitis affecting the posterior segment of the eye.   | 36                                | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 9/27/2019             |
| Drugs               | J7336         | Capsaicin 8% patch, per<br>square centimeter   | per square centimeter      | 1/1/2015                | Qutenza®               | capsaicin 8% patch   | Indicated for the management of neuropathic pain associated with postherpetic neuralgia (PHN).     Indicated for the treatment of neuropathic pain associated with diabetic peripheral neuropathy (DPN) of the feet.  | 1,120                             | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 8/25/2020             |
| Drugs               | J7342         | Installation, ciprofloxacin otic<br>suspension, 6 mg   | 6 mg                       | 1/1/2017                | Otiprio®               | ciprofloxacin otic suspension,<br>for intratympanic or otic use  |   | 10                                | 6 months       | N/A         | N/A                    | Y               | Y                               |          | 9/27/2018             |
| Drugs               | J7351         | Injection, bimatoprost,<br>intracameral implant, 1<br>microgram  | 1 mcg                      | 10/1/2020               | Durysta™               | bimatoprost implant, for intracameral administration   | Indicated for the reduction of intraocular pressure (IOP) in patients with open angle glaucoma (OAG) or<br>ocular hypertension (OHT).   | 20                                | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 9/21/2020             |
| Drugs               | J7352         | Afamelanotide implant, 1 mg  | 1 mg                       | 1/1/2021                | Scenesse*              | afamelanotide implant, for<br>subcutaneous use   | Indicated to increase pain free light exposure in adult patients with a history of phototoxic reactions from<br>erythropoietic protoporphyria (EPP).  | 16                                | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 11/17/2021            |
| Drugs               | J7354         | Cantharidin for topical<br>administration, 0.7%, single<br>unit dose applicator (3.2 mg)                     | 3.2 mg (1 ampule)          | 4/1/2024                | Ycanth™                | cantharidin topical solution   | Indicated for the topical treatment of molluscum contagiosum in adult and pediatric patients 2 years of<br>age and older.   | 4                                 | 2 years        | N/A         | N/A                    | Y               | Y                               |          | 3/22/2024             |
| Drugs               | J7355         | Injection, travoprost,<br>intracameral implant, 1<br>microgram   | 1 mcg                      | 7/1/2024                | iDose® TR              | travoprost intracameral<br>implant, for intracameral<br>administration   | Travoprost intracameral implant is indicated for the reduction of intraocular pressure (IOP) in patients<br>with open-angle glaucoma (OAG) or ocular hypertension (OHT).  | 150                               | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 6/24/2024             |
| Drugs               | J7402         | Mometasone furoate sinus<br>implant, (sinuva), 10<br>micrograms  | 10 mcg                     | 4/1/2021                | Sinuva™                | mometasone furoate sinus<br>implant  | Indicated for the treatment of chronic rhinosinusitis with nasal polyps in patients 2 18 years of age who<br>have had ethmoid sinus surgery.  | 270                               | 18 years       | N/A         | N/A                    | Y               | Y                               |          | 2/23/2023             |
| Immune<br>Globulins | J7504         | Lymphocyte immune globulin,<br>anti-thymocyte globulin,<br>equine, parenteral, 250 mg                        | 250 mg                     | 1/1/2000                | Atgam®                 | lymphocyte immune globulin,<br>anti-thymocyte globulin<br>(equine), sterile solution for<br>intravenous use only                       | Indicated for:<br>•Renal transplant rejection.<br>•Aplastic anemia (moderate to severe) in patients unsuitable for bone marrow transplantation.<br>•Limitations of Use: The usefulness of Atgam has not been demonstrated in patients with aplastic anemia<br>who are suitable candidates for bone marrow transplantation or in patients with aplastic anemia<br>secondary to neoplastic disease. A torage disease, myelofibrosis, Fancon's syndrome, or in patients known<br>to have been exposed to myelotoxic agents or radiation. | 235.2                             | N/A            | N/A         | N/A                    | Y               | Y                               |          | 9/12/2018             |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
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Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

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| Category      | HCPCS<br>Code  | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name       | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age   | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs         | J7613          | Albuterol, inhalation solution,<br>FDA-approved final product,<br>non-compounded,<br>administered through DME,<br>unit dose, 1 mg                              | 1 mg                       | 4/1/2008                | N/A              | albuterol sulfate inhalation<br>solution (0.021%, 0.042% an<br>0.083%)                                   | 0.63 mg/3 mL solution (0.021%) and 1.25 mg/3 mL solution (0.042%) formulations: Indicated for the relief<br>of bronchospasm in patients 2 to 12 years of age with asthma (reversible obstructive airway disease).<br>2.5 mg/3 mL solution (0.083%) formulation: Indicated for the relief of bronchospasm in patients 2 years of<br>age and older with reversible obstructive airway disease and acute attacks of bronchospasm.  | 310                               | 2 years   | Formulation<br>Specific Age<br>Restrictions<br>(see comments) | N/A                    | Y               | Ŷ                               | Formulation Specific:<br>0.63 mg/3 mL solution<br>(0.021%) and 1.25 mg/3 mL<br>solution (0.042%)<br>formulations: 2 to 12 years of<br>age<br>2.5 mg/3 mL solution (0.083%)<br>formulation: 2 years of age<br>and older | 9/21/2022             |
| Drugs         | J7614          | Levalbuterol, inhalation<br>solution, FDA-approved final<br>product, non-compounded,<br>administered through DME,<br>unit dose, 0.5 mg                         | 0.5 mg                     | 4/1/2008                | Xopenex®         | levalbuterol hydrochloride<br>inhalation solution  | Indicated for the treatment or prevention of bronchospasm in adults, adolescents, and children 6 years of<br>age and older with reversible obstructive airway disease.  | 310                               | 6 years   | N/A   | N/A                    | Ŷ               | Ŷ                               |  | 9/23/2022             |
| Drugs         | J7620          | Albuterol, up to 2.5 mg and<br>ipratropium bromide, up to<br>0.5 mg, FDA-approved final<br>product, non-compounded,<br>administered through DME                | 2.5 mg/0.5 mg              | 1/1/2006                | N/A              | ipratropium<br>bromide/albuterol sulfate<br>inhalation solution  | FDA Approved Indication: Indicated for the treatment of bronchospasm associated with COPD in patients<br>requiring more than one bronchodilator.<br>Recommended Uses from the National Heart, Lung, and Blood Institute: Asthma exacerbations for<br>children through 12 years of age and adults.   | 186                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A   | N/A                    | Ŷ               | Y                               | Indication Specific Age<br>Restrictions:<br>Treatment of bronchospasm<br>associated with COPD: 18<br>years of age and older<br>Asthma exacerbations: N/A   | 9/21/2022             |
| Drugs         | J7644          | Ipratropium bromide,<br>inhalation solution, FDA-<br>approved final product, non-<br>compounded, administered<br>through DME, unit dose form,<br>per milligram | 1 mg                       | 1/1/2000                | N/A              | ipratropium bromide<br>inhalation solution, 0.02%  | FDA Approved Indication: Indicated as a bronchodilator for maintenance treatment of bronchospasm<br>associated with chronic obstructive pulmonary disease, including chronic bronchitis and emphysema.<br>Recommended Uses from the National Heart, Lung, and Blood Institute: Asthma exacerbations for<br>children through 12 years of age and adults.   | 93                                | Indication Specific<br>Age Restrictions<br>(see comments) | N/A   | N/A                    | Y               | Y                               | Indication Specific Age<br>Restrictions:<br>Maintenance treatment of<br>bronchospasm associated with<br>chronic obstructive pulmonary<br>disease: 18 years of age and<br>older<br>Asthma exacerbations: N/A            | 9/23/2022             |
| Drugs         | J8499          | Prescription drug, oral, non-<br>chemotherapeutic, Not<br>Otherwise Specified  | 2 grams                    | 1/1/2000                | Flagyl®, Likmez" | metronidazole, oral  | Approved indications for use in the PADP:<br>• Symptomatic Trichomoniasis: Metronidatole is indicated for the treatment of 7. voginalis infection in<br>females and males when the presence of the trichomonad has been confirmed by appropriate laboratory<br>procedures (vet smears and/or cultures).<br>• Asymptomatic Trichomoniasis: Metronidatole is indicated in the treatment of asymptomatic T.<br>voginalis infection in females when the organism is associated with endocervicits, cervical<br>voginalis infection in females when the organism is associated with endocervicits, exvicits, or cervical<br>voginalis infection in females when the organism is associated with endocervicits, exvicits, or cervical<br>voginalis infection in females when the organism is associated with endocervicits, exvicits, or cervical<br>voginalis infection in termales what presence of the trichomonad can interfere with accurate assessment<br>of ahnormal cyclogicals mears, additional smears should be erfored after eradication of the parasite.<br>• Treatment of Asymptomatic Sexual Partners: T. voginalis infection is a venereal disease. Therefore,<br>asymptomatic sexual partners of treated patients should be treated atter endication of the organism<br>been found to be present, in order to prevent reinfection of the partner. The decision as to whether to<br>treat an asymptomatic mean reinfect the sexual partners is not treated. Also, since there can be considerable<br>difficulty in isolating the organism from the asymptomatic male carrier, negative smears and cultures<br>cannot be relied upon in this regard. In any event, the sexual partner should be treated with<br>Metronidazole in cases of reinfection. | 2                                 | N/A   | N/A   | N/A                    | Y               | Y                               |  | 12/1/2023             |
| Drugs         | 19000          | Injection, doxorubicin<br>hydrochloride, 10 mg   | 10 mg                      | 1/1/2000                | Adriamycin®      |  | Indicated:<br>• As a component of multiagent adjuvant chemotherapy for treatment of women with axillary lymph node<br>involvement following resection of primary breast cancer.<br>• For the treatment of a cute lymphoblastic leukemia, acute myeloblastic leukemia, Hodgkin lymphoma,<br>Non-Hodgkin lymphoma, metastatic leukemia, acute myeloblastic leukemia, Hodgkin lymphoma,<br>Non-Hodgkin lymphoma, metastatic breast cancer, metastatic Villms' tumor, metastatic noralina carcinoma, metastatic<br>transitional cell bladder carcinoma, metastatic thyroid carcinoma, metastatic gastric carcinoma,<br>metastatic sontogenic acrinoma.  | 38                                | N/A   | N/A   | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Drugs         | J9015          | Injection, aldesleukin, per<br>single-use via  | per single use vial        | 1/1/2000                | Proleukin®       | aldesleukin for injection, for<br>intravenous infusion   | Indicated for the treatment of adults with metastatic renal cell carcinoma and metastatic melanoma.   | 112                               | 18 years  | N/A   | N/A                    | Y               | Y                               |  | 6/6/2019              |
| Drugs         | J9017          | Injection, arsenic trioxide, 1<br>mg   | 1 mg                       | 1/1/2000                | Trisenox®        |  | <ul> <li>Indicated for induction of remission and consolidation in patients with acute promyelocytic leukemia<br/>(APL) who are refractory to, or have relapsed from, retinoid and anthracycline chemotherapy, and whose<br/>APL is characterized by the presence of the (15,127) translocation or PML/RAR-alpha expression.</li> <li>Indicated in combination with tretinoin for treatment of adults with newly-diagnosed low-risk acute<br/>promyelocytic leukemia (APL) whose APL is characterized by the presence of the t(15,127) translocation or<br/>PML/RAR-alpha gene expression.</li> </ul>   | 651                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A   | N/A                    | Ŷ               | Y                               | Indication specific age<br>restrictions:<br>• In combination with<br>tretinoin: 18 years of age and<br>older<br>• As a single agent: 5 years of<br>age and older   | 9/25/2018             |
| Drugs         | J9019          | Injection, asparaginase<br>(Erwinaze), 1,000 IU  | 1,000 units                | 1/1/2013                | Erwinaze®        | asparaginase erwinia<br>chrysanthemi for injection,<br>for intramuscular (IM) or<br>intravenous (IV) use | Indicated as a component of a multi-agent chemotherapeutic regimen for the treatment of patients with<br>acute lymphoblastic leukemia (ALL) who have developed hypersensitivity to E. coli-derived asparaginase.  | 420                               | 1 year  | N/A   | N/A                    | Y               | Y                               |  | 6/4/2019              |
| Biologicals   | J9021          | Injection, asparaginase,<br>recombinant, (rylaze), 0.1 mg  | 0.1 mg                     | 1/1/2022                | Rylaze™          | asparaginase erwinia<br>chrysanthemi (recombinant)<br>rywn injection, for<br>intramuscular use           | Indicated as a component of a multi-agent chemotherapeutic regimen for the treatment of acute<br>lymphoblastic leukemia (ALL) and lymphoblastic lymphoma (LBL) in adult and pediatric patients 1 month<br>or older who have developed hypersensitivity to E. coli-derived asparaginase.   | 12,200                            | 1 month   | N/A   | N/A                    | Ŷ               | Ŷ                               |  | 12/20/2022            |

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| Category    | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name   | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|-------------|---------------|---|----------------------------|-------------------------|--------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Biologicals | J9022         | Injection, atezolizumab, 10 mg  | 10 mg                      | 1/1/2018                | Tecentriq*   | atezolizumab injection, for<br>intravenous use                        | Indicated for the treatment of patients with:<br>• Non-Small Cell Lung Cancer (NSCLC)<br>O Metastatic non-small cell lung cancer who have disease progression during or following platinum-<br>containing chemotherapy. Patients with ESFR or ALK genomic tumor aberrations should have disease<br>progression on FDA approved therapy for these aberrations prior to receiving Tecentriq.<br>O in combination with beaucizumas placitaxel, and carobplatin, for the firstline treatment of patients with<br>metastatic non-squamous NSCLC with no ESFR or ALK genomic tumor aberrations.<br>O in combination with pacitaxel protein-bound and carobplatin for the first-line treatment of adult<br>patients with metastatic non-squamous NSCLC with no ESFR or ALK genomic tumor aberrations.<br>O for the firstline treatment of adult patients with metastatic NSCCL Workose tumors have high PD-L1<br>eSFR or ALK genomic tumor aberrations.<br>I Combination with acklosed protect [C 2: 50%] or PD-L1 stained tumor-infiltrating immune<br>cells [IC] covering 2: 10% of the tumor area [IC 2: 10%] ), as determined by an FDA-approved test, with no<br>ESFR or ALK genomic tumor aberrations.<br>• In combination with acklosed proc [C 5: 50%] or pD-L1 stained tumor-infiltrating immune<br>cells [IC] covering 2: 10% of the tumor area [IC 2: 10%] ), as determined by an FDA-approved test, with no<br>ESFR or ALK genomic tumor aberrations.<br>• I nombination with collowing rescue [C 5: 50%]<br>• In combination with collowing rescue [C 5: 50%]<br>• In combination with collowing resection and platinum-based chemotherapy for adult patients with<br>Stage II to IIIA NSCLC whose tumors have PD-L1 expression on 2 1% of tumor cells, as determined by an<br>FDA-approved test.<br>• Alveedar Soft Pat Sarcoma (ASPS)<br>O for the treatment of adult and pediatric patients 2 years of age and older with unresectable or<br>metastatic ASPS. | 336                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | NSCLC, SCLC, HCC, melanoma:<br>18 years of age and older<br>ASPS: 2 years of age and older  | 1/23/2023             |
| Biologicals | J9023         | Injection, avelumab, 10 mg  | 10 mg                      | 1/1/2018                | Bavencio*    | avelumab injection, for<br>intravenous use                            | Indicated for:<br>Adults and pediatric patients 12 years and older with metastatic Merkel cell carcinoma (MCC).<br>Patients with locally advanced or metastatic urothelial carcinoma (UC) who have disease progression<br>during or following platinum-containing chemotherapy or have disease progression within 12 months of<br>necadijuvant or advance with platinum-containing chemotherapy.<br>Maintenance treatment of patients with locally advanced or metastatic UC that has not progressed with<br>first-line platinum-containing chemotherapy.<br>First-line treatment, in combination with axitinib, of patients with advanced renal cell carcinoma (RCC).   | 240                               | 12 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 7/28/2020             |
| Drugs       | J9025         | Injection, azacitidine, 1 mg  | 1 mg                       | 1/1/2006                | Vidaza®      | azacitidine for injection, for<br>subcutaneous or intravenous<br>use  | Indicated for the treatment of:<br>- Adult patients with the following FAB myelodysplastic syndrome (MDS) subtypes: refractory anemia (RA)<br>or refractory anemia with ringed sideroblasts (RARS) (if accompanied by neutropenia or thrombocytopenia<br>or requiring transfusions), refractory anemia with excess blasts (RARE), refractory anemia with excess<br>blasts in transformation (RAEB-T) and chronic myelomonocytic leukemia (CMMoL).<br>- Pediatric patients aged 1 month and older with newly diagnosed Juvenile Myelomonocytic Leukemia<br>(MML).   | 3,000                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Ŷ               | Ŷ                               | Indication specific age<br>restrictions:<br>• Adult patients with FAB<br>myelodysplastic syndrome<br>(MDS) subtypes - 18 years of<br>age and older<br>• Pediatric patients with JMML<br>- 1 month and older | 6/9/2022              |
| Biologicals | J9029         | Intravesical instillation,<br>nadofaragene firadenovec-<br>vncg, per therapeutic dose | 1 therapeutic dose         | 7/1/2023                | Adstiladrin® | nadofaragene firadenovec-<br>vncg suspension, for<br>intravesical use | Indicated for the treatment of adult patients with high-risk Bacillus Calmette-Guérin (BCG)-unresponsive<br>non-muscle invasive bladder cancer (NMIBC) with carcinoma in situ (CIS) with or without papillary tumors   | . 1                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 3/22/2024             |
| Biologicals | 19030         | Bcg live intravesical instillation, 1 mg  | per installation           | 1/1/2000                | Tice BCG*    | BCG Live (intravesical)   | Indicated for the treatment and prophylaxis of carcinoma in situ (CIS) of the urinary bladder, and for the<br>prophylaxis of primary or recurrent stage Ta and/or T1 papillary tumors following transurethral resection<br>(TUR). Tice BCG is not recommended for stage TaG1 papillary tumors, unless they are judged to be at high<br>risk of tumor recurrence. Tice BCG is not indicated for papillary tumors of stages higher than T1.  | 250                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               | 6/2024: NC Suggested Max<br>Monthly Units updated to align<br>with NCTracks, which has been<br>set to 250 units/month since<br>7/1/2019.  | 6/7/2024              |
| Drugs       | J9032         | Injection, belinostat, 10 mg  | 10 mg                      | 1/1/2016                | Beleodaq®    | belinostat for injection, for<br>intravenous use                      | Indicated for the treatment of patients with relapsed or refractory peripheral T-cell lymphoma (PTCL).   | 2,500                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 4/10/2019             |
| Drugs       | 19033         | Injection, bendamustine HCl<br>(Treanda), 1 mg  | 1 mg                       | 1/1/2017                | Treanda*     | bendamustine hydrochloride<br>injection, for intravenous use          | not been established.<br>• Indolent B-cell non-Hodgkin lymphoma (NHL) that has progressed during or within six months of<br>treatment with rituximab or a rituximab-containing regimen.  | 1,200                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 9/25/2018             |
| Drugs       | J9034         | Injection, bendamustine HCl<br>(Bendeka), 1 mg  | 1 mg                       | 1/1/2017                | Bendeka®     | bendamustine hydrochloride<br>injection, for intravenous use          |  | 1,200                             | 18 years  | N/A         | N/A                    | Ŷ               | Ŷ                               |   | 9/25/2018             |

