

CDER Therapeutic Biologic Products

This list is intended to include all the Center for Drug Evaluation and Research (CDER) user fee billable therapeutic biological products and potencies approved under Section 351 of the Public Health Service Act. The Orange Book includes a section entitled "Drug Products with Approval under Section 505 of the Act Administered by CBER." Included on that list are several products that have been transferred to CDER which would be considered billable also.

Program fees are assessed for each potency in which the approved (non-revoked, non-suspended) product is manufactured in final dosage form. When evaluating the specific strength or potency of a drug in final dosage form for purposes of assessing program fees for liquid parenteral biological products, CDER intends to take into consideration both the total amount of drug substance in mass or units of activity in a product and the concentration of drug substance (mass or units of activity per unit volume of product). Biologic products considered to have a different strength or potency in a final dosage form will be given separate entries in the Biologics List and assessed separate program fees. An auto-injector that has the same strength or potency as a prefilled syringe or vial will generally be assessed a separate prescription drug program fee. In certain circumstances, products which have been discontinued from marketing but are still licensed are not assessed program fees. Those products are identified on the CDER Discontinued Biologic Product List section.

The potency information contained in this list is based on information in our database. Companies are responsible for alerting CDER to any discrepancies regarding potency information. For product approvals after October 1, 2005, the Biologics License Application Submission Tracking Number (BLA STN) approval date reflects the approval date of the product. For product approvals prior to October 1, 2005, the BLA STN approval date reflects the approval date of the original BLA. Product Number is the FDA assigned number to identify the application products. Each strength is a separate product.

The list is updated three times a year. (Latest Update – June 2026)

***** CDER Billable Biologic Product List *****

Applicant/License No: ABBVIE INC / 1889

Trade Name: CREON

Proper Name: PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 020725 / 0 | 1 | 4/30/2009 | 30,000USP UNITS; 6,000USP UNITS; 19,000USP UNITS (30,000USP UNITS; 6,000USP UNITS; 19,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 020725 / 0 | 2 | 4/30/2009 | 60,000USP UNITS; 12,000USP UNITS; 38,000USP UNITS (60,000USP UNITS; 12,000USP UNITS; 38,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 020725 / 0 | 3 | 4/30/2009 | 120,000USP UNITS; 24,000USP UNITS; 76,000USP UNITS (120,000USP UNITS; 24,000USP UNITS; 76,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |

Applicant/License No: ABBVIE INC / 1889

| | | | |
|------------|---|-----------|--|
| 020725 / 0 | 4 | 7/12/2011 | 15,000USP UNITS; 3,000USP UNITS; 9,500USP UNITS (15,000USP UNITS; 3,000USP UNITS; 9,500USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
|------------|---|-----------|--|

| | | | |
|------------|---|-----------|--|
| 020725 / 0 | 5 | 3/14/2013 | 180,000USP UNITS; 36,000USP UNITS; 114,000USP UNITS (180,000USP UNITS; 36,000USP UNITS; 114,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
|------------|---|-----------|--|

Trade Name: DECNUPAZ**Proper Name:** PIVEKIMAB SUNIRINE-PVZY

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761460 / 0 | 1 | 5/27/2026 | 2 MG (2 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: EMRELIS**Proper Name:** TELISOTUZUMAB VEDOTIN-TLLV

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761384 / 0 | 1 | 5/14/2025 | 20 MG (20 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761384 / 0 | 2 | 5/14/2025 | 100 MG (100 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: HUMIRA**Proper Name:** ADALIMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125057 / 0 | 2 | 11/23/2015 | 40 MG/0.4 ML (40 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: ABBVIE INC / 1889

| | | | |
|------------|----|------------|---|
| 125057 / 0 | 5 | 12/31/2002 | 40 MG/0.8 ML (40 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125057 / 0 | 6 | 12/31/2002 | 10 MG/0.1 ML (10 MG/0.1 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125057 / 0 | 7 | 12/31/2002 | 20 MG/0.2 ML (20 MG/0.2 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125057 / 0 | 8 | 12/31/2002 | 40 MG/0.4 ML (40 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125057 / 0 | 9 | 12/31/2002 | 80 MG/0.8 ML (80 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125057 / 0 | 10 | 12/31/2002 | 80 MG/0.8 ML (80 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: SKYRIZI**Proper Name:** RISANKIZUMAB-RZAA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761105 / 0 | 2 | 4/26/2021 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761105 / 0 | 3 | 4/26/2021 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761105 / 0 | 4 | 6/16/2022 | 360 MG/2.4ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / KIT |

Applicant/License No: ABBVIE INC / 1889

| | | | |
|------------|---|-----------|---|
| 761105 / 0 | 5 | 3/22/2023 | 90 MG/ML (90 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

| | | | |
|------------|---|-----------|-------------------------------|
| 761105 / 0 | 6 | 9/23/2022 | 180 MG/1.2ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / KIT |

| | | | |
|------------|---|----------|---|
| 761105 / 0 | 7 | 9/3/2025 | 180 MG/1.2 ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|---|
| 761262 / 0 | 1 | 6/16/2022 | 600 MG/10 ML (60 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: SURVANTA**Proper Name:** BERACTANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|----------|---|
| 020032 / 0 | 1 | 7/1/1991 | 100 MG/4 ML & 200 MG/ 8 ML (25 MG/ML) |
| | | | SUSPENSION / INTRATRACHEAL / SINGLE-DOSE VIAL |