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|-------------|---------------|--|----------------------------|-------------------------|------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Biologicals | 19035         | Injection, bevacizumab, 10 mg  | 10 mg                      | 1/1/2005                | Avastin*   | bevacizumab injection, for<br>intravenous use                       | Indicated for the treatment of:<br>• Metastatic colorectal cancer, in combination with intravenous 5-fluorouracil-based chemotherapy for<br>first- or second-line treatment.<br>• Metastatic colorectal cancer, in combination with fluoropyrimidine-innotecan- or fluoropyrimidine-<br>bevacitumab product-containing regimen.<br>• Unrescatable, locally advanced, recurrent or metastatic non-squamous non-small cell lung cancer, in<br>combination with carboptatin and pacitized for first-line treatment.<br>• Recurrent glioblastoma in adults.<br>• Metastatic renal cell carcinoma in combination with interferon alfa.<br>• Persistent, recurrent, or metastatic cervical cancer, in combination with pacitized and cisplatin, or<br>pacificate and topotecan.<br>• Epithelial ovarian, fallopian tube, or primary peritoneal cancer:<br>• In combination with actoplatin and pacitized flosomal doxorubicin, or topotecan for platinum-resistant<br>recurrent disease who received on more than 2 pirot chemotherapy regimens.<br>• In combination with carboptatin and pacitized liposomal doxorubicin, or topotecan for platinum-resistant<br>recurrent disease who received no more than 2 pirot, chemotherapy regimens.<br>• In combination with carboptatin and pacitized is carboptatin and gencitabine, followed by Avastin as a<br>single agent, for platinum sensitive recurrent disease.<br>• In combination with carboptatin and pacitizeds, followed by Avastin as a single agent, for stage III or IV<br>disease following initial surgical resection.<br>• In combination with carboptatin and pacitizeds, followed by Avastin as a single agent, for stage III or IV<br>disease following initial surgical resection.<br>• In combination with actoble of the treatment of platents with unresectable or metastatic<br>hepatocellular carcinoma (HCC) who have not received prior systemic therapy.<br>Limitation of Use: Avastin is not indicated for adjuvant treatment of colon cancer. | 420                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/20/2022            |
| Drugs       | J9036         | Injection, bendamustine<br>hydrochloride,<br>(Belrapzo/bendamustine), 1<br>mg              | 1 mg                       | 7/1/2019                | Belrapzo™  | bendamustine hydrochloride<br>injection for intravenous use         | **Macular edema (non-FDA approved indication)<br>indicated for treatment of patients with:<br>• Chronic lymphocytic leukemia (CLL). Efficacy relative to first line therapies other than chlorambucil has<br>not been established.<br>• Indolent B-cell non-Hodgkin lymphoma (NHL) that has progressed during or within six months of<br>treatment with rituximab or a rituximab-containing regimen.  | 1,200                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 8/26/2019             |
| Biologicals | 19039         | Injection, blinatumomab, 1<br>mcg  | 1 mcg                      | 1/1/2016                | Blincyto®  | blinatumomab for injection,<br>for intravenous use                  | Indicated for the treatment of a duit and pediatric patients one month and older with:<br>• Relapsed or refractory CD19-positive B-cell precursor acute lymphoblastic leukemia (ALL).<br>• CD19-positive B-cell precursor acute lymphoblastic leukemia (ALL) in first or second complete remission<br>with minimal residual disease (MRD) greater than or equal to 0.1%.<br>• CD19-positive Philadelphia chromosome-negative B-cell precursor acute lymphoblastic leukemia (ALL) in<br>the consolidation phase of multiphase chemotherapy.  | 980                               | 1 month   | N/A         | N/A                    | Y               | Ŷ                               |  | 7/29/2024             |
| Drugs       | J9040         | Injection, bleomycin sulfate,<br>15 units  | 15 units                   | 1/1/2000                | N/A        | bleomycin for injection   | Considered a palliative treatment shown to be useful in the management of:<br>• Squamous Cell Carcinoma: Head and neck (including mouth, tongue, tonsil, nasopharynx, oropharynx,<br>sinus, palate, lip, buccal mucosa; gingvae, epiglotits, skin, laynx), pensis, cervix, and vulva. The response<br>to bicomycin is poorer in patients with previously irradiated head and neck cancer.<br>• Jumphomas: Hodgkin's disease. non-Hodgkin's disease<br>• Testicular Carcinoma: Ethoryconal cell, choricarcinoma, and teratocarcinoma<br>• Malignant Pleural Eflusion: Bleomycin is effective as a sclerosing agent for the treatment of malignant<br>pleural effusion and prevention of recurrent pleural effusions.   | 27                                | N/A   | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Drugs       | J9041         | Injection, bortezomib, 0.1 mg  | 0.1 mg                     | 1/1/2005                | Velcade®   | bortezomib for injection, for<br>subcutaneous or intravenous<br>use |   | 245                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/12/2022            |
| Biologicals | J9042         | Injection, brentuximab<br>vedotin, 1 mg  | 1 mg                       | 1/1/2013                | Adcetris®  | brentuximab vedotin for<br>injection, for intravenous use           | Indicated for:<br>• Previously untreated Stage III or IV classical Hodgkin lymphoma (cHL), in combination with doxorubicin,<br>vibilastine, and dacarbazine.<br>• Classical Hodgkin lymphoma (cHL) at high risk of relapse or progression as post-autologous<br>hematopoietic stem cell transplantation lauto-HSCT consolitation.<br>• Classical Hodgkin lymphoma (cHL) after failure of auto-HSCT cardidates.<br>• Classical Hodgkin lymphoma (cHL), after failure of auto-HSCT cardidates.<br>• Previously untreated systemic anaplastic large cell lymphoma (ALCL) or other CD30-expressing<br>peripheral T-cell lymphomas (PTCL), including angioimmunoblastic T-cell lymphoma and PTCL not<br>otherwise specified, in combination with cyclophosphamide, doxorubicin, and predisone.<br>• Systemic anaplastic large cell lymphoma (pcALCL) after failure of at least one prior multi-agent<br>chemotherapr regimen.<br>• Primary vutaneous anaplastic large cell lymphoma (pcALCL) or CD30- expressing mycosis fungoides (MF)<br>who have received prior systemic therapy.<br>Indicated for:<br>• Pediatric patients 2 years and older with previously untreated high risk classical Hodgkin lymphoma<br>(cHL), in combination with doxorubicin, vincristine, etoposide, prednisone, and cyclophosphamide.   | 360                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | ¥               | ¥                               | Indication specific age<br>restrictions:<br>• Previously untreated high<br>risk classical Hodgkin<br>lymphoma (cHL): 2 years and<br>older<br>• Other indications: 18 years<br>of age and older | 12/20/2022            |
| Drugs       | J9043         | Injection, cabazitaxel, 1 mg   | 1 mg                       | 1/1/2012                | Jevtana®   | cabazitaxel injection, for<br>intravenous use                       | Indicated in combination with prednisone for treatment of patients with hormone-refractory metastatic<br>prostate cancer previously treated with a docetaxel-containing treatment regimen.  | 240                               | 18 years  | N/A         | Males Only             | Y               | Y                               |  | 9/27/2018             |
| Drugs       | J9045         | Injection, carboplatin, 50 mg  | 50 mg                      | 1/1/2000                | N/A        | carboplatin injection for<br>intravenous use                        | Indicated for the initial treatment of advanced ovarian carcinoma in established combination with other<br>approved chemotherapeutic agents and for the pallative treatment of patients with ovarian carcinoma<br>recurrent after prior chemotherapy, including patients who have previously been treated with cisplatin.   | 36                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019             |
| Drugs       | 19046         | Injection, bortezomib (dr.<br>reddy's), not therapeutically<br>equivalent to j9041, 0.1 mg | 0.1 mg                     | 1/1/2023                | N/A        | bortezomib for injection, for<br>intravenous use (Dr. Reddy's)      | Indicated for:<br>• treatment of adult patients with multiple myeloma<br>• treatment of adult patients with mantle cell lymphoma who have received at least 1 prior therapy   | 245                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 12/12/2022            |

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|-------------|---------------|--|----------------------------|-------------------------|------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|----------------------|
| Drugs       | J9047         | Injection, carfilzomib, 1 mg   | 1 mg                       | 1/1/2014                | Kyprolis®  | carfilzomib for injection, for<br>intravenous use                             | indicated for the treatment of adult patients with relapsed or refractory multiple myeloma who have<br>received one to three lines of therapy in combination with:<br>o Lenalidomide and dexamethasone; or<br>o Dexamethasone; or<br>o Daratumumab and dexamethasone; or<br>o Daratumumab and myaluronidase-fihj and dexamethasone; or<br>o Statumiab and decamethasone<br>indicated as a single agent for the treatment of patients with relapsed or refractory multiple myeloma   | 1060                              | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 7/20/2022            |
| Drugs       | J9048         | Injection, bortezomib<br>(fresenius kabi), not<br>therapeutically equivalent to<br>j9041, 0.1 mg | 0.1 mg                     | 1/1/2023                | N/A        | bortezomib for injection, for<br>intravenous use (Fresenius<br>Kabi)          | who have received one or more lines of therapy.<br>Indicated for:<br>• treatment of adult patients with multiple myeloma<br>• treatment of adult patients with mantle cell lymphoma who have received at least 1 prior therapy  | 245                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 12/12/202            |
| Drugs       | J9049         | Injection, bortezomib<br>(hospira), not therapeutically<br>equivalent to j9041, 0.1 mg           | 0.1 mg                     | 1/1/2023                | N/A        | bortezomib for injection, for<br>subcutaneous or intravenous<br>use (Hospira) |   | 245                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 12/19/202            |
| Drugs       | J9050         | Injection, carmustine, 100 mg  | 100 mg                     | 1/1/2000                | BiCNU®     | carmustine for injection  | Indicated as pailiative therapy as a single agent or in established combination therapy with other approved<br>chemotherapeutic agents in the following:<br>• Brain tumors - gliobiastoma, brainstem glioma, medulloblastoma, astrocytoma, ependymoma, and<br>metastatic brain tumors.<br>• Multiple myeloma - in combination with prednisone.<br>• Hotogkin's disease: as secondary therapy in combination with other approved drugs in patients who<br>relapse while being treated with primary therapy, or who fail to respond to primary therapy.<br>• Non-Hotogkin's disease: coscondary therapy in combination with other approved drugs for<br>patients who relapse while being treated with primary therapy, or who fail to respond to primary therapy.   | 5                                 | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |          | 5/20/2019            |
| Drugs       | J9051         | Injection, bortezomib (maia),<br>not therapeutically equivalent<br>to j9041, 0.1 mg              | 0.1 mg                     | 10/1/2023               | N/A        | bortezomib injection, for<br>intravenous use (Maia)                           | Indicated for:<br>• treatment of adult patients with multiple myeloma<br>• treatment of adult patients with mantle cell lymphoma  | 245                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/28/2023            |
| Drugs       | J9052         | Injection, carmustine (accord),<br>not therapeutically equivalent<br>to j9050, 100 mg            |                            | 1/1/2024                | N/A        | carmustine for injection, for intravenous use (Accord)                        | Carmustine for injection is indicated as palliative therapy as a single agent or in established combination<br>therapy with other approved chemotherapeutic agents in the following:<br>Brain tumors glioblastoma, brainstem glioma, medulloblastoma, astrocytoma, ependymoma, and<br>metastatic brain tumors<br>• Multiple myeloma-in combination with prednisone<br>• Relapsed or refractory Hogkin's lymphoma in combination with other approved drugs<br>• Relapsed or refractory non-Hogkin's lymphomas in combination with other approved drugs   | 5                                 | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 12/22/202            |
| Biologicals | J9055         | Injection, cetuximab, 10 mg  | 10 mg                      | 1/1/2005                | Erbitux*   | cetuximab injection, for<br>intravenous use                                   | Indicated for:<br>• Squamous Cell Carcinoma of the Head and Neck (SCCHN):<br>• Locally or regionally advanced squamous cell carcinoma of the head and neck in combination with<br>radiation therapy.<br>• Recurrent locoregional disease or metastatic squamous cell carcinoma of the head and neck in<br>combination with platinum-based therapy with fluorouracil.<br>• Recurrent or metastatic squamous cell carcinoma of the head and neck progressing after platinum-based<br>therapy.<br>• K-Ras Wild-type, EGFR-expressing Colorectal Cancer (CRC) as determined by an FDA-approved test:<br>- In combination with Folfin for first-line treatment,<br>- In combination with inforteran in patients who are refractory to irinotecan-based chemotherapy,<br>- As a single agent in patients who have failed oxaliplatin- and irinotecan-based chemotherapy or who are<br>intolerant to irinotecan.<br>Limitations of Use: Erbitus is not indicated for treatment of Ras-mutant colorectal cancer or when the<br>results of the Ras mutation tests are unknown.<br>• BRAF V600E Mutation-Positive Metastatic Colorectal Cancer (CRC)<br>- in combination with encorafenib, for the treatment of adult patients with metastatic colorectal cancer<br>(CRC) with a BRAF V600E mutation, as detected by an FDA-approved test, after prior therapy. | 390                               | 18 years    | N/A         | N/A                    | Y               | ¥                               |          | 10/26/2021           |
| Drugs       | J9056         | Injection, bendamustine<br>hydrochloride (vivimusta), 1<br>mg                                    | 1 mg                       | 7/1/2023                | Vivimusta  | bendamustine hydrochloride<br>injection, for intravenous use                  | Indicated for treatment of patients with:<br>• Chronic kymphocycic leukemia (CLL). Efficacy relative to first line therapies other than chlorambucil has<br>not been established.<br>• Indicent 8-cell non-Hodgkin kymphoma (NHL) that has progressed during or within six months of<br>treatment with miximab or arining regimen.  | 1,200                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/22/2023            |
| Drugs       | J9057         | Injection, copanlisib, 1 mg  | 1 mg                       | 1/1/2019                | Aliqopa™   | copanlisib injection, for<br>intravenous use                                  | Indicated for the treatment of adult patients with relapsed follicular lymphoma (FL) who have received at<br>least two prior systemic therapies. Accelerated approval was granted for this indication based on overall<br>response rate. Continued approval for this indication may be contingent upon verification and description<br>of clinical benefit in a confirmatory trial.   | 240                               | 18 years    | N/A         | N/A                    | Y               | Ŷ                               |          | 8/5/2021             |
| Drugs       | J9058         | Injection, bendamustine<br>hydrochloride (apotex), 1 mg  | 1 mg                       | 7/1/2023                | N/A        | bendamustine hydrochloride<br>injection, for intravenous use<br>(Apotex)      |   | 1,440                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 5/23/2024            |

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| Category    | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modifie<br>Date |
|-------------|---------------|---|----------------------------|-------------------------|------------|--|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|----------------------|
| Drugs       | 19059         | Injection, bendamustine<br>hydrochloride (baxter), 1 mg       | 1 mg                       | 7/1/2023                | N/A        | bendamustine hydrochloride<br>injection, for intravenous use<br>(Baxter) |  | 1,200                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/22/2023            |
| Drugs       | 19060         | Injection, cisplatin, powder or<br>solution, per 10 mg        | 10 mg                      | 1/1/2000                | N/A        | cisplatin injection  | Indicated as therapy for:<br>• Metastatic Testicular Tumors: In established combination therapy with other approved<br>chemotherapeutic agents in patients with metastatic testicular tumors who have already received<br>appropriate surgical and/or andiotherapeutic proceedures.<br>• Metastatic Ovarian Tumors: In established combination therapy with other approved chemotherapeutic<br>agents in patients with metastatic ovarian tumors who have already received appropriate surgical and/or<br>radiotherapeutic procedures. An established combination consists of cisplatin and vc/polopsphamide.<br>Cisplatin Injection, as a single agent, is indicated as secondary therapy in patients with metastatic ovarian<br>tumors refractory to standard chemotherapy who have on previously received Cisplatin injection injection herapy.<br>• Advanced Bladder Cancer: Indicated as a single agent for patients with transitional cell bladder cancer<br>which is no longer amenable to local treatments, such as surgery and/or radiotherapy. | 50                                | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/27/2018            |
| Biologicals | J9061         | Injection, amivantamab-vmjw,<br>2 mg                          | 2 mg                       | 1/1/2022                | Rybrevant™ | amivantamab-vmjw injection,<br>for intravenous use                       | Indicated:<br>- as a single agent for the treatment of adult patients with locally advanced or metastatic non-small cell<br>lung cancer (NSCLC) with epidermal growth factor receptor (EGFR) exon 20 insertion mutations, as<br>detected by an FDA-approved test, whose disease has progressed on or after platinum-based<br>chemotherapy.<br>- in combination with carboplatin and pemetrexeef for the first-line treatment of adult patients with<br>locally advanced or metastatic non-small cell lung cancer (NSCLC) with epidermal growth factor receptor<br>(EGFR) exon 20 insertion mutations, as detected by an FDA-approved test.   | 2,800                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 5/3/2024             |
| Biologicals | 19063         | Injection, mirvetuximab soravtansine-gynx, 1 mg               | 1 mg                       | 7/1/2023                | Elahere™   | mirvetuximab soravtansine-<br>gynx injection, for<br>intravenous use     | indicated for the treatment of adult patients with FRα positive, platinum-resistant epithelial ovarian,<br>fallopian tube, or primary peritoneal cancer, who have received one to three prior systemic treatment<br>regimens. Select patients for therapy based on an FDA-approved test.   | 1,800                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/22/2023            |
| Drugs       | J9065         | Injection, cladribine, per 1 mg                               | 1 mg                       | 1/1/2000                | N/A        | cladribine injection   | Indicated for the treatment of active Hairy Cell Leukemia as defined by clinically significant anemia,<br>neutropenia, thrombocytopenia, or disease-related symptoms.  | 91                                | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/4/2019             |
| Drugs       | J9071         | Injection, cyclophosphamide<br>(auromedics), 5 mg             | 5 mg                       | 4/1/2022                | N/A        | cyclophosphamide for<br>injection, for intravenous use<br>(AuroMedics)   | Indicated for the treatment of:<br>Malignant Diseases: malignant lymphomas, Hodgkin's disease, lymphocytic lymphoma, mixed-cell type<br>lymphoma, histoiyotic lymphoma, Burkit's lymphoma; multiple myeloma, leukemias, mycosis fungoides,<br>neuroblastoma, adenocarcinoma of ovary, retinoblastoma, breast carcinoma.  | 2,500                             | N/A         | N/A         | N/A                    | Y               | Y                               |          | 3/17/2022            |
| Drugs       | J9072         | Injection, cyclophosphamide<br>(dr. reddy's), 5 mg            | 5 mg                       | 1/1/2024                | N/A        | cyclophosphamide injection,<br>for intravenous use (Dr.<br>Reddy's)      | Cyclophosphamide injection is indicated for treatment of adult and pediatric patients with:<br>• Malignant Diseases:<br>- malignant lymphomas: Hodgkin's disease, lymphocytic lymphoma, mixed-cell type lymphoma, histiocytic<br>lymphoma, Burkit's lymphoma;<br>- multiple myeloma, leukemias, mycosis fungoides, neuroblastoma, adenocarcinoma of ovary,<br>retinoblastoma, breast carcinoma.<br>Limitations of Use:<br>The safety and effectiveness for the treatment of nephrotic syndrome in adults or other renal disease has<br>not been established.   | 2,500                             | N/A         | N/A         | N/A                    | Y               | ¥                               |          | 12/22/2023           |
| Drugs       | J9073         | Injection, cyclophosphamide<br>(ingenus), 5 mg                | 5 mg                       | 4/1/2024                | N/A        |  | Cyclophosphamide is indicated for treatment of:<br>• Malignant Diseases: malignant lymphomas: Hodgkin's disease, lymphocytic lymphoma, mixed-cell type<br>lymphoma, histiocytic lymphoma, Burkit's lymphoma; multiple myeloma, leukemias, mycosis fungoides,<br>neuroblastoma, adenocarcinoma of ovary, retinoblastoma, breast carcinoma.  | 2,100                             | N/A         | N/A         | N/A                    | Y               | Ŷ                               |          | 3/27/2024            |
| Drugs       | J9074         | Injection, cyclophosphamide<br>(sandoz), 5 mg                 | 5 mg                       | 4/1/2024                | N/A        | cyclophosphamide injection,<br>for intravenous use (Sandoz)              | Cyclophosphamide injection is an alkylating drug indicated for treatment of adult patients with:<br>Malignant Diseases: malignant lymphomas: Hodgkin's lymphoma, lymphocytic lymphoma, mixed-cell type<br>lymphoma, histiocytic lymphoma, Burkit's lymphoma; multiple myedoma, leukemias, mycosis fungoides,<br>neuroblastoma, adenocarcinoma of ovary, retinoblastoma, breast carcinoma.<br>Limitations of Use:<br>This cyclophosphamide product is not indicated for use in pediatric patients due to the alcohol and<br>propylene glycol content in this product. If treatment with cyclophosphamide is indicated in a pediatric<br>patient, use a different cyclophosphamide product.  | 2,100                             | 18 years    | N/A         | N/A                    | Y               | Ŷ                               |          | 5/3/2024             |
| Drugs       | J9075         | Injection, cyclophosphamide,<br>not otherwise specified, 5 mg | 5 mg                       | 4/1/2024                | N/A        |  | Indicated for the treatment of:<br>Malignant Diseases: malignant lymphomas: Hodgkin's disease, lymphocytic lymphoma, mixed-cell type   | 2,500                             | N/A         | N/A         | N/A                    | Y               | Y                               |          | 3/22/2024            |
| Drugs       | J9100         | Injection, cytarabine, 100 mg                                 | 100 mg                     | 1/1/2000                | N/A        | cytarabine injection   | In combination with other approved anticancer drugs, is indicated for remission induction in acute non-<br>lymphocytic leukemia of adults and pediatric patients. It has also been found useful in the treatment of<br>acute lymphocytic leukemia and the blast phase of chronic myelocytic leukemia. Intrathecal<br>administration of cytarabine injection (preservative-free preparations only) is indicated in the prophylaxis<br>and treatment of meningeal leukemia.  | 35                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 7/2/2018             |
| Biologicals | J9118         | Injection, calaspargase pegol-<br>mknl, 10 units              | 10 units                   | 10/1/2019               | Asparlas™  | calaspargase pegol-mknl<br>injection, for intravenous use                | Indicated for the treatment of acute lymphoblastic leukemia in pediatric and young adult patients age 1 month to 21 years.   | 1,500                             | 1 month     | 21 years    | N/A                    | Y               | Y                               |          | 12/3/2019            |