Applicant/License No: ABLYNX NV / 2085**Trade Name:** CABLIVI**Proper Name:** CAPLACIZUMAB-YHDP

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|----------|---|
| 761112 / 0 | 1 | 2/6/2019 | 11 MG (11 MG/VIAL) |
| | | | POWDER / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: ACROTECH BIOPHARMA LLC / 2159**Trade Name:** ZEVALIN**Proper Name:** IBRITUMOMAB TIUXETAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125019 / 0 | 1 | 2/19/2002 | 3.2 MG/2 ML (1.6 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ADC THERAPEUTICS SA / 2166**Trade Name:** ZYNLONTA**Proper Name:** LONCASTUXIMAB TESIRINE-LPYL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761196 / 0 | 1 | 4/23/2021 | 10MG (10MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: AKESO BIOPHARMA CO LTD / 2253**Trade Name:****Proper Name:** PENPULIMAB-KCQX

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761258 / 0 | 1 | 4/23/2025 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ALEXION PHARMACEUTICALS INC / 1743**Trade Name:** KANUMA**Proper Name:** SEBELIPASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125561 / 0 | 1 | 12/8/2015 | 20 MG/10 ML (2 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: SOLIRIS**Proper Name:** ECULIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125166 / 0 | 1 | 3/16/2007 | 300 MG/30 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: STRENSIQ**Proper Name:** ASFOTASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125513 / 0 | 1 | 10/23/2015 | 18 MG/0.45 ML (18 MG/0.45 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 125513 / 0 | 2 | 10/23/2015 | 80 MG/0.8 ML (80 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 125513 / 0 | 3 | 10/23/2015 | 28 MG/0.7 ML (28 MG/0.7 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 125513 / 0 | 4 | 10/23/2015 | 40 MG/ML (40 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: ULTOMIRIS**Proper Name:** RAVULIZUMAB-CWVZ

Applicant/License No: ALEXION PHARMACEUTICALS INC / 1743

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761108 / 0 | 2 | 10/9/2020 | 300 MG/3 ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761108 / 0 | 3 | 10/9/2020 | 1,100 MG/11 ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ALLERGAN INC / 1145**Trade Name:** BOTOX**Proper Name:** BOTULINUM TOXIN TYPE A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---------------|----------|---------------|---|
| 103000 / 0 | 1 | 12/9/1991 | 100 U (100 U/VIAL) POWDER / INTRAMUSCULAR, INTRADERMAL, INTRADETRUSOR / SINGLE-DOSE VIAL |
| 103000 / 5122 | 5 | 11/10/2005 | 200 U (200 U/VIAL) POWDER / INTRAMUSCULAR, INTRADERMAL, INTRADETRUSOR / SINGLE-DOSE VIAL |

Trade Name: BOTOX COSMETIC**Proper Name:** BOTULINUM TOXIN TYPE A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---------------|----------|---------------|---|
| 103000 / 5000 | 2 | 4/12/2002 | 100 U (100 U/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |
| 103000 / 5101 | 4 | 4/14/2005 | 50 U (50 U/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: ALTOR BIOSCIENCE LLC AN INDIRECT WHOLLY OWNED SUBSIDIARY OF IMMUNITYBI**Trade Name:** ANKTIVA**Proper Name:** NOGAPENDEKIN ALFA INBAKICEPT PMLN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761336 / 0 | 1 | 4/22/2024 | 400 MCG/0.4 ML (400 MCG/0.4 ML) SOLUTION / INTRAVESICAL / SINGLE-DOSE VIAL |

Applicant/License No: AMGEN INC / 1080**Trade Name:** AIMOVIG**Proper Name:** ERENUMAB-AOOE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761077 / 0 | 2 | 5/17/2018 | 70 MG (70 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761077 / 1 | 4 | 3/11/2019 | 140 MG (140 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: ARANESP**Proper Name:** DARBEPOETIN ALFA IN POLYSORBATE SOLUTION

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103951 / 0 | 1 | 9/17/2001 | 25 MCG/ML (25 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103951 / 0 | 2 | 9/17/2001 | 40 MCG/ML (40 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103951 / 0 | 3 | 9/17/2001 | 60 MCG/ML (60 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: AMGEN INC / 1080

| | | | |
|------------|----|-----------|--|
| 103951 / 0 | 4 | 9/17/2001 | 100 MCG/ML (100 MCG/ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| <hr/> | | | |
| 103951 / 0 | 6 | 9/17/2001 | 200 MCG/ML (200 MCG/ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| <hr/> | | | |
| 103951 / 0 | 8 | 9/17/2001 | 25 MCG/0.42 ML (25 MCG/0.42 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 9 | 9/17/2001 | 40 MCG/0.4 ML (40 MCG/0.4 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 10 | 9/17/2001 | 60 MCG/0.3 ML (60 MCG/0.3 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 11 | 9/17/2001 | 100 MCG/0.5 ML (100 MCG/0.5 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 12 | 9/17/2001 | 150 MCG/0.3 ML (150 MCG/0.3 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 13 | 9/17/2001 | 200 MCG/0.4 ML (200 MCG/0.4 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 14 | 9/17/2001 | 300 MCG/0.6 ML (300 MCG/0.6 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 15 | 9/17/2001 | 500 MCG/ML (500 MCG/ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| <hr/> | | | |
| 103951 / 0 | 31 | 9/17/2001 | 10 MCG/0.4 ML (10 MCG/0.4 ML) |
| | | | SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: AMGEN INC / 1080**Trade Name:** BLINCYTO**Proper Name:** BLINATUMOMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125557 / 0 | 1 | 12/3/2014 | 35 MCG (35 MCG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: EPOGEN**Proper Name:** EPOETIN ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 103234 / 0 | 6 | 6/1/1989 | 10,000 U/ML (10,000 U/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL |
| 103234 / 0 | 7 | 6/1/1989 | 20,000 U/ML (10,000 U/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL |