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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name              | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|-------------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Biologicals | J9119         | Injection, cemiplimab-rwlc, 1<br>mg  | 1 mg                       | 10/1/2019               | Libtayo®                | cemiplimab-rwlc injection,<br>for intravenous use                         | Indicated<br>• for the treatment of patients with metastatic cutaneous squamous cell carcinoma (CSCC) or locally<br>advanced CSCC who are not candidates for curative surgery or curative radiation.<br>• for the treatment of patients with locally advanced BCC (laBCC) previously treated with a hedgehog  | 700                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 12/20/2022            |
| Drugs       | J9120         | Injection, dactinomycin, 0.5<br>mg   | 0.5 mg                     | 1/1/2000                | Cosmegen®               | dactinomycin for injection,<br>for intravenous use                        | To the readment of parents with recarry advances but (parect) previously tracked with a neugening     indicated for the treatment of:     adult and pediatric patients with Wilms tumor, as part of a multi-phase, combination chemotherapy     regimen     adult and pediatric patients with rhabdomyosarcoma, as part of a multi-phase, combination     chemotherapy regimen  | 42                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 9/25/2018             |
| Drugs       | J9130         | Dacarbazine, 100 mg  | 100 mg                     | 1/1/2000                | N/A                     | dacarbazine for injection   | Indicated for the treatment of metastatic malignant melanoma and as secondary-line therapy when used<br>in combination with other effective agents for Hodkin's disease.  | 91                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 6/10/2019             |
| Biologicals | J9144         | Injection, daratumumab, 10<br>mg and hyaluronidase-fihj                            | 10 mg                      | 1/1/2021                | Darzalex Faspro™        | daratumumab and<br>hyaluronidase-fihj injection,<br>for subcutaneous use  | Indicated for the treatment of adult patients with:<br>• multiple myeloma in combination with bortezomik, melphalan and prednisone in newly diagnosed<br>patients who are ineligible for autologous stem cell transplant<br>• multiple myeloma in combination with lenalidomide and dexamethasone in newly diagnosed patients<br>who are ineligible for autologous stem cell transplant and in patients with relapsed or refractory multiple<br>myeloma who have received at least one prior therapy<br>• multiple myeloma in combination with bortezomib and dexamethasone in patients who have received at<br>least one prior therapy   | 900                               | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |          | 12/16/2021            |
| Biologicals | J9145         | Injection, daratumumab, 10<br>mg   | 10 mg                      | 1/1/2017                | Darzalex*               | daratumumab injection, for<br>intravenous use                             | Indicated for the treatment of adult patients with multiple myeloma:<br>in combination with lenaildomide and deaxmethasione in patients with relapsed or refractory multiple<br>myeloma who have received at least one prior therapy.<br>• in combination with bortezomib and deaxmethasone in patients who have received at least one prior<br>therapy.  | 1,120                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/21/2020             |
| Drugs       | J9150         | Injection, daunorubicin, 10 mg   | 10 mg                      | 1/1/2000                | N/A                     | daunorubicin hydrochloride<br>injection                                   | In combination with other approved anticancer drugs, daunorubicin is indicated for remission induction in<br>acute nonlymphocytic leukemia (myelogenous, monocytic, erythroid) of adults and for remission inductior<br>in acute lymphocytic leukemia of children and adults.   | n 60                              | N/A         | N/A         | N/A                    | Ŷ               | Y                               |          | 6/10/2019             |
| Drugs       | J9153         | Injection, liposomal, 1 mg<br>daunorubicin and 2.27 mg<br>cytarabine               | 1 mg/2.27 mg               | 1/1/2019                | Vyxeos™                 | daunorubicin and cytarabine<br>liposome injection, for<br>intravenous use | Indicated for:<br>- the treatment of adults with newly-diagnosed therapy-related acute myeloid leukemia (t-AML) or AML<br>with myelodysplasia-related changes (AML-MRC).<br>- the treatment of newly-diagnosed therapy-related acute myeloid leukemia (t-AML) or AML with<br>myelodysplasia-related changes (AML-MRC) in pediatric patients 1 year and older.   | 660                               | 1 year      | N/A         | N/A                    | Y               | Y                               |          | 4/26/2021             |
| Drugs       | J9155         | Injection, degarelix, 1 mg   | 1 mg                       | 1/1/2010                | Firmagon®               | degarelix for injection for<br>subcutaneous administration                | Indicated for the treatment of patients with advanced prostate cancer.  | 320                               | 18 years    | N/A         | Males Only             | Y               | Y                               |          | 10/4/2018             |
| Drugs       | J9171         | Injection, docetaxel, 1 mg   | 1 mg                       | 1/1/2010                | Docefrez®,<br>Taxotere® | docetaxel injection<br>concentrate, intravenous<br>infusion               | Indicated for:<br>Breast Cancer (BC): single agent for locally advanced or metastatic BC after chemotherapy failure; and<br>with downobion and cyclophosphamide as adjuvant treatment of operable node-positive BC.<br>* Non-Small Cell Lung Cancer (NSCLC): single agent for locally advanced or metastatic NSCLC after<br>platinum therapy failure; and with isoplain for unreacetable, locally advanced or metastatic untreated<br>NSCLC.<br>+ Normone Refractory Prostate Cancer (HRPC): with prednisone in androgen independent (hormone   | 500                               | N/A         | N/A         | N/A                    | Ŷ               | Y                               |          | 6/8/2019              |
| Drugs       | J9172         | Injection, docetaxel (ingenus)<br>not therapeutically equivalent<br>to j9171, 1 mg | 1 mg                       | 1/1/2024                | Docivyx                 | docetaxel injection, for intravenous use (Ingenus)                        | * Non-more relativity in rotatic cancer. Professional relativity in rotatic cancer. Docetasel injection is indicated for: Breast Cancer (RC): single agent for locally advanced or metastatic BC after chemotherapy failure; and with doxorubicin and cyclophosphamide as adjuvant treatment of operable node-positive BC * Non-small Cell Lung Cancer (NSCLC): single agent for locally advanced or metastatic RSCL after platinum therapy failure; and with isoplatinum therapy failure; advanced to recetable, locally advanced or metastatic NSCL after platinum therapy failure; and with optimizing therapy failure; and with advanced or metastatic wanced or metas | 520                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 5/23/2024             |
| Biologicals | J9173         | Injection, durvalumab, 10 mg   | 10 mg                      | 1/1/2019                | Imfinzi®                | durvalumab injection, for<br>intravenous use                              | Indicated for the treatment of patients with:<br>• unresectable, Stage III non-small cell lung cance (NSCLC) whose disease has not progressed following<br>concurrent platimum-based chemotherapy and radiation therapy<br>• in combination with etoposide and either carboplatin or cisplatin, as first-line treatment of adult<br>patients with extensive-stage small cell lung cancer (ES-SCLC).   | 420                               | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |          | 7/29/2024             |
| Biologicals | J9176         | Injection, elotuzumab, 1 mg  | 1 mg                       | 1/1/2017                | Empliciti®              | elotuzumab for injection, for<br>intravenous use                          | Indicated in:<br>• combination with lenalidomide and dexamethasone for the treatment of adult patients with multiple<br>myeloma who have received one to three prior therapies.   | 5,600                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 5/20/2019             |
| Biologicals | J9177         | Injection, enfortumab vedotin-<br>ejfv, 0.25 mg                                    | 0.25 mg                    | 7/1/2020                | Padcev®                 | enfortumab vedotin-ejfv for<br>injection, for intravenous use             |   | 2,080                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 2/16/2024             |
| Drugs       | J9178         | Injection, epirubicin HCl, 2 mg  | 2 mg                       | 1/1/2004                | Ellence®                | epirubicin hydrochloride<br>injection                                     | Indicated as a component of adjuvant therapy in patients with evidence of axillary node tumor<br>involvement following resection of primary breast cancer.  | 300                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 10/10/2018            |
| Drugs       | J9179         | Injection, eribulin mesylate,<br>0.1 mg  | 0.1 mg                     | 1/1/2012                | Halaven®                | eribulin mesylate injection,<br>for intravenous use                       | Indicated for the treatment of patients with:<br>• Metastatic breast cancer who have previously received at least two chemotherapeutic regimens for the<br>treatment of metastatic disease. Prior therapy should have included an anthracycline and a taxane in   | 160                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/4/2019              |

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|-------------|---------------|---|----------------------------|-------------------------|-------------------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|----------------------|
| Drugs       | J9181         | Injection, etoposide, 10 mg   | 10 mg                      | 1/1/2000                | Etopophos®,<br>Toposar™ | etoposide phosphate for<br>injection, for intravenous use           | Indicated for the treatment of patients with:<br>• Refractory testicular tumors, in combination with other chemotherapeutic drugs.<br>• Small cell lung cancer, in combination with cisplatin, as first-line treatment.   | 300                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 6/10/2019            |
| Drugs       | J9185         | Injection, fludarabine<br>phosphate, 50 mg  | 50 mg                      | 1/1/2000                | N/A                     | fludarabine phosphate for<br>injection for intravenous use          | Indicated for the treatment of adult patients with B-cell chronic lymphocytic leukemia (CLL) who have not<br>responded to or whose disease has progressed during treatment with at least 1 standard alkylating-agent<br>containing regimen. The safety and effectiveness of fludarabine in previously untreated or non-refractory.  |                                   | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/10/2018           |
| Drugs       | J9190         | Injection, fluorouracil, 500 mg   | 500 mg                     | 1/1/2000                | Adrucil®                | fluorouracil injection for<br>intravenous use                       | Indicated for the treatment of patients with:<br>• Adenocarcinoma of the colon and rectum<br>• Adenocarcinoma of the breast<br>• Gastric adenocarcinoma<br>• Pancreatic adenocarcinoma  | 45                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019            |
| Drugs       | J9196         | Injection, gemcitabine<br>hydrochloride (accord), not<br>therapeutically equivalent to<br>J9201, 200 mg | 200 mg                     | 4/1/2023                | N/A                     | gemcitabine injection, for<br>intravenous use (Accord)              | Indicated:<br>• In combination with carboplatin, for the treatment of advanced ovarian cancer that has relapsed at<br>least 6 months after completion of platinum-based therapy.<br>• In combination with paclitaxel, for first-line treatment of metastatic breast cancer after failure of prior<br>anthracycline-containing adjuvant chemotherapy, unless anthracyclines were clinically contraindicated.<br>• In combination with cisplatin for the treatment of non-small cell lung cancer.                               | 64                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 3/16/2023            |
| Drugs       | J9198         | Injection, gemcitabine<br>hydrochloride, (infugem), 100<br>mg   | 100 mg                     | 7/1/2020                | Infugem™                | gemcitabine in sodium<br>chloride injection, for<br>intravenous use | Indicated:<br>• in combination with carboplatin, for the treatment of advanced ovarian cancer that has relapsed at leas<br>6 months after completion of platinum-based therapy.<br>• in combination with paclitaxel, for first-line treatment of metastatic breast cancer after failure of prior<br>anthracycline-containing adjuvant chemotherapy, unless anthracyclines were clinically contraindicated.<br>• in combination with cisplatin for the treatment of non-small cell lung cancer.                                | t<br>128                          | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 6/17/2020            |
| Drugs       | J9200         | Injection, floxuridine, 500 mg  | 500 mg                     | 1/1/2000                | N/A                     | floxuridine for injection, for<br>intra-arterial infusion           | Effective in the palliative management of gastrointestinal adenocarcinoma metastatic to the liver, when<br>given by continuous regional intra-arterial infusion in carefully selected patients who are considered<br>incurable by surgery or other means. Patients with known disease extending beyond an area capable of   | 5                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/26/2018           |
| Drugs       | J9201         | Injection, gemcitabine<br>hydrochloride, not otherwise<br>specified, 200 mg                             | 200 mg                     | 1/1/2000                | Gemzar®                 | gemcitabine for injection, for<br>intravenous use                   | Indicated:<br>• In combination with carboplatin, for the treatment of advanced ovarian cancer that has relapsed at leas<br>6 months after completion of platinum-based therapy.<br>• In combination with paclitaxel, for first-line treatment of metastatic breast cancer after failure of prior<br>anthroxycline-containing adjuvant chemotherapy, unless anthraxyclines were (inicially contraindicated.<br>• In combination with cisplatin for the treatment of non-small cell lung cancer.                                | t<br>64                           | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 1/9/2020             |
| Biologicals | J9203         | Injection, gemtuzumab<br>ozogamicin, 0.1 mg   | 0.1 mg                     | 1/1/2018                | Mylotarg™               | gemtuzumab ozogamicin<br>injection, for intravenous use             | Indicated for:<br>• the treatment of newly-diagnosed CD33-positive acute myeloid leukemia (AML) in adults.<br>• the treatment of newly-diagnosed CD33-positive acute myeloid leukemia (AML) in pediatric patients 1   | 275                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• Newly-diagnosed CD33-<br>positive acute myeloid<br>leukemia: 1 month of age and<br>older | 7/28/2020            |
| Biologicals | J9204         | Injection, mogamulizumab-<br>kpkc, 1 mg   | 1 mg                       | 10/1/2019               | Poteligeo®              | mogamulizumab-kpkc<br>injection, for intravenous use                | Indicated for the treatment of adult patients with relapsed or refractory mycosis fungoides or Sézary<br>syndrome after at least one prior systemic therapy.  | 700                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/27/2019            |
| Drugs       | J9205         | Injection, irinotecan liposome,<br>1 mg   | 1 mg                       | 1/1/2017                | Onivyde**               | irinotecan liposome injection<br>for intravenous use                | Indicated:<br>- in combination with fluorouracil and leucovorin, for the treatment of patients with metastatic<br>adencarinoma of the pancreas after disease progression following gemcitabine-based therapy.<br>- in combination with oxiliplatin, fluorouracil and leucovorin, for the first-line treatment of adult patients<br>with metastatic pancreatic adencarcinoma.<br>Limitation of Use: Onivyde is not indicated as a single agent for the treatment of patients with metastatic<br>adencarcinoma of the pancreas. |                                   | 18 years  | N/A         | N/A                    | Ÿ               | ¥                               |  | 3/22/2024            |
| Drugs       | J9206         | Injection, irinotecan, 20 mg  | 20 mg                      | 1/1/2000                | Camptosar®              | irinotecan injection,<br>intravenous infusion                       | Indicated for:<br>• First-line therapy in combination with 5-fluorouracil and leucovorin for patients with metastatic<br>carcinoma of the colon or rectum.  | 88                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 4/10/2019            |
| Drugs       | J9207         | Injection, ixabepilone, 1 mg  | 1 mg                       | 1/1/2009                | lxempra®                | ixabepilone for injection, for<br>intravenous use                   | Indicated for the treatment   | 180                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 2/23/2023            |

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| medicaid/medi | icaid-ncci-ed | lit-files   |                            |                         |                            |  |  |                                   |   |             |                        |                 |                                 |  |                       |
|---------------|---------------|---|----------------------------|-------------------------|----------------------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                 | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs         | J9208         | Injection, ifosfamide, 1 gram   | 1 g                        | 1/1/2000                | lfex®                      | ifosfamide for injection,<br>intravenous use                                     | Indicated for use in combination with certain other approved antineoplastic agents for third-line<br>chemotherapy of general testicular cancer. It should be used in combination with mesna for prophylaxis<br>of hemorrhagic cystits.                         | 30                                | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 6/4/2019              |
| Drugs         | J9209         | Injection, mesna, 200 mg  | 200 mg                     | 1/1/2000                | Mesnex*                    | mesna injection solution   | Indicated as a prophylactic agent in reducing the incidence of ifosfamide-induced hemorrhagic cystitis.  | 90                                | 18 years  | N/A         | N/A                    | Y               | ¥                               |  | 8/5/2021              |
| Biologicals   | J9210         | Injection, emapalumab-Izsg, 1<br>mg   | 1 mg                       | 10/1/2019               | Gamifant™                  | emapalumab-Izsg injection,<br>for intravenous use                                | Indicated for the treatment of adult and pediatric (newborn and older) patients with primary<br>hemophagocytic lymphohistiocytosis (HLH) with refractory, recurrent or progressive disease or intolerance<br>with conventional HLH therapy.                    | 9 14,000                          | N/A   | N/A         | N/A                    | ¥               | Ÿ                               |  | 5/27/2020             |
| Drugs         | J9211         | Injection, idarubicin<br>hydrochloride, 5 mg                                | 5 mg                       | 1/1/2000                | Idamycin®                  | idarubicin hydrochloride for<br>injection  | Indicated in combination with other approved antileukemic drugs for the treatment of acute myeloid<br>leukemia in adults. This includes French-American-British (FAB) classifications M1 through M7.   | 36                                | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/31/2018            |
| Biologicals   | J9214         | Injection, interferon, alfa-2b, recombinant, 1 million units                | 1 million units            | 1/1/2000                | Intron® A                  | interferon alfa-2b<br>recombinant for injection                                  | Indicated for: hairy cell leukemia, malignant melanoma, follicular lymphoma, condylomata acuminata,<br>AIDS-related Kaposi's sarcoma, chronic hepatitis C and chronic hepatitis B. Please see package insert for<br>additional information on each indication. | 1,050                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific: 18 years<br>and older for all indications<br>except chronic Hepatitis B and<br>C. | 6/4/2019              |
| Biologicals   | J9215         | Injection, interferon, alfa-n3,<br>(human leukocyte derived),<br>250,000 IU | 250,000 IU                 | 1/1/2000                | Alferon® N                 | interferon alfa-n3 injection   | Indicated for condyloma acuminata.   | 100                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/4/2018             |
| Biologicals   | J9216         | Injection, interferon, gamma-<br>1b, 3 million units                        | 3 million units            | 1/1/2000                | Actimmune®                 | interferon gamma-1b<br>injection, for subcutaneous<br>use                        | Indicated for:<br>• Reducing the frequency and severity of serious infections associated with Chronic Granulomatous<br>Disease (CG)<br>• Delaying time to disease progression in patients with severe, malignant osteoporosis (SMO)                            | 18.67                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | ¥                               | Indication specific age<br>restrictions:<br>CGD: 1 year and older<br>SMO: 1 month and older            | 5/6/2019              |
| Drugs         | J9217         | Leuprolide acetate (for depot<br>suspension), 7.5 mg                        | 7.5 mg                     | 1/1/2000                | Eligard®, Lupron<br>Depot® | leuprolide acetate for<br>injectable suspension, for<br>doses 7.5 mg and greater | Eligard: Indicated for the treatment of advanced prostate cancer.<br>Lupron Depot: Indicated for the treatment of advanced prostatic cancer.   | 6                                 | 18 years  | N/A         | Males Only             | Y               | Y                               |  | 2/19/2024             |
| Drugs         | J9218         | Leuprolide acetate, per 1 mg  | per 1 mg                   | 1/1/2000                | N/A                        | leuprolide acetate injection   | Indicated in the palliative treatment of advanced prostatic cancer.  | 31                                | N/A   | N/A         | Males Only             | Y               | Y                               |  | 2/19/2024             |
| Drugs         | J9223         | Injection, lurbinectedin, 0.1 mg  | 0.1 mg                     | 1/1/2021                | Zepzelca™                  | lurbinectedin for injection,<br>for intravenous use                              | Indicated for the treatment of adult patients with metastatic small cell lung cancer (SCLC) with disease<br>progression on or after platinum-based chemotherapy.   | 160                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/28/2020            |
| Drugs         | J9225         | Histrelin implant (Vantas), 50<br>mg  | 50 mg                      | 1/1/2006                | Vantas®                    | histrelin acetate<br>subcutaneous implant  | Indicated for the palliative treatment of advanced prostate cancer.  | 1                                 | 18 years  | N/A         | Males Only             | Y               | Y                               |  | 2/19/2024             |