Proper Name: EPOETIN ALFA - PRESERVATIVE FREE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103234 / 0 | 1 | 6/1/1989 | 2000 U/ML (2000 U/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103234 / 0 | 2 | 6/1/1989 | 3000 U/ML (3000 U/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103234 / 0 | 3 | 6/1/1989 | 4000 U/ML (4000 U/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103234 / 0 | 4 | 6/1/1989 | 10,000 U/ML (10,000 U/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: AMGEN INC / 1080**Trade Name:** EVENITY**Proper Name:** ROMOSOZUMAB-AQQG

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761062 / 0 | 1 | 4/9/2019 | 105 MG/1.17 ML (90 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: IMDELLTRA**Proper Name:** TARLATAMAB-DLLE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761344 / 0 | 1 | 5/16/2024 | 1 MG/VIAL (1 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761344 / 0 | 2 | 5/16/2024 | 10 MG/VIAL (10 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: NEULASTA**Proper Name:** PEGFILGRASTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125031 / 0 | 1 | 1/31/2002 | 6 MG/0.6 ML (6 MG/0.6 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: NEULASTA ONPRO**Proper Name:** PEGFILGRASTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--------------|----------|---------------|--|
| 125031 / 181 | 2 | 9/23/2015 | 6 MG/0.6 ML (6 MG/0.6 ML) SOLUTION / SUBCUTANEOUS / KIT |

Trade Name: NEUPOGEN

Applicant/License No: AMGEN INC / 1080

Proper Name: FILGRASTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 103353 / 0 | 1 | 2/20/1991 | 300 MCG/ML (300 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103353 / 0 | 2 | 2/20/1991 | 300 MCG/0.5 ML (300 MCG/0.5 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 103353 / 0 | 3 | 2/20/1991 | 480 MCG/1.6 ML (300 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103353 / 0 | 4 | 2/20/1991 | 480 MCG/0.8 ML (480 MCG/0.8 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: NPLATE

Proper Name: ROMIPLOSTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--------------|----------|---------------|--|
| 125268 / 0 | 1 | 8/22/2008 | 250 MCG (250 MCG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 125268 / 0 | 2 | 8/22/2008 | 500 MCG (500 MCG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 125268 / 165 | 3 | 7/22/2019 | 125 MCG (125 MCG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: PROCRIT

Proper Name: EPOETIN ALFA - PRESERVATIVE FREE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: AMGEN INC / 1080

103234 / 0 5 6/1/1989 40,000 U/ML (40,000 U/ML)

SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL

Trade Name: PROLIA**Proper Name:** DENOSUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

125320 / 0 1 6/1/2010 60 MG/ML (60 MG/ML)

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Trade Name: REPATHA**Proper Name:** EVOLOCUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

125522 / 0 1 8/27/2015 140 MG/ML (140 MG/ML)

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

125522 / 0 3 8/27/2015 140 MG/ML (140 MG/ML)

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

Trade Name: VECTIBIX**Proper Name:** PANITUMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

125147 / 0 1 9/27/2006 100 MG/5 ML (20 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

125147 / 0 3 9/27/2006 400 MG/20 ML (20 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: XGEVA**Proper Name:** DENOSUMAB

Applicant/License No: AMGEN INC / 1080

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125320 / 7 | 2 | 11/18/2010 | 120 MG/1.7 ML (70 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: AMICUS THERAPEUTICS US LLC / 2224**Trade Name:** POMBILITI**Proper Name:** CIPAGLUCOSIDASE ALFA-ATGA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761204 / 0 | 1 | 9/28/2023 | 105 MG (105 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: AMPHASTAR PHARMACEUTICAL INC / 2179**Trade Name:** AMPHADASE**Proper Name:** HYALURONIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021665 / 0 | 1 | 10/26/2004 | 150 UNITS/ML (150 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: ARGENX BV / 2217**Trade Name:** VYVGART**Proper Name:** EFGARTIGIMOD ALFA-FCAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761195 / 0 | 1 | 12/17/2021 | 400 MG/20 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: VYVGART HYTRULO

Applicant/License No: ARGENX BV / 2217**Proper Name:** EFGARTIGIMOD ALFA AND HYALURONIDASE-QVFC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761304 / 0 | 1 | 6/20/2023 | 1008 MG AND 11200 UNITS/5.6 ML (180 MG AND 2000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761304 / 0 | 2 | 4/10/2025 | 1000 MG AND 10000 UNITS/5 ML (200 MG AND 2000 UNITS/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: ASCENDIS PHARMA ENDOCRINOLOGY DIVISION AS / 2165**Trade Name:** SKYTROFA**Proper Name:** LONAPEGSOMATROPIN-TCGD

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761177 / 0 | 1 | 8/26/2021 | 3 MG (3 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 2 | 8/26/2021 | 3.6 MG (3.6 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 3 | 8/26/2021 | 4.3 MG (4.3 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 4 | 8/26/2021 | 5.2 MG (5.2 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 5 | 8/26/2021 | 6.3 MG (6.3 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 6 | 8/26/2021 | 7.6 MG (7.6 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |