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| medicaid/medi | caid-ncci-ed  | it-files  |                            |                         |                           |  |  |                                   |   |             |                        |                 |                                 |  |
|---------------|---------------|---|----------------------------|-------------------------|---------------------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|
| Category      | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name                | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments Last Modified Date  |
| Drugs         | J9226         | Histrelin implant (Supprelin<br>LA), 50 mg  | 50 mg                      | 1/1/2008                | Supprelin <sup>®</sup> LA | histrelin acetate<br>subcutaneous implant  | Indicated for the treatment of children with central precocious puberty (CPP).   | 1                                 | 2 years   | N/A         | N/A                    | ¥               | ¥                               | 2/19/2024  |
| Biologicals   | J9227         | Injection, isatuximab-irfc, 10<br>mg  | 10 mg                      | 10/1/2020               | Sarclisa*                 | isatuximab-irfc injection, for intravenous use   | Indicated<br>• in combination with pomalidomide and dexamethasone, for the treatment of adult patients with<br>multiple myeloma who have received at least two prior therapies including lenalidomide and a  | 700                               | 18 years  | N/A         | N/A                    | Y               | Y                               | 4/26/2021  |
| Biologicals   | J9228         | Injection, ipilimumab, 1 mg   | 1 mg                       | 1/1/2012                | Yervoy®                   | ipilimumab injection, for<br>intravenous use   | Indicated for:<br>• Adjuvant treatment of patients with cutaneous melanoma with pathologic involvement of regional<br>lymph nodes of more than 1 mm who have undergone complete resection, including total   | 2,800                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions: 3/21/2023<br>• Melanoma as a single agent                         |
| Biologicals   | J9229         | Injection, inotuzumab<br>ozogamicin, 0.1 mg   | 0.1 mg                     | 1/1/2019                | Besponsa™                 | inotuzumab ozogamicin<br>injection, for intravenous use  | Indicated for the treatment of relapsed or refractory CD22-positive B-cell precursor acute lymphoblastic<br>leukemia (ALL) in adult and pediatric patients 1 year and older.   | 108                               | 1 year  | N/A         | N/A                    | Y               | Y                               | 5/3/2024   |
| Drugs         | J9245         | Injection, melphalan<br>hydrochloride, not otherwise  | 50 mg                      | 1/1/2000                | Alkeran®                  | melphalan hydrochloride for<br>injection   | Indicated for the palliative treatment of patients with multiple myeloma for whom oral therapy is not<br>appropriate.  | 3                                 | 18 years  | N/A         | N/A                    | Y               | Y                               | 6/17/2020  |
| Drugs         | J9246         | Injection, melphalan<br>(evomela), 1 mg   | 1 mg                       | 7/1/2020                | Evomela®                  | melphalan for injection, for intravenous use   | Indicated for:<br>• use as a high-dose conditioning treatment prior to hematopoietic progenitor (stem) cell transplantation<br>in patients with multiple myeloma.  | 500                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               | 9/28/2021  |
| Drugs         | J9258         | Injection, paclitaxel protein-<br>bound particles (teva), not<br>therapeutically equivalent to<br>J9264, 1 mg             | 1 mg                       | 1/1/2024                | N/A                       | paclitaxel protein-bound<br>particles for injectable<br>suspension (albumin-bound),<br>for intravenous use (Teva)                | Paciltaxel Protein-Bound Particles for Injectable Suspension (albumin-bound) is a microtubule inhibitor<br>indicated for the treatment of:<br>• Metastatic treaset cancer, after failure of combination chemotherapy for metastatic disease or relapse<br>within 6 months of adjuvant chemotherapy. Prior therapy should have included an anthracycline unless<br>clinically contraindicated.<br>• Locally advanced or metastatic non-small cell lung cancer (NSCLC), as first-line treatment in combination<br>with carboplatin, in patients who are not candidates for curative surgery or radiation therapy.<br>• Metastatic adenocarcinoma of the pancreas as first-line treatment, in combination with gemcitabine. | 1,600                             | 18 years  | N/A         | N/A                    | Y               | Ŷ                               | 7/29/2024  |
| Drugs         | J9259         | injection, pacifizarel protein-<br>bound particles (american<br>regent), not therapeutically<br>equivalent to j9264, 1 mg | 1 mg                       | 7/1/2023                | N/A                       | paclitaxel protein-bound<br>particles for injectable<br>suspension, (albumin-bound),<br>for intravenous use<br>(American Regent) | Indicated for the treatment of:<br>• Metastatic breast cancer, after failure of combination chemotherapy for metastatic disease or relapse<br>within 6 months of adjuvant chemotherapy. Prior therapy should have included an anthracycline unless<br>clinically contraindicated.<br>• Locally advanced or metastatic non-small cell lung cancer (NSCLC), as first-line treatment in combination<br>with carobaltatin, in patients who are not candidates for curative surgery or radiation therapy.<br>• Metastatic adenocarcinoma of the pancreas as first-line treatment, in combination with gemcitabine.  | 1,600                             | 18 years  | N/A         | N/A                    | Y               | Ŷ                               | 6/22/2023  |
| Drugs         | J9260         | Methotrexate sodium, 50 mg  | 50 mg                      | 1/1/2000                | N/A                       | methotrexate sodium<br>injection, 50 mg  | <ul> <li>Methotrexate is indicated in the treatment of gestational choriocarcinoma, chorioadenoma destruens<br/>and hydatidiform mole.</li> <li>In acute lymphocytic leukemia, methotrexate is indicated in the prophylaxis of meningeal leukemia and<br/>is used in maintenance therapy in combination with other chernotherapeutic agents. Methotrexate is also</li> </ul>   | 3,000                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• Cancer chemotherapy: None<br>• Polyarticular-course juvenile |

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| nedicaid/medi | HCPCS<br>Code | HCPCS Description                                       | HCPCS Code Billing<br>Unit         | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
|---------------|---------------|---|------------------------------------|-------------------------|------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Drugs         | J9261         | Injection, nelarabine, 50 mg                            | 50 mg                              | 1/1/2007                | Arranon®   | nelarabine injection, for<br>intravenous use  | Indicated for the treatment of patients with T-cell acute lymphoblastic leukemia and T-cell lymphoblastic<br>lymphoma in adult and pediatric patients age 1 year and older whose disease has not responded to or has<br>relapsed following treatment with at least two chemotherapy regimens. | 450                               | 1 year      | N/A         | N/A                    | Y               | ¥                               |          | 12/16/2021            |
| Drugs         | J9263         | Injection, oxaliplatin, 0.5 mg                          | 0.5 mg                             | 1/1/2004                | Eloxatin®  | oxaliplatin injection for<br>intravenous use  | Indicated for:<br>• Adjuvant treatment of stage III colon cancer in patients who have undergone complete resection of the<br>primary tumor.<br>• Treatment of advanced colorectal cancer.   | 1,500                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/4/2019              |
| Drugs         | J9264         | Injection, paclitaxel protein-<br>bound particles, 1 mg | 1 mg                               | 1/1/2006                | Abraxane®  | paclitaxel protein-bound<br>particles for injectable<br>suspension, (albumin-bound),<br>for intravenous use | Indicated for the treatment:<br>• Metastatic breast cancer, after failure of combination chemotherapy for metastatic disease or relapse<br>within six months of adjuvant chemotherapy. Prior therapy should have included an anthracycline unless<br>clinically contraindicated.              | 1,600                             | 18 years    | N/A         | N/A                    | Y               | Ŷ                               |          | 5/25/2023             |
| Biologicals   | J9266         | Injection, pegaspargase, per<br>single dose vial        | per single dose vial<br>(3,750 IU) | 1/1/2000                | Oncaspar*  |   | Indicated as a component of a multi-agent chemotherapeutic regimen for treatment of patients with:<br>• First line acute lymphoblastic leukemia<br>• Acute lymphoblastic leukemia and hypersensitivity to asparaginase  | 6                                 | 1 year      | N/A         | N/A                    | ¥               | ¥                               |          | 8/24/2018             |
| Drugs         | J9267         | Injection, paclitaxel, 1 mg                             | 1 mg                               | 1/1/2015                | Taxol®     | paclitaxel injection  | Indicated for breast cancer, ovarian cancer, non-small cell lung cancer, and AID5-related karposi sarcoma.<br>See package insert for full details of each indication.   | 875                               | 18 years    | N/A         | N/A                    | Ŷ               | Ŷ                               |          | 9/27/2018             |

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•Medically Unlikely Edits (MUEs) are used by NC Medicaid to reduce the improper payment for medical drug claims. A MUE for a HCPCS/CPT code is the maximum units of service that a provider would report under most circumstances for a single baeficiary on a single date of service. Not all HCPCS/CPT codes have a MUE. CMS publishes MUE values on its website: https://www.ems.gov/medicare/coding-biling/ncci-

medicaid/medicaid-ncci-edit-files

| nedicaid/medi |               | inc-mes                                       |                            |                         |                        |  |   |                                   |   |             |   |                 | Rebating            |  | 1 1                   |
|---------------|---------------|---|----------------------------|-------------------------|------------------------|--|---|-----------------------------------|---|-------------|---|-----------------|---------------------|--|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description                             | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name             | Generic Name                                       | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions  | NDC<br>Required | Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs         | J9268         | Injection, pentostatin, per 10<br>mg          | 10 mg                      | 7/15/2001               | Nipent®                | pentostatin for injection                          | Indicated as single-agent treatment for both untreated and alpha-interferon-refractory hairy cell leukemia<br>patients with active disease as defined by clinically significant anemia, neutropenia, thrombocytopenia, or<br>disease-related symptoms.  | 3                                 | 18 years  | N/A         | N/A   | ¥               | Ŷ                   |  | 9/21/2018             |
| Biologicals   | J9269         | Injection, tagraxofusp-erzs, 10<br>micrograms | 10 mcg                     | 10/1/2019               | Elzonris™              | tagraxofusp-erzs injection,<br>for intravenous use | Indicated for the treatment of blastic plasmacytoid dendritic cell neoplasm (BPDCN) in adults and in<br>pediatric patients 2 years and older.   | 2,000                             | 2 years   | N/A         | N/A   | Y               | Y                   |  | 10/3/2019             |
| Biologicals   | J9271         | injection, pembrolizumab, 1<br>mg             | 1 mg                       | 1/1/2016                | Keytruda®              |  | Nelanoma:<br>1. Indicated for the treatment of patients with unresectable or metastatic melanoma.<br>2. Indicated for the adjuvant treatment of adult and pediatric (12 years and older) patients with Stage IIB,<br>IIC, or III melanoma following complete resection.<br>Non-Small Cell Lung Cancer (NSCLC):<br>1. Indicated in combination with pemetrexed and platinum chemotherapy, as first-line treatment of<br>patients with metastatic nonsquarmous MSCLC, with no EGFR or ALK genomic tumor aberrations.<br>2. Indicated as a single agent for the treatment of patients with metastatic NSCL whose tumors express<br>PO-L1 (TFS ± 1%) as determined by an FDA-approved test, with disease progression on or after platinum-<br>containing chemotherapy. Patients with EGFR or ALK genomic tumor aberrations.<br>3. Indicated as a single agent for the first-line treatment of patients with stage III NSCLC, who are not<br>candidates for singular rescue or definitive chemocalation, or metastic NSCL who are not<br>candidates for singular rescue on definitive chemocalation, protestatic NSCL who are not<br>candidates for surgical rescue on definitive chemocalation, protestatic NSCL who are not<br>candidates for SUP-L1 [Tumor Proportion SCore (TFS) 21%] as determined by an FDA-approved test, with no EGFR<br>or ALK genomic Lumor aberrations. | 400                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A   | Y               | Y                   | The safety and effectiveness of<br>Keytruda as a single agent<br>have been established in<br>pediatric patients with<br>melanoma, CHL, PMBCL, MCC,<br>MSI-H or dMMR cancer, and<br>TMB-H cancer. The safety and<br>effectiveness of Keytruda in<br>pediatric patients have not<br>been established in the other<br>approved indications. | 7/29/2024             |
| Biologicals   | J9272         | Injection, dostarlimab-gxly, 10<br>mg         | 10 mg                      | 1/1/2022                | Jemperli               | dostariimab-gely injection,<br>for intravenous use | Endometrial Cancer (EC)<br>indicated for the treatment of adult patients with mismatch repair deficient (dMMR) recurrent or<br>advanced endometrial cancer, as determined by an FDA-approved test, that has progressed on or<br>following pior treatment with a platinum-containing regimen in any setting and are not candidates for<br>curative surgery or radiation.<br>indicated in combination with carboplatin and pacitiaxel, followed by Jemperli as a single agent, for the<br>treatment of adult patients with primary advanced or recurrent endometrial cancer that is mismatch<br>repair deficient (dMMR), as determined by an FDA-approved test, or microsatellite instability-high (MSI-<br>H).<br>Mismatch Repair Deficient Recurrent or Advanced Solid Tumors<br>I indicated for the treatment of adult patients with mismatch repair deficient (dMMR) recurrent or<br>advanced as a single agent, for solid tumors, as determined by an FDA-approved test, that have<br>progressed on or following prior treatment and who have no satisfactory alternative treatment options.   | 150                               | 18 years  | N/A         | Endometrial<br>Cancer: Females<br>only<br>Solid Tumors:<br>None | Y               | Y                   |  | 9/13/2023             |
| Biologicals   | J9273         | Injection, tisotumab vedotin-<br>tftv, 1 mg   | 1 mg                       | 4/1/2022                | Tivdak™                |  | Indicated for the treatment of adult patients with recurrent or metastatic cervical cancer with disease<br>progression on or after chemotherapy.  | 400                               | 18 years  | N/A         | N/A   | Y               | Y                   |  | 3/21/2022             |
| Biologicals   | J9274         | Injection, tebentafusp-tebn, 1<br>microgram   | 1 mcg                      | 10/1/2022               | Kimmtrak®              |  | Indicated for the treatment of HLA-A*02:01-positive adult patients with unresectable or metastatic uveal<br>melanoma.   | 500                               | 18 years  | N/A         | N/A   | Y               | Y                   |  | 9/15/2022             |
| Drugs         | J9280         | Injection, mitomycin, 5 mg                    | 5 mg                       | 1/1/2000                | Mutamycin <sup>®</sup> | mitomycin for injection, 5 mg                      | Mitomycin is not recommended as single-agent, primary therapy. It has been shown to be useful in the<br>therapy of disseminated adenocarcinoma of the stomach or pancreas in proven combinations with other<br>approved chemotherapeutic agents and as palliative treatment when other modalities have failed.<br>Mitomycin is not recommended to replace appropriate surgery and/or radiotherapy.  | 10                                | 18 years  | N/A         | N/A   | Y               | Y                   |  | 6/7/2019              |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
 The HCPCS Code effective date represents the date the HCPCS code was established

• Procedure codes for covered devices and vaccines are not required to be from a rebating labeler/manufacturer as they are not classified as covered outpatient drugs.

| medicaid/medi | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments                           | Last Modified<br>Date |
|---------------|---------------|---|----------------------------|-------------------------|------------|--|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|------------------------------------|-----------------------|
| Drugs         | J9281         | Mitomycin pyelocalyceal instillation, 1 mg  | 1 mg                       | 1/1/2021                | Jelmyto™   | mitomycin for pyelocalyceal solution                       | Indicated for the treatment of adult patients with low-grade Upper Tract Urothelial Cancer (LG-UTUC).  | 400                               | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |                                    | 12/28/2020            |
| Biologicals   | J9286         | Injection, glofitamab-gobm,<br>2.5 mg   | 2.5 mg                     | 1/1/2024                | Columvi*   | glofitamab-gxbm injection,<br>for intravenous use          | Indicated for the treatment of adult patients with relapsed or refractory diffuse large B-cell lymphoma,<br>not otherwise specified (DLBCL, NOS) or large B-cell lymphoma (LBCL) arising from follicular lymphoma,<br>after two or more lines of systemic therapy.   | 24                                | 18 years    | N/A         | N/A                    | Ŷ               | ¥                               |                                    | 12/22/2023            |
| Drugs         | J9293         | Injection, mitoxantrone<br>hydrochloride, per 5 mg                                    | 5 mg                       | 1/1/2000                | N/A        | mitoxantrone hydrochloride<br>injection, solution          | Indicated:<br>• For reducing neurologic disability and/or the frequency of clinical relapses in patients with secondary<br>(thronic) progressive, progressive relapsing, or worsening relapsing-remitting multiple sclerosis (i.e.,<br>patients whose neurologic status is significantly abnormal between relapses).<br>Mittoantrone is on indicated in the treatment of patients with primary progressive multiple sclerosis.<br>• In combination with corticosteroids is indicated as initial chemotherapy for the treatment of patients<br>with pair relate to advanced hormone-refractory prostate cancer.<br>• In combination with other approved drug(s) is indicated in the initial therapy of acute nonlymphocytic<br>leukemia (ANLI) in adults. This category includes myelogenous, promyelocytic, monocytic, and erythroid<br>acute leukemias.   | 30                                | 18 years    | N/A         | N/A                    | v               | Y                               | Lifetime Maximum Dose: 70<br>units | 10/31/2018            |
| Drugs         | J9294         | Injection, pemetrexed<br>(hospira), not therapeutically<br>equivalent to j9305, 10 mg |                            | 4/1/2023                | N/A        | pemetrexed for injection, for<br>intravenous use (Hospira) | Indicated:<br>• In combination with cisplatin for the initial treatment of patients with locally advanced or metastatic,<br>non-squamous, non-small cell lung cancer (NSCLC).<br>• As a single agent for the maintenance treatment of patients with locally advanced or metastatic, non -<br>squamous NSCL Whose disease has not progressed after four cycles of platinum-based first-line<br>chemotherapy.<br>• As a single agent for the treatment of patients with recurrent, metastatic non-squamous, NSCLC after<br>prior chemotherapy.<br>Limitations of Use: Penetrexed for Injection is not indicated for the treatment of patients with squamous<br>cell, non-small cell lung cancer.<br>• Initial treatment, in combination with cisplatin, of patients with malignant pleural mesothelioma whose<br>disease is unresectable or who are otherwise not candidates for curative surgery. |                                   | 18 years    | N/A         | N/A                    | Y               | Ŷ                               |                                    | 3/16/2023             |