Applicant/License No: ASCENDIS PHARMA ENDOCRINOLOGY DIVISION AS / 2165

| | | | |
|------------|---|-----------|-----------------------------------|
| 761177 / 0 | 7 | 8/26/2021 | 9.1 MG (9.1 MG) |
| | | | POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 8 | 8/26/2021 | 11 MG (11 MG) |
| | | | POWDER / SUBCUTANEOUS / CARTRIDGE |
| 761177 / 0 | 9 | 8/26/2021 | 13.3 MG (13.3 MG) |
| | | | POWDER / SUBCUTANEOUS / CARTRIDGE |

Applicant/License No: ASTELLAS PHARMA US INC / 2124**Trade Name:** PADCEV**Proper Name:** ENFORTUMAB VEDOTIN-EJFV

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761137 / 0 | 1 | 12/18/2019 | 20 MG (20 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761137 / 0 | 2 | 12/18/2019 | 30 MG (30 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: VYLOY**Proper Name:** ZOLBETUXIMAB-CLZB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761365 / 0 | 1 | 10/18/2024 | 100 MG (100 MG) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761365 / 0 | 2 | 3/24/2025 | 300 MG (300 MG) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ASTRAZENECA AB / 2059**Trade Name:** BEYFORTUS**Proper Name:** NIRSEVIMAB-ALIP

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761328 / 0 | 1 | 7/17/2023 | 50 MG/0.5 ML (50 MG/0.5 ML) SOLUTION / INTRAMUSCULAR / PREFILLED SYRINGE |
| 761328 / 0 | 2 | 7/17/2023 | 100 MG/ML (100 MG/ML) SOLUTION / INTRAMUSCULAR / PREFILLED SYRINGE |

Trade Name: IMJUDO**Proper Name:** TREMELIMUMAB-ACTL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761270 / 0 | 1 | 11/10/2022 | 25 MG/1.25 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761270 / 0 | 2 | 11/10/2022 | 300 MG/15 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: SAPHNELO**Proper Name:** ANIFROLUMAB-FNIA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761123 / 0 | 1 | 7/30/2021 | 300 MG/2 ML (150 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761451 / 0 | 1 | 4/24/2026 | 300 MG/2 ML (150 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ASTRAZENECA AB / 2059

761451 / 0 2 4/24/2026 120 MG/0.8 ML (120 MG/0.8 ML)

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Trade Name: SAPHNELO PEN**Proper Name:** ANIFROLUMAB-FNIA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761451 / 0 | 3 | 4/24/2026 | 120 MG/0.8 ML (120 MG/0.8 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: TEZSPIRE**Proper Name:** TEZEPELUMAB-EKKO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761224 / 0 | 1 | 12/17/2021 | 210 MG/1.91 ML (110 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL & PREFILLED SYRINGE |

| | | | |
|------------|---|----------|--|
| 761224 / 0 | 2 | 2/1/2023 | 210 MG/1.91 ML (110 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: ASTRAZENECA UK LTD / 2043**Trade Name:** FASENRA**Proper Name:** BENRALIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761070 / 0 | 1 | 11/14/2017 | 30 MG/ML (30 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

| | | | |
|------------|---|-----------|--|
| 761070 / 0 | 2 | 10/3/2019 | 30 MG/ML (30 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: ASTRAZENECA UK LTD / 2043

761070 / 0 3 4/8/2024 10 MG/0.5 ML (10 MG/0.5 ML)

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Trade Name: IMFINZI**Proper Name:** DURVALUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761069 / 0 | 1 | 5/1/2017 | 120 MG/2.4 ML (50 MG/ML) |
| 761069 / 0 | 2 | 5/1/2017 | 500 MG/10 ML (50 MG/ML) |

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: AUXILIUM PHARMACEUTICALS INC / 1816**Trade Name:** XIAFLEX**Proper Name:** CLOSTRIDIAL COLLAGENASE HISTOLYTICUM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125338 / 0 | 1 | 2/2/2010 | 0.9 MG (0.9 MG/VIAL) |

POWDER / INTRALESIONAL / SINGLE-DOSE VIAL

Applicant/License No: BAXTER HEALTHCARE CORP / 0140**Trade Name:** MYXREDLIN**Proper Name:** INSULIN HUMAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 208157 / 0 | 1 | 6/20/2019 | 100 UNITS/100 ML (100 UNITS/100 ML) |

SOLUTION / INTRAVENOUS / SINGLE DOSE CONTAINER

Applicant/License No: BAYER HEALTHCARE PHARMACEUTICALS INC / 1778**Trade Name:** BETASERON**Proper Name:** INTERFERON BETA-1B

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103471 / 0 | 1 | 7/23/1993 | 0.3 MG (0.3 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: BEONE MEDICINES USA INC / 2232**Trade Name:** TEVIMBRA**Proper Name:** TISLELIZUMAB-JSGR

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761232 / 0 | 1 | 3/13/2024 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: BIOGEN INC / 1697**Trade Name:** AVONEX**Proper Name:** INTERFERON BETA 1A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103628 / 0 | 2 | 5/17/1996 | 30 MCG/0.5 ML (30 MCG/0.5 ML) SOLUTION / INTRAMUSCULAR / PREFILLED SYRINGE |
| 103628 / 0 | 3 | 5/17/1996 | 30 MCG/0.5 ML (30 MCG/0.5 ML) SOLUTION / INTRAMUSCULAR / AUTOINJECTOR-PREFILLED PEN |

Trade Name: PLEGRIDY**Proper Name:** PEGINTERFERON BETA-1A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: BIOGEN INC / 1697

| | | | |
|------------|---|-----------|---|
| 125499 / 0 | 1 | 8/15/2014 | 63 MCG/0.5 ML (63 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125499 / 0 | 2 | 8/15/2014 | 125 MCG/0.5 ML (125 MCG/0.5 ML) SOLUTION / INTRAMUSCULAR, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125499 / 0 | 3 | 8/15/2014 | 63 MCG/0.5 ML (63 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125499 / 0 | 4 | 8/15/2014 | 94 MCG/0.5 ML (94 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125499 / 0 | 5 | 8/15/2014 | 94 MCG/0.5 ML (94 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125499 / 0 | 6 | 8/15/2014 | 125 MCG/0.5 ML (125 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: TYSABRI**Proper Name:** NATALIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125104 / 0 | 1 | 11/23/2004 | 300 MG/15 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: BIOMARIN PHARMACEUTICAL INC / 1649**Trade Name:** ALDURAZYME**Proper Name:** LARONIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125058 / 0 | 1 | 4/30/2003 | 2.9 MG/5 ML (0.58 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: BRINEURA**Proper Name:** CERLIPONASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761052 / 0 | 1 | 4/27/2017 | 150 MG/5 ML (30 MG/ML) SOLUTION / INTRAVENTRICULAR / SINGLE-DOSE VIAL |

Trade Name: NAGLAZYME**Proper Name:** GALSULFASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125117 / 0 | 1 | 5/31/2005 | 5 MG/ 5ML (1 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: PALYNZIQ**Proper Name:** PEGVALIASE-PQPZ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761079 / 0 | 1 | 5/24/2018 | 2.5 MG (5 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761079 / 0 | 2 | 5/24/2018 | 10 MG (20 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: BIOMARIN PHARMACEUTICAL INC / 1649

761079 / 0 3 5/24/2018 20 MG (20 MG/ML)

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Trade Name: VIMIZIM**Proper Name:** ELOSULFASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

125460 / 0 1 2/14/2014 5 MG/5 ML (1 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: BOEHRINGER INGELHEIM PHARMACEUTICALS INC / 2006**Trade Name:** PRAXBIND**Proper Name:** IDARUCIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761025 / 0 1 10/16/2015 2500 MG/50 ML (50 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: BRISTOL MYERS SQUIBB CO / 1713**Trade Name:** EMLICITI**Proper Name:** ELOTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761035 / 0 1 11/30/2015 300 MG (300 MG/VIAL)

POWDER / INTRAVENOUS / SINGLE-DOSE VIAL

761035 / 0 2 11/30/2015 400 MG (400 MG/VIAL)

POWDER / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: NULOJIX**Proper Name:** BELATACEPT

Applicant/License No: BRISTOL MYERS SQUIBB CO / 1713

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125288 / 0 | 1 | 6/15/2011 | 250 MG (250 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: OPDIVO**Proper Name:** NIVOLUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125554 / 0 | 1 | 12/22/2014 | 40 MG/4 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125554 / 0 | 2 | 12/22/2014 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125554 / 0 | 3 | 12/22/2014 | 240 MG/24 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125554 / 0 | 4 | 8/27/2021 | 120 MG/12 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: OPDIVO QVANTIG**Proper Name:** NIVOLUMAB AND HYALURONIDASE-NVHY

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761381 / 0 | 1 | 12/27/2024 | 600 MG AND 10,000 UNITS/5 ML (120 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: OPDUALAG**Proper Name:** NIVOLUMAB AND RELATLIMAB-RMBW

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: BRISTOL MYERS SQUIBB CO / 1713

| | | | |
|------------|---|-----------|--|
| 761234 / 0 | 1 | 3/18/2022 | 240 MG AND 80 MG/20 ML (12 MG AND 4 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ORENCIA**Proper Name:** ABATACEPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--------------|----------|---------------|--|
| 125118 / 0 | 1 | 12/23/2005 | 250 MG (250 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125118 / 122 | 2 | 7/29/2011 | 50 MG/0.4 ML (50 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125118 / 122 | 3 | 7/29/2011 | 87.5 MG/0.7 ML (87.5 MG/0.7 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125118 / 122 | 4 | 7/29/2011 | 125 MG/ML (125 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125118 / 122 | 5 | 7/29/2011 | 125 MG/ML (125 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: YERVOY**Proper Name:** IPILIMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125377 / 0 | 1 | 3/25/2011 | 50 MG/10 ML (5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125377 / 0 | 2 | 3/25/2011 | 200 MG/40 ML (5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: BTG INTERNATIONAL INC / 1861**Trade Name:** VORAXAZE**Proper Name:** GLUCARPIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125327 / 0 | 1 | 1/17/2012 | 1,000 U (1,000 U/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: CELGENE CORPORATION / 2114**Trade Name:** REBLOZYL**Proper Name:** LUSPATERCEPT-AAMT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761136 / 0 | 1 | 11/8/2019 | 25 MG (25 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761136 / 0 | 2 | 11/8/2019 | 75 MG (75 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: CELLTRION INC / 1996**Trade Name:** ZYMFENTRA**Proper Name:** INFLIXIMAB-DYYB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761358 / 0 | 1 | 10/20/2023 | 120 MG/ML (120 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761358 / 0 | 2 | 10/20/2023 | 120 MG/ML (120 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: CHECKPOINT THERAPEUTICS INCORPORATED / 2275**Trade Name:** UNLOXCYT**Proper Name:** COSIBELIMAB-IPDL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761297 / 0 | 1 | 12/13/2024 | 300 MG/5 ML (60 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: CHIESI FARMACEUTICI SPA / 2245**Trade Name:** ELFABRIO**Proper Name:** PEGUNIGALSIDASE ALFA-IWXJ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761161 / 0 | 1 | 5/9/2023 | 20 MG/10 ML (2 ML/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761161 / 0 | 2 | 5/17/2024 | 5 MG/2.5 ML (2 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: LAMZEDE**Proper Name:** VELMANASE ALFA-TYCV