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The Max Daily Units for radiopharmaceuticals represents one therapeutic dose or diagnostic dose.
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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age                             | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|------------|---|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Biologicals | J9295         | Injection, necitumumab, 1 mg   | 1 mg                       | 1/1/2017                | Portrazza™ | necitumumab injection, for<br>intravenous use                       | Indicated, in combination with gemcitabine and cisplatin, for first-line treatment of patients with<br>metastatic squamous non-small cell lung cancer.<br>Limitation of Use: Portrazza is not indicated for treatment of non-squamous non-small cell lung cancer.  | 3,200                             | 18 years                                | N/A         | N/A                    | Y               | Ŷ                               |  | 7/2/2018              |
| Drugs       | J9296         | Injection, pemetrexed<br>(accord), not therapeutically<br>equivalent to j9305, 10 mg | 10 mg                      | 4/1/2023                | N/A        | pemetrexed injection, for intravenous use (Accord)                  | Indicated:<br>• In combination with pembrolizumab and platinum chemotherapy, for the initial treatment of patients<br>with metastatic non-squamous non-small cell lung cancer (NSCLC), with no EGFR or ALK genomic turnor<br>aberrations.<br>• In combination with cisplatin for the initial treatment of patients with locally advanced or metastatic,<br>non-squamous, NSCLC.<br>• as a single agent for the maintenance treatment of patients with locally advanced or metastatic, non-<br>squamous NSCLC whose disease has not progressed after four cycles of platinum-based first-line<br>chemotherapy.<br>• as a single agent for the treatment of patients with recurrent, metastatic non-squamous, NSCLC after<br>prior chemotherapy. | 300                               | 18 years                                | N/A         | N/A                    | Y               | Ŷ                               |  | 3/16/2023             |
| Drugs       | J9297         | Injection, pemetrexed<br>(sandoz), not therapeutically<br>equivalent to j9305, 10 mg | 10 mg                      | 4/1/2023                | N/A        | pemetrexed injection, for<br>intravenous use (Sandoz)               | Indicated:<br>• in combination with pembrolizumab and platinum chemotherapy, for the initial treatment of patients<br>with metastatic non-squamous non-small cell lung cancer (NSCLC), with no EGFR or ALK genomic tumor   | 300                               | 18 years                                | N/A         | N/A                    | Y               | Y                               |  | 3/16/2023             |
| Biologicals | J9298         | Injection, nivolumab and relatlimab-rmbw, 3 mg/1 mg                                  | 3 mg/1 mg                  | 10/1/2022               | Opdualag™  | nivolumab and relatlimab-<br>rmbw injection, for<br>intravenous use | Indicated for the treatment of adult and pediatric patients 12 years of age or older with unresectable or<br>metastatic melanoma.  | 320                               | 12 years                                | N/A         | N/A                    | Y               | Y                               |  | 9/15/2022             |
| Biologicals | J9299         | Injection, nivolumab, 1 mg   | 1 mg                       | 1/1/2016                | Opdivo®    | nivolumab injection, for<br>intravenous use                         | Indicated for:<br>Melanoma:  | 1,260                             | Indication Specific<br>Age Restrictions | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:         | 5/3/2024              |
| Biologicals | J9301         | Injection, obinutuzumab, 10<br>mg  | 10 mg                      | 1/1/2015                | Gazyva®    | obinutuzumab Injection, for<br>intravenous use                      | Indicated:<br>• In combination with chlorambucil, for the treatment of patients with previously untreated chronic<br>lymphocytic leukemia.<br>• In combination with bendamustine followed by Gazyua monotherapy, for the treatment of patients with<br>follicular lymphoma who relapsed after, or are refractory to, a rituzmia-containing regimen.  | 400                               | 18 years                                | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 7/16/2018             |
| Biologicals | J9302         | Injection, ofatumumab, 10 mg   | 10 mg                      | 1/1/2011                | Arzerra*   | ofatumumab injection, for<br>intravenous use                        | Indicated for the treatment of chronic lymphocytic leukemia (CLL):<br>• in combination with chlorambucil, for the treatment of previously untreated patients with CLL for whom<br>fludarabine-based therapy is considered<br>inappropriate.<br>• in combination with fludarabine and cyclophosphamide for the treatment of patients with relapsed CLL<br>• for extended treatment of patients who are in complete or partial response after at least two lines of<br>therapy for recurrent or progressive CLL.<br>• for the treatment of patients with CLL refractory to fludarabine and alemtuzumab.  | 1,000                             | 18 years                                | N/A         | N/A                    | Y               | ¥                               | Pregnancy: May cause fetal B-<br>cell depletion. | 7/16/2018             |
| Biologicals | J9303         | Injection, panitumumab, 10<br>mg   | 10 mg                      | 1/1/2008                | Vectibix*  | panitumumab injection, for<br>intravenous use                       | Indicated for the treatment of wild-type RAS (defined as wild-type in both KRAS and NRAS as determined<br>by an FDA-approved test for this use) metastatic colorectal cancer (mCRC):<br>- In combination with Foldra for first-line treatment.<br>- As monotherapy following disease progression after prior treatment with fluoropyrimidine, oxaliplatin,<br>and irinotecan-containing chemotherapy.  | 270                               | 18 years                                | N/A         | N/A                    | Ŷ               | Y                               |  | 6/4/2019              |
| Drugs       | J9304         | Injection, pemetrexed<br>(pemfexy), 10 mg  | 10 mg                      | 10/1/2020               | Pemfexy™   | pemetrexed injection, for<br>intravenous use                        | Indicated:<br>• In combination with cisplatin for the initial treatment of patients with locally advanced or metastatic<br>non-squamous, non-small cell lung cancer (NSCLC).<br>• as a single agent for the maintenance treatment of patients with locally advanced or metastatic non-<br>squamous NSCLC whose disease has not progressed after four cycles of platinum-based first-line<br>chemotherapy.  | 300                               | 18 years                                | N/A         | N/A                    | Ŷ               | Ŷ                               |  | 1/23/2023             |
| Drugs       | J9305         | Injection, pemetrexed, not<br>otherwise specified, 10 mg                             | 10 mg                      | 10/1/2020               | Alimta®    | pemetrexed for injection, for<br>intravenous use                    | Indicated:<br>• In combination with cisplatin for the initial treatment of patients with locally advanced or metastatic,   | 300                               | 18 years                                | N/A         | N/A                    | Y               | Y                               |  | 12/12/2022            |
| Biologicals | J9306         | Injection, pertuzumab, 1 mg  | 1 mg                       | 1/1/2014                | Perjeta®   | pertuzumab injection, for<br>intravenous use                        | Indicated for:<br>• Use in combination with trasturumab and docetaxel for treatment of patients with HER2-positive<br>metastatic breast cancer (MBC) who have not received prior anti-HER2 thrapy or chemotherapy for<br>metastatic disease.<br>• Use in combination with trasturumab and chemotherapy as  | 1,260                             | 18 years                                | N/A         | N/A                    | Ŷ               | Y                               |  | 7/2/2018              |
| Drugs       | J9307         | Injection, pralatrexate, 1 mg  | 1 mg                       | 1/1/2011                | Folotyn®   | pralatrexate injection, for<br>intravenous use                      | Indicated for the treatment of patients with relapsed or refractory peripheral T-cell lymphoma.  | 400                               | 18 years                                | N/A         | N/A                    | Ŷ               | Y                               |  | 8/24/2018             |
| Biologicals | 19308         | Injection, ramucirumab, 5 mg   | 5 mg                       | 1/1/2016                | Cyramza®   | ramucirumab injection, for intravenous use                          | Indicated:<br>• As a single agent or in combination with paclitaxel, for treatment of advanced gastric or gastro-<br>esophageal junction adenocarcinoma, with disease progression on or after prior fluoropyrimidine- or<br>platinum-containing chemotherapy.<br>• In combination with docetaxel, for treatment of metastatic non-small cell lung cancer with disease  | 900                               | 18 years                                | N/A         | N/A                    | Y               | Y                               |  | 6/17/2020             |

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| medicalu/medii | caid-ncci-ed  | dit-files  | i.                         |                         |                 |  | r  |                                   |   |             |                        |                 |                                 |  |                       |
|----------------|---------------|--|----------------------------|-------------------------|-----------------|--|--|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category       | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name      | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Biologicals    | 19309         | Injection, polatuzumab<br>vedotin-piiq, 1 mg                                       | 1 mg                       | 1/1/2020                | Polivy®         | polatuzumab vedotin-piiq for<br>injection, for intravenous use                           | Indicated:<br>Indicated:<br>In combination with bendamustine and a rituximab product for the treatment of adult patients with<br>relapsed or refractory diffuse large B-cell lymphoma, not otherwise specified, after at least two prior<br>therapies.<br>In combination with a rituximab product, cyclophosphamide, doxorubicin, and prednisone (R-CHP) for<br>the treatment of adult patients who have previously untreated diffuse large B-cell lymphoma (DLBCL), not<br>otherwise specified (NGS) or high-grade B-cell lymphoma (HGBL) and who have an International<br>Prognostic Index source 02 or greater. | 560                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 5/25/2023             |
| Biologicals    | J9311         | Injection, rituximab 10 mg and<br>hyaluronidase                                    | 10 mg                      | 1/1/2019                | Rituxan Hycela® | rituximab and hyaluronidase<br>human injection, for<br>subcutaneous use                  | Indicated for the treatment of adult patients with:<br>• Follicular Lymphoma (FL):<br>o Relapsed or fractaory, follicular lymphoma as a single agent<br>o Previously untreated follicular lymphoma in combination with first line chemotherapy and, in patients<br>achieving a complete or patial response to rituximab in combination with chemotherapy, as single-agent<br>maintenance therapy   | 700                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 4/19/2019             |
| Biologicals    | J9312         | Injection, rituximab, 10 mg  | 10 mg                      | 1/1/2019                | Rituxan®        | rituximab injection, for<br>intravenous use  | Indicated for the treatment of adult patients with:<br>• Non-Hodgkin's Lymphona (NHL)<br>- Relapsed or refractory, low grade or follicular, CD20-positive B-cell NHL as a single agent.<br>- Previously untreated follicular, CD20-positive, B-cell NHL in combination with first line chemotherapy  | 500                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication Specific:<br>• CLL, RA, PV: 18 years of age<br>and older<br>• GPA and MPA: 2 years of age | 1/13/2022             |
| Drugs          | J9314         | Injection, pemetrexed (teva),<br>not therapeutically equivalent<br>to J9305, 10 mg | 10 mg                      | 1/1/2023                | N/A             | pemetrexed for injection, for<br>intravenous use (Teva)                                  | Indicated:<br>in combination with pembrolizumab and platinum chemotherapy, for the initial treatment of patients<br>with metastatic non-squamous non-small cell lung cancer (NSCLC), with no epidermal growth factor<br>receptor (EGFR) or anaplastic lymphoma kinase (ALK) genomic tumor aberrations.<br>in combination with cisplatin for the initial treatment of patients with locally advanced or metastatic,<br>non-squamous NSCLC.  | 300                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/12/2022            |
| Biologicals    | J9316         | Injection, pertuzumab,<br>trastuzumab, and<br>hyaluronidase-zzxf, per 10 mg        | 10 mg                      | 1/1/2021                | Phesgo™         | pertuzumab, trastuzumab,<br>and hyaluronidase-zzxf<br>injection, for subcutaneous<br>use | <ul> <li>as a single agent for the maintenance treatment of patients with locally advanced or metastatic, non-<br/>indicated for:</li> <li>Use in combination with chemotherapy as:</li> <li>o neoadjuvant treatment of patients with HEC2-positive, locally advanced, inflammatory, or early stage<br/>breast cance (ether greater than 2 m in diameter or node positive) as part of a complete treatment</li> </ul>  | 300                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 12/28/2020            |
| Biologicals    | J9317         | Injection, sacituzumab<br>govitecan-hziy, 2.5 mg                                   | 2.5 mg                     | 1/1/2021                | Trodelvy™       | sacituzumab govitecan-hziy<br>for injection, for intravenous<br>use                      | Indicated for the treatment of adult patients with:<br>• Unresectable locally advanced or metastatic triple-negative breast cancer (mTNBC) who have received<br>two or more prior systemic therapies, at least one of them for metastatic disease.<br>• Locally advanced or metastatic urothelial cancer (mUC) who have previously received a platinum-  | 2,304                             | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 3/16/2023             |
| Drugs          | J9318         | Injection, romidepsin, non-<br>lyophilized, 0.1 mg                                 | 0.1 mg                     | 10/1/2021               | N/A             | romidepsin for injection, for<br>intravenous use (non-<br>lyophilized)                   | Indicated for:<br>• The treatment of cutaneous T-cell lymphoma (CTCL) in adult patients who have received at least one<br>prior systemic therapy.  | 2,200                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 1/13/2022             |
| Drugs          | J9319         | Injection, romidepsin,<br>lyophilized, 0.1 mg                                      | 0.1 mg                     | 10/1/2021               | Istodax®        | romidepsin for injection, for<br>intravenous use (lyophilized)                           | Indicated for:   | 1600                              | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/29/2021             |
| Drugs          | J9320         | Injection, streptozocin, 1 gram  | 1 g                        | 1/1/2000                | Zanosar®        | streptozocin powder, for<br>solution   | Indicated in the treatment of metastatic islet cell cancer of pancreas.  | 20                                | N/A   | N/A         | N/A                    | Y               | Y                               |  | 6/7/2019              |
| Biologicals    | J9321         | Injection, epcoritamab-bysp,<br>0.16 mg  | 0.16 mg                    | 1/1/2024                | Epkinly™        | epcoritamab-bysp injection,<br>for subcutaneous use                                      | indicated for the treatment of:<br>Diffuse Large B-cell Lymphoma and High-grade B-cell Lymphoma<br>- adult patients with relapsed or refractory diffuse large B-cell lymphoma (DLBCL), not otherwise<br>specified, including DLBCL arising from indolent lymphoma, and high-grade B-cell lymphoma after two or   | 1,500                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 7/29/2024             |
| Drugs          | J9323         | Injection, pemetrexed ditromethamine, 10 mg  | 10 mg                      | 7/1/2023                | N/A             | pemetrexed ditromethamine<br>for injection, for intravenous<br>use (Hospira)             | Indicated:<br>In combination with cisplatin for the initial treatment of patients with locally advanced or metastatic,<br>non-squamous, non-small cell lung cancer (NSCLC).<br>• As a single agent for the maintenance treatment of patients with locally advanced or metastatic, non-<br>squamous NSCLC whose disease has not progressed after four cycles of platinum-based first-line<br>chemotherapy.<br>• As a single agent for the treatment of patients with recurrent, metastatic non-squamous, NSCLC after<br>prior chemotherapy.   | 300                               | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 6/22/2023             |
| Drugs          | J9324         | Injection, pemetrexed<br>(pemrydi rtu), 10 mg                                      | 10 mg                      | 1/1/2024                | Pemrydi RTU®    | pemetrexed injection, for<br>intravenous use (Shilpa)                                    | Pemetrexed Injection is indicated:<br>- in combination with pembrolizumab and platinum chemotherapy, for the initial treatment of patients<br>with metastatic non-squamous non-small cell lung cancer (NSCLC), with no EGFR or ALK genomic tumor<br>aberrations.<br>- in combination with cisplatin for the initial treatment of patients with locally advanced or metastatic,<br>non-squamous NSCLC.  | 300                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 5/3/2024              |
| Biologicals    | J9325         | Injection, talimogene<br>laherparepvec, per 1 million<br>plaque forming units      | 1 million PFU              | 1/1/2017                | Imlygic®        | talimogene laherparepvec<br>suspension for intralesional<br>injection                    | Indicated for the local treatment of unresectable cutaneous, subcutaneous, and nodal lesions in patients<br>with melanoma recurrent after initial surgery.<br>Limitations of Use: Imlygic has not been shown to improve overall survival or have an effect on visceral   | 800                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 7/16/2018             |
| Drugs          | J9328         | Injection, temozolomide, 1 mg  | 1 mg                       | 1/1/2010                | Temodar®        | temozolomide for injection,<br>for intravenous use                                       | Indicated in adult patients for:<br>• Treatment of newly diagnosed glioblastoma multiforme (GBM) concomitantly with radiotherapy and<br>then as maintenance treatment.   | 6,200                             | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 10/26/2023            |

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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name       | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Drugs       | 19330         | Injection, temsirolimus, 1 mg                                | 1 mg                       | 1/1/2009                | Torisel®         | temsirolimus injection, for<br>intravenous use  | Indicated for the treatment of advanced renal cell carcinoma.   | 125                               | N/A         | N/A         | N/A                    | ¥               | Y                               |  | 9/25/2018             |
| Drugs       | J9331         | Injection, sirolimus protein-<br>bound particles, 1 mg       | 1 mg                       | 1/1/2000                | Fyarro™          | sirolimus protein-bound<br>particles for injectable<br>suspension (albumin-bound),<br>for intravenous use | Indicated for the treatment of adult patients with locally advanced unresectable or metastatic malignant<br>perivascular epithelioid cell tumor (PEComa).   | 1,200                             | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 6/6/2022              |
| Biologicals | J9332         | Injection, efgartigimod alfa-<br>fcab, 2mg                   | 2 mg                       | 7/1/2022                | Vyvgart**        | efgartigimod alfa-fcab<br>injection, for intravenous use  | Indicated for the treatment of generalized myasthenia gravis (gMG) in adult patients who are anti-<br>acetylcholine receptor (AChR) antibody positive.  | 2,400                             | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |  | 6/6/2022              |
| Biologicals | 19333         | Injection, rozanolixizumab-<br>noli, 1 mg                    | 1 mg                       | 1/1/2024                | Rystiggo®        | rozanolixizumab-noli<br>injection, for subcutaneous<br>use  | Indicated for the treatment of generalized myasthenia gravis (gMG) in adult patients who are anti-<br>acetylcholine receptor (AChR) or antimuscle-specific tyrosine kinase (MuSK) antibody positive.  | 4,200                             | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |  | 12/22/2023            |
| Biologicals | J9334         | Injection, efgartigimod alfa, 2<br>mg and hyaluronidase-qvfc | 2 mg                       | 1/1/2024                | Vyvgart® Hytrulo | efgartigimod alfa and<br>hyaluronidase-qvfc injection,<br>for subcutaneous use                            | Indicated for the treatment of adult patients with:<br>- generalized myasthenia gravis (gMG) who are anti-acetylcholine receptor (AChR) antibody positive<br>- chronic inflammatory demyelinating polyneuropathy (CIDP)   | 2,016                             | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 7/29/2024             |
| Drugs       | J9340         | Injection, thiotepa, 15 mg                                   | 15 mg                      | 1/1/2000                | N/A              | thiotepa injection, powder,<br>lyophilized, for solution  | Thiotepa has been tried with varying results in the palliation of a wide variety of neoplastic diseases.<br>However, the most consistent results have been seen in the following tumors: adenocarcinoma of the<br>breast; adenocarcinoma of the ovary, for controlling intracavitary effusions secondary to diffuse or<br>localized neoplastic diseases of various serosal cavities; for the treatment of superficial applicary<br>carcinoma of the urinary bladder. Thiotepa has been effective against other lymphomas, such as<br>lymphosarcoma and Hodgkin's disease. | 20                                | 18 years    | N/A         | N/A                    | ¥               | Ŷ                               |  | 9/21/2018             |
| Biologicals | J9345         | Injection, retifanlimab-dlwr, 1<br>mg                        | 1 mg                       | 10/1/2023               | Zynyz™           | retifanlimab-dlwr injection,<br>for intravenous use   | Indicated for the treatment of adult patients with metastatic or recurrent locally advanced Merkel cell<br>carcinoma.   | 1,000                             | 18 years    | N/A         | N/A                    | Y               | Y                               | 9/2023: NC Suggested Max<br>Monthly Units updated from<br>500 units to 1,000 units | 9/28/2023             |
| Biologicals | J9347         | Injection, tremelimumab-actl<br>1 mg                         | 1 mg                       | 7/1/2023                | Imjudo*          | tremelimumab-actl injection,<br>for intravenous use   | Indicated:<br>• in combination with durvalumab, for the treatment of adult patients with unresectable hepatocellular<br>carcinoma (uHCC).<br>• in combination with durvalumab and platinum-based chemotherapy for the treatment of adult patients   | 300                               | 18 years    | N/A         | N/A                    | Y               | Y                               |  | 6/22/2023             |
| Biologicals | J9348         | Injection, naxitamab-gqgk, 1<br>mg                           | 1 mg                       | 7/1/2021                | Danyelza®        | naxitamab-gqgk injection, for<br>intravenous use  | Indicated, in combination with granulocyte-macrophage colony-stimulating factor (GM-CSF), for the<br>treatment of pediatric patients 1 year of age and older and adult patients with relapsed or refractory high-<br>risk neuroblastoma in the bone or bone marrow who have demonstrated a partial response, minor<br>response, or stable disease to prior therapy.   | 800                               | 1 year      | N/A         | N/A                    | ¥               | Y                               |  | 6/28/2021             |