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761278 / 0 | 1 | 2/16/2023 | 10 MG/VIAL (10 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: CHIESI USA INC / 2150**Trade Name:** CUROSURF**Proper Name:** PORACTANT ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 020744 / 0 | 1 | 11/18/1999 | 120 MG/1.5ML & 240 MG/3ML (80 MG/ML) SUSPENSION / INTRATRACHEAL / SINGLE-DOSE VIAL |

Trade Name: MYALEPT**Proper Name:** METRELEPTIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125390 / 0 | 1 | 2/24/2014 | 11.3 MG (11.3 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: REVCOVI**Proper Name:** ELAPEGADEMASE-LVLR

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761092 / 0 | 1 | 10/5/2018 | 2.4 MG/1.5 ML (1.6 MG/ML) SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: CIS BIO INTERNATIONAL / 2204**Trade Name:** PULMOTECH MAA**Proper Name:** TECHNETIUM TC 99M ALBUMIN AGGREGATED

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 210089 / 0 | 1 | 3/20/2020 | 2 MG (2 MG) POWDER / INTRAVENOUS, INTRAPERITONEAL / MULTI-DOSE VIAL |

Applicant/License No: CITIUS ONCOLOGY INC / 2290**Trade Name:** LYMPHIR**Proper Name:** DENILEUKIN DIFTITOX-CXDL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761312 / 0 | 1 | 8/7/2024 | 300 MCG/VIAL (300 MCG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: COHERUS BIOSCIENCES INC / 2023**Trade Name:** LOQTORZI**Proper Name:** TORIPALIMAB-TPZI

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761240 / 0 | 1 | 10/27/2023 | 240 MG/6 ML (40 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: CSL BEHRING LLC / 1767**Trade Name:** ANDEMBRY**Proper Name:** GARADACIMAB-GXII

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761367 / 0 | 1 | 6/16/2025 | 200 MG/1.2 ML (200 MG/1.2 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761367 / 0 | 2 | 6/16/2025 | 200 MG/1.2 ML (167 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: DAIICHI SANKYO INC / 2128**Trade Name:** DATROWAY**Proper Name:** DATOPOTAMAB DERUXTECAN-DLNK

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761394 / 0 | 1 | 1/17/2025 | 100 MG (100 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761464 / 0 | 1 | 6/23/2025 | 100 MG (100 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ENHERTU**Proper Name:** FAM-TRASTUZUMAB DERUXTECAN-NXKI

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761139 / 0 | 1 | 12/20/2019 | 100 MG (100 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: DENALI THERAPEUTICS INC / 2385**Trade Name:** AVLAYAH**Proper Name:** TIVIDENOFUSP ALFA-EKNM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761485 / 0 | 1 | 3/24/2026 | 150 MG (150 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: DIGESTIVE CARE INC / 2184**Trade Name:** PERTZYE**Proper Name:** PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 022175 / 0 | 1 | 5/17/2012 | 30,250USP UNITS; 8,000USP UNITS; 28,750USP UNITS (30,250USP UNITS; 8,000USP UNITS; 28,750USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022175 / 0 | 2 | 5/17/2012 | 60,500USP UNITS; 16,000USP UNITS; 57,500USP UNITS (60,500USP UNITS; 16,000USP UNITS; 57,500USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022175 / 0 | 3 | 10/6/2016 | 15,125USP UNITS; 4,000USP UNITS; 14,375USP UNITS (15,125USP UNITS; 4,000USP UNITS; 14,375USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022175 / 0 | 4 | 7/13/2017 | 90,750USP UNITS; 24,000USP UNITS; 86,250USP UNITS (90,750USP UNITS; 24,000USP UNITS; 86,250USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |

Applicant/License No: DOMPE FARMACEUTICI SPA / 2074**Trade Name:** OXERVATE**Proper Name:** CENEGERMIN-BKBJ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761094 / 0 | 1 | 8/22/2018 | 20 MCG/ML (0.002%) SOLUTION / OPHTHALMIC / MULTI-DOSE VIAL |

Applicant/License No: DYAX CORPORATION / 1789**Trade Name:** KALBITOR**Proper Name:** ECALLANTIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125277 / 0 | 1 | 12/1/2009 | 10 MG/ML (10 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: TAKHZYRO**Proper Name:** LANADELUMAB-FLYO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761090 / 0 | 1 | 8/23/2018 | 300 MG/2 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: EISAI INC / 1862**Trade Name:** LEQEMBI**Proper Name:** LECANEMAB-IRMB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761269 / 0 | 1 | 1/6/2023 | 500 MG/5 ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

| | | | |
|------------|---|----------|--|
| 761269 / 0 | 2 | 1/6/2023 | 200 MG/2 ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
|------------|---|----------|--|