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| medicaid/med | icaid-ncci-ed | lit-files   |                           |                           |            |  |   |                                   |             |             |                        |                 |                                 |          |                       |
|--------------|---------------|---|---------------------------|---------------------------|------------|--|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Category     | HCPCS<br>Code | HCPCS Description                                     | HCPCS Code Billin<br>Unit | g HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
| Biologicals  | J9349         | Injection, tafasitamab-cxix, :<br>mg                  | 2 2 2 mg                  | 4/1/2021                  | Monjuvi*   | tafasitamab-cxix for injection,<br>for intravenous use             | Indicated in combination with lenalidomide for the treatment of adult patients with relapsed or refractory<br>diffuse large B-cell lymphoma (DLBCL) not otherwise specified, including DLBCL arising from low grade<br>lymphoma, and who are not eligible for autologous stem cell transplant (ASCT).   | 5,400                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 3/25/2021             |
| Biologicals  | J9350         | Injection, mosunetuzumab-<br>axgb, 1 mg               | 1 mg                      | 7/1/2023                  | Lunsumio™  | mosunetuzumab-axgb<br>injection, for intravenous use               | Indicated for the treatment of adult patients with relapsed or refractory follicular lymphoma after two or<br>more lines of systemic therapy.   | 123                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/22/2023             |
| Drugs        | J9351         | injection, topotecan, 0.1 mg                          | 5 0.1 mg                  | 1/1/2011                  | Hycamtin®  | topotecan for injection  | Indicated for:<br>• Metastatic carcinoma of the ovary after disease progression on or after initial or subsequent<br>chemotherapy.<br>• Small cell lung cancer platinum-sensitive disease in patients who progressed after first-line<br>chemotherapy.<br>• Combination therapy with cisplatin for Stage IV-B, recurrent, or persistent carcinoma of the cervix which<br>is not amenable to curative treatment. | 400                               | 18 years    | N/A         | N/A                    | Y               | ¥                               |          | 9/12/2018             |
| Drugs        | J9352         | Injection, trabectedin, 0.1 m                         | g 0.1 mg                  | 1/1/2017                  | Yondelis*  | trabectedin for injection, for intravenous use                     | Indicated for the treatment of patients with unresectable or metastatic liposarcoma or leiomyosarcoma<br>who received a prior anthracycline-containing regimen.   | 80                                | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/12/2018             |
| Biologicals  | 19353         | Injection, margetuximab-<br>cmkb, 5 mg                | 5 mg                      | 7/1/2021                  | Margenza™  | margetuximab-cmkb<br>injection, for intravenous use                | Indicated, in combination with chemotherapy, for the treatment of adult patients with metastatic HER2-<br>positive breast cancer who have received two or more prior anti-HER2 regimens, at least one of which was<br>for metastatic disease.   | 900                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 6/28/2021             |
| Biologicals  | J9354         | Injection, ado-trastuzumab<br>emtansine, 1 mg         | 1 mg                      | 1/1/2014                  | Kadcyla*   | ado-trastuzumab emtansine<br>for injection, for intravenous<br>use |   | 1,160                             | 18 years    | N/A         | N/A                    | Y               | ¥                               |          | 6/4/2019              |
| Biologicals  | J9355         | Injection, trastuzumab,<br>excludes biosimilar, 10 mg | 10 mg                     | 1/1/2000                  | Herceptin* | trastuzumab for injection, for<br>intravenous use                  | Indicated for:<br>• The treatment of HER2-overexpressing breast cancer.<br>• The treatment of HER2-overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma.<br>Select patients for therapy based on an FDA-approved companion diagnostic for Herceptin.   | 196                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 9/12/2018             |

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| medicaid/medi | caid-ncci-ed  | lit-files   |                            |                         | 1                     |   |   |                                   |             | 1 1         |                        |                 |                                 |          |                       |
|---------------|---------------|---|----------------------------|-------------------------|-----------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description                                       | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name            | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
| Biologicals   | 19356         | Injection, trastuzumab, 10 mg<br>and Hyaluronidase-oysk | 10 mg                      | 7/1/2019                | Herceptin<br>Hylecta™ | trasturumab and<br>hyaluronidase-oysk injection<br>for subcutaneous use | Indicated in adults for the treatment of HER2-overexpressing breast cancer. Select patients for therapy<br>based on an FDA-approved companion diagnostic for trastuzumab.   | 120                               | 18 years    | N/A         | N/A                    | ¥               | ¥                               |          | 6/3/2019              |
| Drugs         | J9357         | Injection, valrubicin,<br>intravesical, 200 mg          | 200 mg                     | 1/1/2000                | Valstar <sup>®</sup>  | valrubicin solution,<br>concentrate, for intravesica<br>use             | Indicated for intravesical therapy of Bacillus Calmette-Guérin (BCG) -refractory carcinoma in situ (CIS) of<br>the urinary bladder in patients for whom immediate cystectomy would be associated with unacceptable<br>morbidity or mortality.   | 20                                | 18 years    | N/A         | N/A                    | Ÿ               | ¥                               |          | 9/12/2018             |
| Biologicals   | J9358         | Injection, fam-trastuzumab<br>deruxtecan-nxki, 1 mg     | 1 mg                       | 7/1/2020                | Enhertu®              | fam-trastuzumab deruxteca<br>nxki for injection, for<br>intravenous use | Indicated for the treatment of:<br>• adult patients with unresectable or metastatic HER2-positive breast cancer who have received a prior<br>anti-HER2-based regimen either:<br>- in the metastatic setting, OR   | 1,800                             | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 5/23/2024             |
| Biologicals   | 19359         | Injection, loncastuximab<br>tesirine-lpyl, 0.075 mg     | 0.075 mg                   | 4/1/2022                | Zynlonta™             | loncastuximab tesirine-lpyl<br>for injection, for intravenou<br>use     |   | 800                               | 18 years    | N/A         | N/A                    | Y               | Y                               |          | 3/17/2022             |
| Drugs         | 19360         | Injection, vinblastine sulfate, 1<br>mg                 | 1 mg                       | 1/1/2009                | N/A                   | vinblastine sulfate injectior   | Indicated in the palliative treatment of the following:<br>Frequently Responsive Malignancies -<br>Generalized Hodgkin's disease (Stages III and IV, Ann Arbor modification of Rye staging system)<br>+ Uymphocytic (hymphoma (and utarian and diffuse, poorly and well differentiated)<br>+ Histocytic lymphoma<br>- Mycosis fungeides (advanced stages)<br>• Advanced carcinoma of the testis<br>• Kaposi's sarcoma<br>+ Letterer-Stwe disease (histiocytosis X)<br>Less Frequently Responsive Malignancies -<br>• Choriocarcinoma resistant to other chemotherapeutic agents<br>• Charcinoma of the breast, unresponsive to appropriate endocrine surgery and hormonal therapy | 250                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 9/12/2018             |

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| medicaid/medi | icaid-ncci-ed | lit-files   |                            |                         |                        |   |   |                                   |             |             |                        |                 |                                 |          |                       |
|---------------|---------------|---|----------------------------|-------------------------|------------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
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| Drugs         | J9370         | Vincristine sulfate, 1 mg   | 1 mg                       | 1/1/2000                | Vincasar PFS®          | vincristine sulfate injection solution                              | Indicated in acute leukemia. Vincasar PFS has also been shown to be useful in combination with other<br>oncolytic agents in Hodgkin's disease, non Hodgkin's malignant lymphomas, rhabdomyosarcoma,<br>neuroblastoma, and Wilms' tumor.   | 20                                | N/A         | N/A         | N/A                    | Y               | Y                               |          | 9/12/2018             |
| Biologicals   | J9376         | Injection, pozelimab-bbfg, 1<br>mg  | 1 mg                       | 4/1/2024                | Veopoz™                | pozelimab-bbfg injection, for<br>intravenous or subcutaneous<br>use |   | 4,000                             | 1 year      | N/A         | N/A                    | Y               | Y                               |          | 4/12/2024             |
| Biologicals   | J9381         | Injection, teplizumab-mzwv,<br>mcg  | 5 5 mcg                    | 7/1/2023                | Tzield™                | teplizumab-mzwv injection,<br>for intravenous use                   | Indicated to delay the onset of Stage 3 type 1 diabetes (TID) in adults and pediatric patients aged 8 years<br>and older with Stage 2 TID.  | 9,600                             | 8 years     | N/A         | N/A                    | Y               | Y                               |          | 6/22/2023             |
| Drugs         | 19390         | Injection, vinorelbine tartrate<br>per 10 mg                                      | , 10 mg                    | 1/1/2000                | Navelbine <sup>®</sup> | vinorelbine tartrate injection,<br>for intravenous use              | Indicated:<br>• In combination with cisplatin for first-line treatment of patients with locally advanced or metastatic non-<br>small cell lung cancer (NSCLC).<br>• As a single agent for first-line treatment of patients with metastatic NSCLC.   | 40                                | 18 years    | N/A         | N/A                    | Y               | ¥                               |          | 9/27/2018             |
| Drugs         | 19393         | Injection, fulvestrant (teva),<br>not therapeutically equivalen<br>to j335, 25 mg | t 25 mg                    | 1/1/2023                | N/A                    | fulvestrant injection, for<br>intramuscular use (Teva)              | Indicated for the treatment of:<br>• Hormone receptor (HRP.)-positive, human epidermal growth factor receptor 2 (HER2)-negative advanced<br>breast cancer in postmenopausal women not previously treated with endocrine therapy.<br>• HRP-positive advanced breast cancer in postmenopausal women with disease progression following<br>endocrine therapy.<br>• HRP-positive, HER2-negative advanced or metastatic breast cancer in postmenopausal women in<br>combination with ridociclib, as initial endocrine based therapy or following disease progression on<br>endocrine therapy.<br>• HRP-positive, HER2-negative advanced or metastatic breast cancer in combination with palbociclib or<br>abemaciclib in women with disease progression after endocrine therapy. | 60                                | 18 years    | N/A         | Females Only           | Y               | Y                               |          | 12/6/2022             |

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|--------------|---------------|--|----------------------------|-------------------------|-----------------------|---|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Drugs        | J9394         | Injection, fulvestrant<br>(fresenus kabi) not<br>therapeutically equivalent to<br>j9395, 25 mg | 25 mg                      | 1/1/2023                | N/A                   | fulvestrant injection, for<br>intramuscular use (Fresenius<br>Kabi) | Monotherapy<br>Fulvestrant Injection is indicated forthe treatment of:<br>+ Hormone receptor(HR)-positive, human epidermal growth factor receptor2 (HER2)-negative advanced<br>breast cancer in postmenopausal women not previously treated with endocrine therapy, or<br>+ HR-positive advanced breast cancer in postmenopausal women with disease progression following<br>endocrine therapy.<br>Combination Therapy<br>Fulvestrant Injection is indicated for the treatment of:<br>+ HR-positive.HR2:negative advanced or metastatic breast cancer in postmenopausal women in<br>combination with ribocible as initial endocrine based therapy or following disease progression on<br>endocrine therapy.<br>+ HR-positive.HR2:negative advanced or metastatic breast cancer in combination with palbociclib or<br>abemaciclib in women with disease progression after endocrine therapy. | 60                                | 18 years    | N/A         | Females Only           | ¥               | ¥                               |          | 12/6/2022             |
| Drugs        | J9395         | Injection, fulvestrant, 25 mg  | 25 mg                      | 1/1/2004                | Faslodex <sup>®</sup> | fulvestrant injection, for<br>intramuscular use                     | Indicated for the treatment of HR-positive advanced breast cancer in postmenopausal women with<br>disease progression following endocrine therapy.<br>Indicated for the treatment of HR-positive, HER2-negative advanced or metastatic breast cancer in<br>combination with palbociclib in women with disease progression after endocrine therapy.<br>Indicated for the treatment of hormone receptor (HR)-positive, human epidermal growth receptor 2<br>(HER2)-negative advanced breast cancer in postmenopausal women not previously treated with endocrine<br>therapy.<br>Indicated for the treatment of HR-positive, HER2-negative advanced or metastatic breast cancer in<br>combination with abemaciclib in women with disease progression after endocrine therapy.  | 60                                | 18 years    | N/A         | Females only           | Y               | Y                               |          | 10/10/2018            |
| Biologicals  | J9400         | injection, ziv-aflibercept, 1 mg   | 1 mg                       | 1/1/2014                | Zaltrap*              | ziv-afilbercept injection for<br>Intravenous infusion               | Indicated in combination with 5-fluorouracil, leucovorin, irinotecan-(FOLFIRI), for the treatment of<br>patients with metastatic colorectal cancer (mCRC) that is resistant to or has progressed following an<br>oxaliplatin-containing regimen.  | 1,800                             | 18 years    | N/A         | N/A                    | Ŷ               | Y                               |          | 6/7/2019              |

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| medicaid/medi |               | dit-files   | 1                          | 1                       | 1          | 1  |   | 1                                 |                                   | T           |                        | 1               | Rebating            |  | 1                     |
|---------------|---------------|---|----------------------------|-------------------------|------------|--|---|-----------------------------------|-----------------------------------|-------------|------------------------|-----------------|---------------------|--|-----------------------|
| Category      | HCPCS<br>Code | HCPCS Description                                 | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age                       | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs         | 009et         | Injection, porfimer sodium, 75<br>mg              | 75 mg                      | 1/1/2000                | Photofrin® | porfimer sodium injection  | Indicated for:<br>Esophageal Cancer<br>- Palliation of patients with completely obstructing esophageal cancer, or of patients with partially<br>obstructing esophageal cancer who, in the opinion of<br>their physician, cannot be satisfactorily treated with Nd:YAG laser therapy<br>Endobranchial Cancer<br>- Treatment of microinvasive endobronchial non-small cell lung cancer (NSCLC) in patients for whom<br>surgery and radiotherapy are not indicated<br>- Reduction of obstruction and palliation of symptoms in patients with completely or partially obstructing<br>endobronchial NSCLC<br>High-Grade Dysplasia in RSCL<br>- Ablation of high-grade dysplasia (HGD) in Barrett's esophagus (BE) patients who do not undergo<br>esophagectomy | 8                                 | 18 years                          | N/A         | N/A                    | Y               | Ŷ                   |  | 6/6/2019              |
| Biologicals   | J9999         | Not otherwise classified,<br>antineoplastic drugs | 1 mcg                      | 1/1/1986                | Anktiva*   |  | Nogapendekin alfa inbakicept-pmln solution is indicated with Bacillus Calmette-Guérin (BCG) for the<br>treatment of adult patients with BCG-unresponsive non-muscle invasive bladder cancer (NMIBC) with<br>carcinoma in situ (CIS) with or without papillary tumors.   | 2,000                             | 18 years                          | N/A         | N/A                    | Ÿ               | Ÿ                   |  | 6/25/2024             |
| Biologicals   | 19999         | Not otherwise classified,<br>antineoplastic drugs | 1 mg                       | 1/1/1986                | imdelltra™ | tarlatamab-dlle for injection,<br>for intravenous use              | Tarlatamab-dlle for injection is indicated for the treatment of adult patients with extensive stage small cell<br>lung cancer (ES-SCLC) with disease progression on or after platinum-based chemotherapy.   | 31                                | 18 years                          | N/A         | N/A                    | Ŷ               | Y                   |  | 6/25/2024             |
| Biologicals   | 19999         | Not otherwise classified,<br>antineoplastic drugs | 1 mcg                      | 1/1/1986                | Besremi®   | ropeginterferon alfa-2b-njft<br>injection, for subcutaneous<br>use | Indicated for the treatment of adults with polycythemia vera.   | 1,500                             | 18 years                          | N/A         | N/A                    | ¥               | Ŷ                   | 1/2024: Procedure code<br>updated from J3590 to J9999<br>to align with product's FDA-<br>approved inclation effective<br>2/1/2024. | 1/26/2024             |
| Biologicals   | P9041         | Infusion, albumin (human),<br>5%, 50 mL           | 50 mL                      | 1/1/2001                | Albutein®  | albumin (human), 5%  | Albutein: Indicated for:<br>+ Hypovolemia<br>- Cardiopulmonary bypass procedures<br>+ Hypoalbuminemia<br>+ Plasma exchange  | 1,550                             | None (use only if clearly needed) | N/A         | N/A                    | Y               | Y                   |  | 5/23/2024             |

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| medicaid/mee | dicaid-ncci-ed | it-files   |                            |                         |   |  |   |                                   |   |             |                        |                 |                                 |   |                       |
|--------------|----------------|--|----------------------------|-------------------------|---|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Category     | HCPCS<br>Code  | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name  | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
| Biologicals  | P9045          | Infusion, albumin (human),<br>5%, 250 mL   | 250 mL                     | 1/1/2002                | Albuked™-5,<br>Albuminex®,<br>AlbuRx®,<br>Albutein®,<br>Flexbumin | albumin (human) U.S.P., 5%<br>solution for injection - 250<br>mL | Albuked-5:<br>Mubked-5: sindicated for:<br>• Emergency treatment of hypovolemic shock<br>• Burn threapy<br>• Cardiopulmonary bypass<br>• Acute liver failure<br>• Sequestration of protein rich fluids  | 620                               | Pediatric Use: Ensure<br>dose is appropriate<br>for body weight. The<br>safety of albumin<br>solutions has been<br>demonstrated in<br>children provided the | N/A         | N/A                    | Y               | Y                               |   | 4/23/2024             |
| Biologicals  | P9046          | Infusion, albumin (human),<br>25%, 20 mL   | 20 mL                      | 1/1/2002                | Albutein®   | albumin (human) U.S.P., 25%<br>solution for injection - 20 mL    | Alburéin 25% is indicated for:<br>+ Hypovolemia<br>< Cardiopulmonary bypass procedures<br>+ Acute nephrosis<br>+ Hypoalburinemia<br>• Ovarian hyperstimulation syndrome<br>+ Neonatal hyperstimulation syndrome<br>Neonatal hyperstimulations   | 775                               | Pediatric Use: No<br>human or animal<br>data. Use only if<br>clearly needed.  | N/A         | N/A                    | Y               | Y                               |   | 4/23/2024             |
| Biologicals  | P9047          | Infusion, albumin (human),<br>25%, 50 mL   | 50 mL                      | 1/1/2002                | Albuked,<br>Albuminar®,<br>Albutein®,<br>Flexbumin,<br>Kedbumin™  | albumin (human), 25%   | Albuket: Indicated for:<br>Emergency treatment of hypovolemic shock<br>Burn therapy<br>Hypoproteinemia with or without edema<br>Adult respiratory distress syndrome (ARDS)<br>Cardiopulmonary bypass<br>- Acute liver failure   | 310                               | Product Specific Age<br>Restrictions<br>(see comments)  | N/A         | N/A                    | Y               | Y                               | Product specific age<br>restrictions:<br>• Kedbumin: 12 years of age<br>and older<br>• Albuked: 18 years of age and<br>older<br>• Albuminar: None | 5/23/2024             |
| Drugs        | Q0138          | Injection, ferumoxytol, for<br>treatment of iron deficiency<br>anemia, 1 mg (non-ESRD use)   | 1 mg                       | 1/1/2010                | Feraheme*   | ferumoxytol injection, for<br>intravenous use (non-ESRD<br>use)  | <ul> <li>Indicated for the treatment of iron deficiency anemia in adult patients with chronic kidney disease<br/>(CKD).</li> <li>Treatment of iron deficiency anemia in adult patients who have intolerance to oral iron or have had<br/>unsatisfactory response to oral iron.</li> </ul>   | 1,020                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 10/26/2018            |
| Drugs        | Q0139          | Injection, ferumoxytol, for<br>treatment of iron deficiency<br>anemia, 1 mg (for ESRD on<br>dialysis)  | 1 mg                       | 1/1/2010                | Feraheme®   | ferumoxytol injection, for<br>intravenous use (ESRD use)         | Indicated for the treatment of iron deficiency anemia in adult patients<br>• With chronic kidney disease (CKD) or<br>• Who have intolerance to oral iron or have had unsatisfactory response to oral iron.  | 1,020                             | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 10/26/2018            |
| Drugs        | Q0144          | Azithromycin dihydrate, oral,<br>capsule/powder, 1 g   | 1 g                        | 1/1/2000                | Zithromax®  | azithromycin, oral   | Approved indication for use in the PADP:<br>4 Sexually Transmitted Diseases<br>Other FDA approved indications:<br>Indicated for the treatment of mild to moderate infections caused by designated, susceptible bacteria:<br>+ Acute bacterial sinusitis in adults<br>+ Acute bacterial sinusitis in adults.   | 2                                 | N/A   | N/A         | N/A                    | Y               | Ŷ                               |   | 6/7/2019              |
| Biologicals  |                | Injection, pemivibart, for the<br>pre-exposure prophylaxis only,<br>for certain adults and<br>adolescents (12 years of age<br>and older weighing at least 40<br>kg) with no known SARS-CoV-2<br>exposure, and who either have<br>moderate-to-severe immune<br>compromise due to a medical<br>condition or receipt of<br>immunosuppressive<br>medications or treatments,<br>and are unlikely to mount an<br>adequate immune response to<br>COVID-19 vaccination, 4500<br>mg | 4500 mg (1 dose)           | 3/22/2024               | Pemgarda  | pemivibart injection, for<br>intravenous use                     | The U.S. FDA has issued an EUA for the emergency use of the unapproved product Pemgarda (pemivibart),<br>a SABS-CoV-2 spike protein-directed attachment inhibitor, for the pre-exposure prophylaxis of coronavirus<br>disease 2019 (COVID-19) in adults and adolecsent [J2 years of age and older weighing at least 40 kgi:<br>• who are not currently infected with SABS-CoV-2 and who have not had a known recent exposure to an<br>individual infected with SABS-CoV-2 and who have not had a known recent exposure to an<br>• who are not currently infected with SABS-CoV-2 and who have not had a known recent exposure to an<br>individual infected with SABS-CoV-2 and who have not had a known recent exposure to an<br>• who have moderate-to-severe immune compromise due to a medical condition or receipt of<br>immunouppressive medications or treatments and are unlikely to mount an adequate immune response<br>to COVID-19 actionation.<br>Pemgarda has been authorized by FDA for the emergency use described above. Pemgarda is not FDA-<br>approved for any use, including use for pre-exposure prophylaxis of COVID-19. | 1                                 | 12 years  | N/A         | N/A                    | Y               | N                               |   | 5/3/2024              |