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761375 / 0 | 1 | 8/29/2025 | 500 MG/5 ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: EISAI INC / 1862

| | | | |
|------------|---|-----------|---|
| 761375 / 0 | 2 | 8/29/2025 | 200 MG/2 ML (100 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

| | | | |
|------------|---|-----------|--|
| 761375 / 0 | 3 | 8/29/2025 | 360 MG/1.8 ML (200 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: ELI LILLY AND CO / 1891**Trade Name:** BASAGLAR KWIKPEN**Proper Name:** INSULIN GLARGINE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 205692 / 0 | 1 | 12/16/2015 | 300 UNITS/3 ML (100 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: BASAGLAR TEMPO PEN**Proper Name:** INSULIN GLARGINE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 205692 / 0 | 2 | 11/15/2019 | 300 UNITS/3 ML (100 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: CYRAMZA**Proper Name:** RAMUCIRUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125477 / 0 | 1 | 4/21/2014 | 100 MG/10 ML (10 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125477 / 0 | 2 | 4/21/2014 | 500 MG/50 ML (10 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ELI LILLY AND CO / 1891

Trade Name: EBGLYSS

Proper Name: LEBRIKIZUMAB-LBKZ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761306 / 0 | 1 | 9/13/2024 | 250 MG/2 ML (250 MG/2 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761306 / 0 | 2 | 9/13/2024 | 250 MG/2 ML (250 MG/2 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: EMGALITY

Proper Name: GALCANEZUMAB-GNLM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761063 / 0 | 1 | 9/27/2018 | 120 MG/ML (120 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761063 / 0 | 2 | 9/27/2018 | 120 MG/ML (120 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761063 / 3 | 3 | 6/4/2019 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: HUMALOG

Proper Name: INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 020563 / 0 | 1 | 6/14/1996 | 1000 UNITS/10 ML (100 UNITS/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL |

Applicant/License No: ELI LILLY AND CO / 1891

| | | | |
|------------|---|----------|-------------------------------------|
| 020563 / 0 | 2 | 8/6/1998 | 300 UNITS/3 ML (100 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / CARTRIDGE |

| | | | |
|------------|---|-----------|---|
| 020563 / 0 | 6 | 9/20/2019 | 300 UNITS/3 ML (100 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: HUMALOG KWIKPEN**Proper Name:** INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020563 / 0 | 3 | 9/6/2007 | 300 UNITS/3 ML (100 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 205747 / 0 | 1 | 5/26/2015 | 600 UNITS/3 ML (200 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: HUMALOG KWIKPEN JUNIOR**Proper Name:** INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020563 / 0 | 4 | 6/6/2017 | 300 UNITS/3 ML (100 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: HUMALOG MIX 50/50 KWIKPEN**Proper Name:** INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021018 / 0 | 2 | 12/22/1999 | 150 UNITS/3 ML; 150 UNITS/3 ML (50 UNITS/ML; 50 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: ELI LILLY AND CO / 1891**Trade Name:** HUMALOG MIX 75/25**Proper Name:** INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021017 / 0 | 1 | 12/22/1999 | 750 UNITS/10 ML; 250 UNITS/10 ML (75 UNITS/ML; 25 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: HUMALOG MIX 75/25 KWIKPEN**Proper Name:** INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021017 / 0 | 2 | 9/6/2007 | 225 UNITS/3 ML; 75 UNITS/3 ML (75 UNITS/ML; 25 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: HUMATROPE**Proper Name:** SOMATROPIN RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 019640 / 0 | 5 | 2/4/1999 | 6 MG (6 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 019640 / 0 | 6 | 2/4/1999 | 12 MG (12 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 019640 / 0 | 7 | 2/4/1999 | 24 MG (24 MG) POWDER / SUBCUTANEOUS / CARTRIDGE |

Trade Name: HUMULIN R**Proper Name:** INSULIN HUMAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: ELI LILLY AND CO / 1891

018780 / 0 4 3/31/1994 10000 UNITS/20 ML (500 UNITS/ML)

SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL

Trade Name: HUMULIN R KWIKPEN**Proper Name:** INSULIN HUMAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

018780 / 0 2 12/29/2015 1500 UNITS/3 ML (500 UNITS/ML)

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

Trade Name: KISUNLA**Proper Name:** DONANEMAB-AZBT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761248 / 0 1 7/2/2024 350 MG/20 ML (17.5 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: LYUMJEV**Proper Name:** INSULIN LISPRO-AABC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761109 / 0 1 6/15/2020 1000 UNITS/10 ML (100 UNITS/ML)

SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL

761109 / 0 5 6/15/2020 300 UNITS/3 ML (100 UNITS/ML)

SOLUTION / SUBCUTANEOUS / CARTRIDGE

Trade Name: LYUMJEV JUNIOR KWIKPEN**Proper Name:** INSULIN LISPRO-AABC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: ELI LILLY AND CO / 1891

| | | | |
|--|---|-----------|---------------------------------|
| 761109 / 0 | 3 | 6/15/2020 | 300 UNITS/3 ML (100 UNITS/ML) |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | |