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| medicaid/medi<br>Category | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name        | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)   | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments | Last Modified<br>Date |
|---------------------------|---------------|---|----------------------------|-------------------------|-------------------|--|--|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------------------|----------|-----------------------|
| Drugs                     | Q2009         | Injection, fosphenytoin, 50 mg<br>phenytoin equivalent  | 50 mg                      | 1/1/2001                | Cerebyx*          | fosphenytoin sodium<br>injection, for intravenous or<br>intramuscular use                | Indicated for the treatment of generalized tonic-clonic status epilepticus and prevention and treatment of<br>seizures occurring during neurosurgery. Cerebyx can also be substituted, as short-term use, for oral<br>phenytoin. Cerebyx should be used only when oral phenytoin administration is not possible.   | 164                               | N/A         | N/A         | N/A                    | Y               | Y                               |          | 3/21/2022             |
| Biologicals               | Q2043         | Sipuleucel-T, minimum of 50<br>million autologous CD54+ cells<br>activated with PAP-GM-CSF,<br>including leukapheresis and all<br>other preparatory procedures,<br>per infusion | 250 mL                     | 7/1/2011                | Provenge®         | sipuleucel-T, suspension for<br>intravenous infusion                                     | Indicated for the treatment of asymptomatic or minimally symptomatic metastatic castrate-resistant<br>(hormone refractory) prostate cancer.  | 3                                 | N/A         | N/A         | Males Only             | Y               | ¥                               |          | 7/16/2018             |
| Drugs                     | Q2050         | Injection, doxorubicin<br>hydrochioride, liposomal, not<br>otherwise specified, 10 mg   | 10 mg                      | 7/1/2013                | Doxil®            | doxorubicin hydrochloride<br>liposome injection, for<br>intravenous use                  | Indicated for:<br>• Ovarian cancer after failure of platinum-based chemotherapy.<br>• AIDS-related Kaposi's Sarcoma after failure of prior systemic chemotherapy or intolerance to such<br>therapy.<br>• Multiple Myeloma in combination with bortezomib in patients who have not previously received<br>bortezomib and have received at least one prior therapy.  | 30                                | 18 years    | N/A         | N/A                    | ¥               | Å                               |          | 6/10/2019             |
| Biologicals               | Q4081         | Injection, epoetin alfa, 100<br>units (for ESRD on dialysis) (for<br>renal dialysis facilities and<br>hospital use)   | 100 units                  | 1/1/2007                | Epogen®, Procrit® | epoetin alfa injection, for<br>intravenous or subcutaneous<br>use (for ESRD on dialysis) | Indicated for treatment of anemia due to<br>- Chronic Kidney Disease (CKD) in patients on dialysis and not on dialysis.<br>- Zidovudine in patients with HIV-infection.<br>The effects of concomitant myelosuppressive chemotherapy, and upon initiation, there is a minimum of<br>two additional months of planned chemotherapy.<br>• Reduction of allogeneic RBC transfusions in patients undergoing elective, noncardiac, nonvascular<br>surgery. | 1,960                             | 1 month     | N/A         | N/A                    | Y               | Y                               |          | 1/12/2022             |

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| Category    | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modifie<br>Date |
|-------------|---------------|--|----------------------------|-------------------------|------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|----------------------|
| Biologicals | Q5101         | Injection, filgrastim-sndz,<br>biosimilar, (Zarxio), 1<br>microgram                          | 1 mcg                      | 4/1/2018                | Zarxio*    | fligrastim-sndz injection, for<br>subcutaneous or intravenous<br>use                            | Indicated to:<br>• Decrease the incidence of infection, as manifested by febrile neutropenia, in patients with nonmyeloid<br>malignancies receiving myelosuppressive anticancer drugs associated with a significant incidence of sever<br>neutropenia with feve.<br>• Reduce the time to neutrophil recovery and the duration of fever, following induction or consolidation<br>chemotherapy treatment of patients with acute myeloid leukemia (AML).<br>• Reduce the duration of neutropenia and neutropenia-related clinicalsequele, e.g., febrile neutropenia,<br>in patients with nonmyeloid malignancies undergoing myeloslative chemotherapy followed by bone<br>marrow transplantation (BMT).<br>• Mobilize autologous hematopoietic progenitor cells into the peripheral blood for collection by<br>leukapheresis.<br>• Reduce the incidence and duration of sequelae of severe neutropenia (e.g., fever, infections,<br>oropharyngeal ulcers) in symptomatic patients with congenital neutropenia, cyclic neutropenia, or<br>idiopathic neutropenia.   | 59,520                            | N/A   | N/A         | N/A                    | Y               | Y                               |   | 6/6/2019             |
| Biologicals | Q5103         | Injection, infliximab-dyyb,<br>biosimilar, (Inflectra), 10 mg                                | 10 mg                      | 4/1/2018                | Inflectra® | infliximab-dyyb lyophilized<br>concentrate for injection, for<br>intravenous use                | Indicated for:<br>Crohn's Disease:<br>• reducing signs and symptoms and inducing and maintaining clinical remission in adult patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.  | 300                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>Crohn's Disease and Ulcerative<br>Colitis: 6 years of age and   | 7/29/2024            |
| Biologicals | Q5104         | Injection, infliximab-abda,<br>biosimilar, (Renflexis), 10 mg                                | 10 mg                      | 4/1/2018                | Renflexis® | infliximab-abda for injection,<br>for intravenous use   | Indicated for:<br>Crohr's Disease:<br>- Reducing signs and symptoms and inducing and maintaining clinical remission in adult patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.<br>- Reducing the number of draining netrococtaneous and rectorograph fistulas and maintaining fistula<br>- Reducing the number of draining netrococtaneous and rectorograph fistulas and maintaining fistula  | 300                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>Crohn's Disease: 6 years and<br>older<br>Ulcerative Collist: 6 years<br>and older<br>Rheumatoid Arthritis in<br>combination with<br>methotrexate: 18 years and<br>older<br>• Ankytosing Spondylits: 18<br>years and older | 7/29/2024            |
| Biologicals | Q5105         | Injection, epoetin alfa-epbx,<br>biosimilar, (retacrit) (for esrd<br>on dialysis), 100 units | 100 units                  | 7/1/2018                | Retacrit™  | epoetin alfa-epbx injection,<br>for intravenous or<br>subcutaneous use for ESRD<br>on dialysis) | Indicated for the treatment of anemia due to: O Gronic kidney disease (XD) in patients on dialysis and not on dialysis. O Zdoudne in patients with HIV-infection. O The effects of concomitant myelosuppressive chemotherapy, and upon initiation, there is a minimum of two additional months of planned chemotherapy. Indicated for the reduction of allogeneic RRC transfusions in patients undergoing elective, noncardiac, nonvascular surgery. Limitations of Use: Retarcit has not been shown to improve quality of life, fatigue, or patient well-being. Not indicated for use in: In patients with cancer receiving hormonal agents, biologic products, or radiotherapy, unless also receiving concomitant myelosuppressive chemotherapy. In patients with cancer receiving myelosuppressive chemotherapy when the anticipated outcome is cure. In patients with cancer receiving myelosuppressive chemotherapy in whom the anemia can be managed by transfusion. In patients undergoing cardiac vascular surgery. In patients undergoing cardiac vascular surgery. As a substitute for RBC transfusions in patients who require immediate correction of anemia. | 1,960                             | 1 month   | N/A         | N/A                    | Y               | Y                               |   | 1/12/2022            |
| Biologicals | Q5106         | Injection, epoetin alfa-epbx,<br>biosimilar, (retacrit) (for non-<br>esrd use), 1000 units   | 1,000 units                | 7/1/2018                | Retacrit™  | epoetin alfa-epbx injection,<br>for intravenous or<br>subcutaneous use (for non-<br>ESRD use)   | Indicated for the treatment of anemia due to:<br>o Chronic kidney disease (CKD) in patients on dialysis and not on dialysis.<br>o Zidovudine in patients with HIV:Infection.<br>o The effects of concomitant myelosuppressive chemotherapy, and upon initiation, there is a minimum of<br>two additional months of planned chemotherapy.<br>Indicated for the reduction of allogeneic RBC transfusions in patients undergoing elective, noncardiac,<br>nonvascular surgery.<br>Umitations of Use: Retacrit has not been shown to improve quality of life, fatigue, or patient well-being.   | 630                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Y                               | Indication specific age<br>restrictions:<br>• CKD not on dialysis: 1 month<br>of age and older<br>• Anemia due to concomitant<br>myelosuppressive<br>chemotherapy: 5 years of age<br>and older<br>• Zidovudine-treated, anemia,<br>patients with HV infection: 8      | 1/12/2022            |
| Biologicals | Q5107         | Injection, bevacizumab,<br>(mvasi), 10 mg  | 10 mg                      | 1/1/2019                | Mvasi™     | bevacizumab-awwb injection<br>for intravenous use   | Indicated for the treatment of:<br>• Metastatic colorectal cancer, in combination with intravenous fluorouracil-based chemotherapy for first-<br>or scoond-line treatment.<br>• Metastatic colorectal cancer, in combination with fluoropyrimidine-irinotecan- or fluoropyrimidine-<br>onalipatir-based chemotherapy for second-line treatment in patients who have progressed on a first-line<br>bevacizmab product-containing regimen.  | 420                               | 18 years  | N/A         | N/A                    | Y               | Y                               | process with my meetion. o  | 7/20/2022            |

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| Visio         Visio         Vision         Vision </th <th>Category</th> <th>aid-ncci-ed</th> <th>HCPCS Description</th> <th>HCPCS Code Billing</th> <th></th> <th>Brand Name</th> <th>Generic Name</th> <th>FDA Approved Indications</th> <th>NC Suggested Max</th> <th>Minimum Age</th> <th>Maximum Age</th> <th>Gender</th> <th>NDC</th> <th>Rebating<br/>Labeler</th> <th>Comments</th> <th>Last Modified</th>  | Category    | aid-ncci-ed | HCPCS Description  | HCPCS Code Billing |                | Brand Name | Generic Name                | FDA Approved Indications   | NC Suggested Max | Minimum Age | Maximum Age | Gender       | NDC      | Rebating<br>Labeler | Comments | Last Modified |
|--|-------------|-------------|--|--------------------|----------------|------------|-----------------------------|--|------------------|-------------|-------------|--------------|----------|---------------------|----------|---------------|
| No.         Participant Performance Science         Participant Performance Science  | Category    | Code        | HCPC3 Description  | Unit               | Effective Date | Brand Name | Generic Name                | (See Package Insert for full FDA approved indication descriptions)   | Monthly Units    | winimum Age | waximum Age | Restrictions | Required |                     | Comments | Date          |
| bit  | Biologicals | Q5108       |  | 0.5 mg             | 10/1/2018      | Fulphila** |                             | myeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically significant<br>incidence of febrile neutropenia.<br>Limitations of Use:<br>Liphihia is not indicated for the mobilization of peripheral blood progenitor cells for hematopoletic stem   |                  | N/A         | N/A         | N/A          | ¥        | Y                   |          | 3/21/2023     |
| No.11       Nighteds, programments, objection, programments, objection, section, and section in programments and cover programment and cover programeter programeter and cover programment and cover prog  | Biologicals | Q5110       | biosimilar, (Nivestym), 1                                  | 1 mcg              | 10/1/2018      | Nivestym™  | subcutaneous or intravenous | <ul> <li>Decrease the incidence of infection, as manifested by febrile neutropenia, in patients with nonmyeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a significant incidence of severe neutropenia with fever.</li> <li>Reduce the time to neutrophil recovery and the duration of fever, following induction or consolidation chemotherapy treatment of patients with acute myeloid leukemia (AML).</li> </ul> | 59,520           | N/A         | N/A         | N/A          | Y        | Y                   |          | 12/28/2018    |
| 9 S12       bpc/clon, Trastlaurnab-dth, 10 mg       10 mg       71/2019       Ontuant       instalurunab dth, 0 miller, 0 for intravenous up       The trastment of HE2 overcepressing metatatic gastric or gastroesophaged junction adencacriom       136       18 years       N/A       N/A       V       V       V       (512)       (512)         Biological biolitar, (berrunab-bith), 10 mg       10 mg       71/2019       herum*       millected for:<br>intertement of HE2 overcepressing metatatic gastric or gastroesophaged junction adencacriom       136       18 years       N/A       N/A       V       V       V       (22)       (22)         Biological biolitar, (berrunab-bith), 10 mg       10 mg       71/2019       Herum*       indicated for:<br>instrume dthE2 overcepressing metatatic gastric or gastroesophaged junction adencacriom       136       18 years       N/A       N/A       V       V       V       (22)       (22)         Biological biolitar, (berrunab-bith), 10 mg       10 mg       71/2019       Levin       Indicated for:<br>instrume dthE2 overcepressing metatatic gastric or gastroesophaged junction adencacriom       136       18 years       N/A       N/A       V       V       V       V       V       V       V       V       V       V       V       V       V       V       V       V       V       V <t< td=""><td>Biologicals</td><td>Q5111</td><td></td><td>0.5 mg</td><td>1/1/2019</td><td>Udenyca*</td><td></td><td>non-myeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically<br/>significant incidence of febrile neutropenia.</td><td>36</td><td>N/A</td><td>N/A</td><td>N/A</td><td>Y</td><td>Y</td><td></td><td>5/23/2024</td></t<>   | Biologicals | Q5111       |  | 0.5 mg             | 1/1/2019       | Udenyca*   |                             | non-myeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically<br>significant incidence of febrile neutropenia.  | 36               | N/A         | N/A         | N/A          | Y        | Y                   |          | 5/23/2024     |
| $10 \text{ gs} 13$ $\frac{10 \text{ gs}}{10 \text{ gs}} \frac{10 \text{ gs}}{$ | Biologicals | Q5112       |  | 10 mg              | 7/1/2019       | Ontruzant® |                             | <ul> <li>The treatment of HER2-overexpressing breast cancer.</li> </ul>  | 196              | 18 years    | N/A         | N/A          | Y        | Y                   |          | 5/25/2020     |
| Biologicals       Q5114       Injection, Trasturumab-dikt, (Ogivri), 10 mg       10 mg       7/1/2019       Ogivri <sup>w</sup> resturanab-kist for intravenous use       *The treatment of HER2-overexpressing meast cancer.       196       18 years       N/A       N/A       Y       Y       Y       P       12/4/2019         Biologicals       0.5114       Injection, frusturanab-dikt, (Ogivri), 10 mg       10 mg       7/1/2019       Ogivri <sup>w</sup> resturanab-dikt for intravenous use       *The treatment of HER2-overexpressing meastatic gastric or gastroesophageal junction adenocarcinoma.       196       18 years       N/A       N/A       Y       Y       Y       P       P       12/4/2019         Injection, rituximab-abbs,       10 mg       7/1/2019       Ogivri <sup>w</sup> rituinab-abbs injection, for intravenous use       *Interestment of AUE patients with:       root       18 years       N/A       N/A       Y       Y       P       P       12/4/2019         Ninesteine       0.011       Injection, rituximab-abbs,       10 mg       7/1/2019       rituinab-abbs injection, for       Indicated for the treatment of adult patients with:       root       18 years       N/A       N/A       Y       Y       P       12/4/2019   | Biologicals | Q5113       |  | 10 mg              | 7/1/2019       | Herzuma®   |                             | the treatment of HER2-overexpressing breast cancer.     the treatment of HER2-overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma.   | 196              | 18 years    | N/A         | N/A          | Y        | ¥                   |          | 4/29/2020     |
|  | Biologicals | Q5114       |  | 10 mg              | 7/1/2019       | Ogivri*    |                             | The treatment of HER2-overexpressing breast cancer.     The treatment of HER2-overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma.   | 196              | 18 years    | N/A         | N/A          | ¥        | ¥                   |          | 12/4/2019     |
| biosimilar, (Truzima), 10 mg 1/1/2015 100mm intravenous use • Non-Hodgkin's Lymphoma (NHL) 300 10 years 10/1 10/1 1 1  | Biologicals | Q5115       | Injection, rituximab-abbs,<br>biosimilar, (Truxima), 10 mg | 10 mg              | 7/1/2019       | Truxima®   |                             |  | 500              | 18 years    | N/A         | N/A          | Y        | Y                   |          | 12/4/2019     |

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| medicaid/med | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments  | Last Modified<br>Date |
|--------------|---------------|--|----------------------------|-------------------------|------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|---|-----------------------|
| Biologicals  | Q5116         | Injection, trastuzumab-gyyp,<br>biosimilar, (trazimera), 10 mg   | 10 mg                      | 10/1/2019               | Trazimera™ | trastuzumab-qyyp for<br>injection, for intravenous use | Indicated for:<br>• The treatment of HER2-overexpressing breast cancer.<br>• The treatment of HER2-overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma.   | 196                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 3/26/2020             |
| Biologicals  | Q5117         | Injection, trastuzumab-anns,<br>biosimilar, (kanjinti), 10 mg    | 10 mg                      | 10/1/2019               | Kanjinti™  | trastuzumab-anns for<br>injection, for intravenous use | Indicated for:<br>• The treatment of HER2 overexpressing breast cancer.<br>• The treatment of HER2 overexpressing metastatic gastric or gastroesophageal junction adenocarcinoma.<br>Select patients for therapy based on an FDA-approved companion diagnostic for a trastuzumab product.   | 196                               | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |   | 12/14/2021            |
| Biologicals  | Q5118         | Injection, bevacizumab-bvzr,<br>biosimilar, (Zirabev), 10 mg     | 10 mg                      | 10/1/2019               | Zirabev™   | bevacizumab-bvzr injection,<br>for intravenous use     | Indicated for the treatment of:<br>• Metastatic colorectal cancer, in combination with intravenous fluorouracil-based chemotherapy for first<br>or second-line treatment.<br>• Metastatic colorectal cancer, in combination with fluoropyrimidine-irinotecan- or fluoropyrimidine-<br>oxaliplatin-based chemotherapy for second-line treatment in patients who have progressed on a first-line<br>bevacicumab product-containing regimen.   | 420                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 7/20/2022             |
| Biologicals  | Q5119         | Injection, rituximab-pvvr,<br>biosimilar, (ruxience), 10 mg      | 10 mg                      | 7/1/2020                | Ruxience™  | rituximab-powr injection, for<br>intravenous use       | Indicated for the treatment of adult patients with:<br>• Non-Hodgkin's Lymphoma (NHL):<br>o Relaped or refractory, low grade or folicular, CD20-positive B-cell NHL as a single agent.<br>o Previously untreated folicular, CD20-positive, B-cell NHL in combination with first line chemotherapy<br>and, in patients achieving a complete or partial response to a rituximab product in combination with<br>chemotherapy, as single-agent maintenance therapy.<br>o Non-progressing (including stable disease), low-grade, CD20-positive, B-cell NHL as a single agent after<br>first-line cyclophosphamide, vincristine, and prednisone (CVP) chemotherapy.<br>o Previously untreated diffuse large B-cell, CD20-positive NHL in combination with (cyclophosphamide,<br>doxorubicin, vincristine, and prednisone) (CHDP) or other anthracycline-based chemotherapy regimens.<br>- Chronic Lymphorytic Leukemia (CL1):<br>o Previously untreated and previously treated CD20-positive CLL in combination with fludarabine and<br>cyclophosphamide (FC).<br>in adult patients in combination with glucocorticoids.<br>severely-active RA who have inadequate response to one or more TNF antagonist therapies.  | 500                               | 18 years  | N/A         | N/A                    | Y               | Y                               |   | 12/16/2021            |
| Biologicals  | Q5120         | Injection, pegfilgrastim-bmez<br>(ziextenzo), biosimilar, 0.5 mg |                            | 7/1/2020                | Ziextenzo™ | pegfilgrastim-bmez injection,<br>for subcutaneous use  | Indicated to:<br>• decrease the incidence of infection, as manifested by febrile neutropenia, in patients with nonmyeloid   | 36                                | N/A   | N/A         | N/A                    | Y               | Y                               |   | 3/22/2024             |
| Biologicals  | Q5121         | Injection, infloimab-axxq,<br>biosimilar, (avsola), 10 mg        | 10 mg                      | 7/1/2020                | Avsola**   |  | malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically significant<br>Indicated for:<br>Indicate for:<br>ereducing signs and symptoms and inducing and maintaining clinical remission in adult patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.<br>• reducing signs and symptoms and inducing and maintaining clinical remission in pediatric patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.<br>• reducing signs and symptoms and inducing and maintaining clinical remission in pediatric patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.<br>• reducing signs and symptoms, inducing and maintaining clinical remission and mucosal healing, and<br>an inadequate response to conventional therapy.<br>Pediatric Cortons of the severely active disease who have had an inadequate response to conventional therapy.<br>• Reducing signs and symptoms, inducing and maintaining clinical remission in pediatric patients with<br>moderately to severely active disease who have had an inadequate response to conventional therapy.<br>• Pediatric Ulcerative Colitis:<br>• reducing signs and symptoms and inducing and maintaining clinical remission in pediatric patients with<br>moderately to severely active disease.<br>• Nanylosing Spondytilis:<br>• reducing signs and symptoms in patients with active disease.<br>• Boriatic Arthritis:<br>• reducing signs and symptoms in patients with active disease.<br>• Soriatic Arthritis:<br>• reducing signs and symptoms of active arthritis, inhibiting the progression of structural damage, and<br>improving physical function.<br>• Plaque Fornical<br>• readure signs and symptoms of active arthritis, inhibiting the progression of structural damage, and<br>improving physical function.<br>• Plaque Fornical | 300                               | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | ¥                               | Indication specific age<br>restrictions:<br>Crohr's disease and ulcerate<br>RA, ankylosing spondylits,<br>psoriatic arthritis and plaque<br>psoriasis: 18 years of age and<br>older<br>5/2024: NC Suggested Max<br>Monthly Units updated to align<br>with MUE values effective<br>5/6/2024. |                       |

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|             | icaid-ncci-edi |  | HCPCS Code Billing         | HCPCS                   |            |  | FDA Annual Indiantiana  | NO Commented                      |             |             | Gender                 | NDC             | Rebating            |          | Last Modified         |
|-------------|----------------|--|----------------------------|-------------------------|------------|--|---|-----------------------------------|-------------|-------------|------------------------|-----------------|---------------------|----------|-----------------------|
| Category    | Code           | HCPCS Description  | HCPCS Code Billing<br>Unit | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Labeler<br>Required | Comments | Last Modified<br>Date |
| Biologicals | Q5122          | Injection, pegfilgrastim-apgf<br>(nyvepria), biosimilar, 0.5 mg      | 0.5 mg                     | 1/1/2021                | Nyvepria™  | pegfilgrastim-apgf injection,<br>for subcutaneous use                | Indicated to decrease the incidence of infection, as manifested by febrile neutropenia, in patients with non<br>myeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically significant<br>incidence of febrile neutropenia.<br>Umitations of Use:<br>Nyvepria is not indicated for the mobilization of peripheral blood progenitor cells for hematopoietic stem<br>cell transplantation.  | 36                                | N/A         | N/A         | N/A                    | Ŷ               | Y                   |          | 3/21/2023             |
| Biologicals | Q5123          | Injection, rituximab-arrx,<br>biosimilar, (riabni), 10 mg            | 10 mg                      | 7/1/2021                | Riabni™    | rituximab-anx injection, for<br>intravenous use                      | Indicated for the treatment of:<br>• Adult patients with non-Hodgkin's Lymphoma (NHL).<br>• Relapsed or refractory, low grade or follicular, CD20-positive B-cell NHL as a single agent.<br>• O Previously untreated follicular, CD20-positive, B-cell NHL in combination with first line chemotherapy<br>and, in patients achieving a complete or partial response to a rituzimab product in combination with<br>chemotherapy, as single-agent maintenance therapy.<br>• Non-progressing (including stable disease), low-grade, CD20-positive, B-cell NHL as a single agent after<br>first-line cyclophosphamide, vincristine, and prednisone (CVP) chemotherapy.<br>• Previously untreated diffue large B-cell, CD20-positive NHL in combination with cyclophosphamide,<br>doxorubicin, vincristine, and prednisone (CVP) or other antracycline-based chemotherapy regimens.<br>• Adult patients with Chronic Lymphocytic Leukemia (CLL).<br>• Oreviously untreated and previously treated CD20-positive CLL in combination with fludarabine and<br>cyclophosphamide (FC).<br>• Granulomatosis with Polyanglitis (GPA) (Wegener's Granulomatosis) and Microscopic Polyangitis (MPA).  | 500                               | 18 years    | N/A         | N/A                    | Y               | Y                   |          | 7/20/2022             |
| Biologicals | Q5124          | Injection, ranibizumab-nuna,<br>biosimilar, (byooviz), 0.1 mg        | 0.1 mg                     | 4/1/2022                | Byooviz™   | ranibizumab-nuna injection,<br>for intravitreal use                  | Indicated for the treatment of patients with:<br>- Neovascular (Wet) Age-Related Macular Degeneration (AMD)<br>- Macular Edema Following Reinal Vein Occlusion (RVD)<br>- Myopic Choroidal Neovascularization (mCNV)  | 20                                | 18 years    | N/A         | N/A                    | Y               | Y                   |          | 6/20/2022             |
| Biologicals | Q5125          | Injection, filgrastim-ayow,<br>biosimilar, (releuko), 1<br>microgram | 1 mcg                      | 10/1/2022               | Releuko*   | filgrastim-ayow injection, for<br>subcutaneous or intravenous<br>use |   | 59,520                            | N/A         | N/A         | N/A                    | Y               | Y                   |          | 9/15/2022             |
| Biologicals | Q5126          | Injection, bevacizumab-maly,<br>biosimilar, (alymsys), 10 mg         | 10 mg                      | 1/1/2023                | Alymsys®   | bevacizumab-maly injection,<br>for intravenous use                   | Indicated for the treatment of:<br>• Metastatic colorectal cancer, in combination with intravenous fluorouracii-based chemotherapy for first<br>or second-line treatment.<br>• Metastatic colorectal cancer, in combination with fluoropyrimidine-innotecan- or fluoropyrimidine-<br>oralipatitr-based chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment in patients who have progressed chemotherapy for second-line treatment of colon cancer.<br>• Unresectable, locally advanced, recurrent or metastatic non-squamous non-small cell lung cancer, in combination with carboplatin and patitixas for first-line treatment.<br>• Recurrent glioblastoma in adults.<br>• Metastatic recurrent, or metastatic cervical cancer, in combination with paclitaxel and cisplatin, or paclitaxel and topotecan.<br>• Epithelial oxoriacin, faliopian tube, or primary peritoneal cancer in combination with paclitaxel, pegylated liposomal downbicin, or topotecan for platinum-resistant recurrent disease who received no more than 2 prior chemotherapy regimens. | 420                               | 18 years    | N/A         | N/A                    | Y               | Y                   |          | 12/12/2022            |
| Biologicals | Q5127          | Injection, pegfilgrastim-fpgk<br>(stimufend), biosimilar, 0.5 mg     | 0.5 mg                     | 4/1/2023                | Stimufend® | pegfilgrastim-fpgk injection,<br>for subcutaneous use                | In combination with atexolizumab for the treatment of patients with unresectable or metastatic<br>hepatocellular carcinoma (HCC) who have not received prior systemic therapy.<br>Indicated to: Decrease the incidence of infection, as manifested by febrile neutropenia, in patients with non-myeloid<br>malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically significant<br>incidence of febrile neutropenia.<br>Increase survival in patients acutely exposed to myelosuppressive doses of radiation (Hematopoietic<br>Subsyndrome of Acute Radiation Syndrome).<br>Limitations of Use<br>Stimufend is not indicated for the mobilization of peripheral blood progenitor cells for hematopoietic stem   | 36                                | N/A         | N/A         | N/A                    | Y               | Y                   |          | 10/26/2023            |

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| Category    | HCPCS<br>Code | HCPCS Description   | HCPCS Code Billing<br>Unit   | HCPCS<br>Effective Date | Brand Name | Generic Name   | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
|-------------|---------------|---|------------------------------|-------------------------|------------|--|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Biologicals | Q5128         | Injection, ranibizumab-eqrn<br>(cimerli), biosimilar, 0.1 mg                              | 0.1 mg                       | 4/1/2023                | Cimerli™   | ranibizumab-eqm injection,<br>for intravitreal use   | Indicated for the treatment of patients with:<br>- Neovascular (Wet) Age-Related Macular Degeneration (AMD)<br>- Macular Edema (IONE)<br>- Diabetic Macular Edema (IONE)<br>- Diabetic Retinopathy (DR)<br>- Myopic Choroidal Neovascularization (mCNV)   | 20                                | 18 years  | N/A         | N/A                    | ¥               | Y                               |  | 3/16/2023             |
| Biologicals | Q5129         | Injection, bevacizumab-adcd<br>(vegzelma), biosimilar, 10 mg                              | 10 mg                        | 4/1/2023                | Vegzelma*  | bevacizumab-adcd injection,<br>for intravenous use   | sacificated and topotecan.<br>• Epithelial ovarian, fallopian tube, or primary peritoneal cancer:<br>o in combination with achoplatin and pacitaxel, followed by Vegzelma as a single agent, for stage III or IV<br>disease following initial surgical resection<br>o in combination with pacificaxel, pegylated liposomal doxerubicin, or topotecan for platinum-resistant<br>recurrent disease who received no more than 2 prior themotherapy regimens<br>o in combination with carboplatin and pacificatel or carboplatin and gemcitabine, followed by Vegzelma as<br>a single agent, for platinum-sensitive recurrent disease<br>**Added at Request of the State Per NCCN Guidelines:<br>• in combination with ateriobizumab for the treatment of patients with unresectable or metastatic  | 420                               | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 5/25/2023             |
| Biologicals | Q5130         | Injection, pegfilgrastim-pbbk<br>(fylnetra), biosimilar, 0.5 mg                           | 0.5 mg                       | 4/1/2023                | Fylnetra®  | pegfilgrastim-pbbk injection,<br>for subcutaneous use  | hepatocellular carcinoma (HCC) who have not received prior systemic therapy.<br>Indicated to decrease the incidence of infection, as manifested by febrile neutropenia, in patients with<br>nonmyeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically<br>significant incidence of febrile neutropenia.<br>Limitations of Use:<br>Fynetra is not indicated for the mobilization of peripheral blood progenitor cells for hematopoietic stem<br>cell transplantation.   | 36                                | N/A   | N/A         | N/A                    | Y               | Y                               |  | 5/25/2023             |
| Biologicals | Q5133         | Injection, tocilizumab-bavi<br>(tofidence), biosimilar, 1 mg                              | 1 mg                         | 4/1/2024                | Tofidence™ | tocilizumab-bavi injection, fo<br>intravenous use  | Variability and a service of the service of th | 1,600                             | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | Y               | Ŷ                               | RA: 18 years of age and older<br>PJIA, SJIA: 2 years of age and<br>older | 5/23/2024             |
| Drugs       | Q9991         | Injection, buprenorphine<br>extended-release (Sublocade),<br>less than or equal to 100 mg | less than or equal to 100 mg | 7/1/2018                | Sublocade™ | buprenorphine extended-<br>release injection, for<br>subcutaneous use, less than<br>or equal to 100 mg | Indicated for the treatment of moderate to severe opioid use disorder in patients who have initiated  | 2                                 | 18 years  | N/A         | N/A                    | Y               | Ŷ                               |  | 9/27/2018             |
| Drugs       | Q9992         | Injection, buprenorphine<br>extended-release (Sublocade),<br>greater than 100 mg          | greater than 100 mg          | 7/1/2018                | Sublocade™ | buprenorphine extended-<br>release injection, for<br>subcutaneous use, greater<br>than 100 mg          | Indicated for the treatment of moderate to severe opioid use disorder in patients who have initiated<br>treatment with a transmucosal buprenorphine-containing product, followed by dose adjustment for a<br>minimum of 7 days.   | 2                                 | 18 years  | N/A         | N/A                    | Y               | Y                               |  | 9/27/2018             |

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| medicaid/medicaid-ncci-edit-files |               |  |   |                         |             |   |   |                                   |   |             |                        |                 |                                 |  |                       |
|-----------------------------------|---------------|--|---|-------------------------|-------------|---|---|-----------------------------------|---|-------------|------------------------|-----------------|---------------------------------|--|-----------------------|
| Category                          | HCPCS<br>Code | HCPCS Description  | HCPCS Code Billing<br>Unit  | HCPCS<br>Effective Date | Brand Name  | Generic Name  | FDA Approved Indications<br>(See Package Insert for full FDA approved indication descriptions)  | NC Suggested Max<br>Monthly Units | Minimum Age   | Maximum Age | Gender<br>Restrictions | NDC<br>Required | Rebating<br>Labeler<br>Required | Comments   | Last Modified<br>Date |
| Drugs                             | 50013         | Esketamine, nasal spray, 1 mg                              | 1 mg  | 1/1/2021                | Spravato™   | esketamine nasal spray                                      | <ul> <li>Indicated, in conjunction with an oral antidepressant, for the treatment of treatment-resistant<br/>depression (TRD) in adults.</li> <li>Indicated for depressive symptoms in adults with major depressive disorder (MDD) with acute suicidal<br/>ideation or behavior.</li> <li>Limitations of Use: Spravato is not approved as an anesthetic agent. The safety and effectiveness of</li> </ul>   | 728                               | 18 years  | N/A         | N/A                    | Ŷ               | Y                               |  | 12/28/2020            |
|                                   |               |  |   |                         |             |   | Spravato as an anesthetic agent have not been established.  |                                   |   |             |                        |                 |                                 |  |                       |
| Drugs                             | 50028         | Injection, famotidine, 20 mg                               | 20 mg   | 1/1/2000                | Pepcid*     | famotidine injection  | Indicated in some hospitalized patients with pathological hypersecretory conditions or intractable ulcers,<br>or as an alternative to the oral dosage forms for short term use in patients who are unable to take oral<br>medication for the following conditions:<br>1. Short term treatment of active duodenal ulcer. Most adult patients heal within 4 weeks; there is rarely<br>reason to use famoltima et ill duodage for longer than to 8 a weeks. Xudies have not assessed the safety<br>of famotidine in uncomplicated active duodenal ulcer for periods of more than eight weeks.<br>2. Maintenance therary for duodenal ulcer patients are teduced dosage after healing of an active ulcer.<br>Controlled studies in adults have not extended beyond one year.<br>3. Short term treatment of active duoden generation and the safety of famotidine in uncomplicated active beneign gastric ulcer. Most adult patients heal within 6 weeks. Studies<br>have not assessed the safety or efficacy of famotidime in uncomplicated active beneign gastric ulcer for<br>periods of more than 8 weeks.<br>4. Short term treatment of gastroesophageal reflux disease (GERD). Famotidine is indicated for short term<br>treatment of patients with symptoms of GERD.<br>5. Fraentotine is also indicated for the short term treatment of esophagitis due to GERD including erosive<br>or ulcerative disease diagnosed by endoscopy.<br>6. Treatment of pathological hypersecretory conditions (e.g., Zollinger-Ellison Syndrome, multiple<br>endocrine adenomas). | 62                                | 1 year  | N/A         | N/A                    | ¥               | ¥                               | 11/2020 Coverage effective<br>1//2019 per DHB request<br>11/2023 Permanent code<br>S0028 effective 12/1/2023 per<br>DHB request  | 11/10/2023            |
| Drugs                             | S0080         | Injection, pentamidine<br>isethionate, 300 mg              | 300 mg  | 1/1/2000                | Pentam® 300 | pentamidine isethionate for<br>injection                    | Indicated for the treatment and prevention of pneumonia caused by Pneumocystis carinii.   | 42                                | 4 months  | N/A         | N/A                    | Y               | Y                               |  | 8/24/2018             |
| Biologicals                       | 50145         | Injection, pegylated interferon<br>alfa-2a, 180 mcg per mL | 180 mcg   | 7/1/2005                | Pegasys®    | peginterferon alfa-2a<br>injection, for subcutaneous<br>use | Chronic Hepatitis C (CHC):<br>•Adult Patients: In combination therapy with other hepatitis C virus drugs for adults with compensated<br>liver disease. Pegasys monotherapy is indicated only if patient has contraindication or significant<br>intolerance to other HCV drugs.<br>•Pediatric Patients: In combination with ribavirin for pediatric patients 5 years of age and older with<br>compensated liver disease.<br>Chronic Hepatitis B (CHB):<br>•Adult Patients: Treatment of adults with HBeAg-positive and HBeAg-negative chronic hepatitis B (CHB)<br>infection who have compensated liver disease and evidence of viral replication and liver inflammation.<br>•Pediatric Patients: Treatment of non-cirrhotic pediatric patients 3 years of age and older with HBeAg-<br>positive CHB and evidence of viral replication and elevations in serum alanine aminotransferase (ALT).   | 5                                 | Indication Specific<br>Age Restrictions<br>(see comments) | N/A         | N/A                    | ¥               | Y                               | Indication specific age<br>restrictions:<br>• Chronic Hepatitis C: 5 years<br>of age and older<br>• Chronic Hepatitis B: 3 years<br>of age and older   | 7/2/2018              |
| Drugs                             | S0189         | Testosterone pellet, 75 mg                                 | 75 mg   | 1/1/2002                | Testopel®   | testosterone pellets for subcutaneous implantation          | Indicated for replacement therapy in conditions associated with a deficiency or absence of endogenous<br>testosterone:<br>Primary hypogonadism (congenital or acquired) - testicular failure due to cryptorchidism, bilateral<br>torsion, orchitis, vanishing testes syndrome; or orchiectomy.<br>Hypogonadotropic hypogonadism (congenital or acquired) - gonadotropic LHRH deficiency, or pituitary -<br>hypothalamic injury from tumors; runum ao radiation.   | 6                                 | N/A   | N/A         | Males Only             | Y               | Y                               |  | 9/21/2018             |
| Drugs                             | S0190         | Mifepristone, oral, 200 mg                                 | 200 mg  | 1/1/2000                | Mifeprex*   | mifepristone tablets, for oral<br>use                       | Indicated, in a regimen with misoprostol, for the medical termination of intrauterine pregnancy through 70 days gestation.  | 1                                 | N/A   | N/A         | Females Only           | Y               | Y                               |  | 3/15/2019             |
| Drugs                             | S0191         | Misoprostol, oral, 200 mcg                                 | 200 mcg   | 1/1/2000                | Cytotec®    |   | To use generation<br>Indicated, in a regimen with mifepristone, for the medical termination of intrauterine pregnancy through<br>70 days gestation.   | 4                                 | N/A   | N/A         | Females Only           | Y               | Y                               | Only covered for non-FDA<br>approved indication in the<br>PADP program   | 11/30/2021            |
| Drugs                             | 54993         | Contraceptive pills for birth<br>control                   | 1 pack (1 pack = 21- or<br>28-tablet pack; 3 packs<br>= 91-tablet pack) | 4/1/2002                | N/A         | contraceptive pills for birth<br>control                    | Indicated as birth control.   | 14 in a 12-month<br>interval      | 8 years   | 55 years    | Females Only           | ¥               | Ŷ                               | 3/2024: Effective 2/1/2024,<br>HCPCS billing unit of 1 pack,<br>Clarified to be defined as 1<br>pack = 21- or 28-tablet pack.<br>Suggested max monthly<br>updated to match NCTracks 14<br>packs per year, effective<br>7/1/2019. Use of code limited<br>to LHDS. |                       |