Trade Name: LYUMJEV KWIKPEN**Proper Name:** INSULIN LISPRO-AABC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--|----------|---------------|--|
| 761109 / 0 | 2 | 6/15/2020 | 300 UNITS/3 ML (100 UNITS/ML) |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | |

| | | | |
|--|---|-----------|---------------------------------|
| 761109 / 0 | 6 | 6/15/2020 | 600 UNITS/3 ML (200 UNITS/ML) |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | |

Trade Name: OMVOH**Proper Name:** MIRIKIZUMAB-MRKZ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---|----------|---------------|--|
| 761279 / 0 | 1 | 10/26/2023 | 300 MG/15 ML (20 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

| | | | |
|--|---|------------|-------------------------|
| 761279 / 0 | 2 | 10/26/2023 | 100 MG/ML (100 MG/ML) |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | |

| | | | |
|---|---|-----------|-------------------------|
| 761279 / 0 | 3 | 4/29/2024 | 100 MG/ML (100 MG/ML) |
| SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | | | |

| | | | |
|---|---|-----------|---------------------------|
| 761279 / 0 | 4 | 1/15/2025 | 200 MG/2 ML (100 MG/ML) |
| SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | | | |

| | | | |
|--|---|-----------|---------------------------|
| 761279 / 0 | 5 | 1/15/2025 | 200 MG/2 ML (100 MG/ML) |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | |

Applicant/License No: ELI LILLY AND CO / 1891

Trade Name: TALTZ

Proper Name: IXEKIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125521 / 0 | 1 | 3/22/2016 | 80 MG/ML (80 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125521 / 0 | 2 | 3/22/2016 | 80 MG/ML (80 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: TRULICITY

Proper Name: DULAGLUTIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|-------------|----------|---------------|---|
| 125469 / 0 | 3 | 9/18/2014 | 0.75 MG/0.5 ML (0.75 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125469 / 0 | 4 | 9/18/2014 | 1.5 MG/0.5 ML (1.5 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125469 / 36 | 5 | 9/3/2020 | 3 MG/0.5 ML (3 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125469 / 36 | 6 | 9/3/2020 | 4.5 MG/0.5 ML (4.5 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: ELUSYS THERAPEUTICS INC / 1907**Trade Name:** ANTHIM**Proper Name:** OBILTOXAXIMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125509 / 1 | 1 | 3/18/2016 | 600 MG/6 ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: EMD SERONO INC / 1773**Trade Name:** BAVENCIO**Proper Name:** AVELUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761049 / 0 | 1 | 3/23/2017 | 200 MG/10 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: GONAL-F**Proper Name:** FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020378 / 0 | 4 | 2/28/2001 | 1,050 IU/VIAL (600 IU/ML) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL |
| 020378 / 0 | 5 | 3/26/2004 | 450 IU/VIAL (600 IU/ML) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: GONAL-F RFF REDI-JECT**Proper Name:** FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: EMD SERONO INC / 1773

| | | | |
|------------|---|-----------|--|
| 021684 / 0 | 1 | 5/25/2004 | 300 IU/0.5ML (300 IU/0.5ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

| | | | |
|------------|---|-----------|--|
| 021684 / 0 | 2 | 5/25/2004 | 450 IU/0.75 ML (450 IU/0.75 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

| | | | |
|------------|---|-----------|--|
| 021684 / 0 | 3 | 5/25/2004 | 900 IU/1.5 ML (600 IU/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: OVIDREL**Proper Name:** CHORIOGONADOTROPIN ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021149 / 0 | 2 | 9/20/2000 | 0.25 MG /0.5 ML (0.25 MG /0.5 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: REBIF**Proper Name:** INTERFERON BETA 1A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103780 / 0 | 1 | 3/7/2002 | 8.8 MCG/0.2 ML (8.8 MCG/0.2 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 103780 / 0 | 2 | 3/7/2002 | 8.8 MCG/0.2 ML (8.8 MCG/0.2 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 103780 / 0 | 3 | 3/7/2002 | 22 MCG/0.5 ML (22 MCG/0.5 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 103780 / 0 | 4 | 3/7/2002 | 22 MCG/0.5 ML (22 MCG/0.5 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: EMD SERONO INC / 1773

| | | | |
|------------|---|----------|---|
| 103780 / 0 | 5 | 3/7/2002 | 44 MCG/0.5 ML (44 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 103780 / 0 | 6 | 3/7/2002 | 44 MCG/0.5 ML (44 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: SEROSTIM**Proper Name:** SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020604 / 0 | 1 | 8/23/1996 | 6 MG/VIAL (6 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 020604 / 0 | 2 | 8/23/1996 | 5 MG/VIAL (5 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 020604 / 0 | 3 | 7/25/1997 | 4 MG/VIAL (4 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL |

Applicant/License No: EMERGENT MANUFACTURING OPERATIONS BALTIMORE LLC / 2083**Trade Name:** EBANGA**Proper Name:** ANSUVIMAB-ZYKL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761172 / 0 | 1 | 12/21/2020 | 400 MG/VIAL (400 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: RAXIBACUMAB**Proper Name:** RAXIBACUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: EMERGENT MANUFACTURING OPERATIONS BALTIMORE LLC / 2083

125349 / 0 1 12/14/2012 1700 MG/34 ML (50 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: ETON PHARMACEUTICALS INC / 2393**Trade Name:** INCRELEX**Proper Name:** MECASERMIN RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|------------------------|
| 021839 / 0 | 1 | 8/30/2005 | 40 MG/4 ML (10MG/ML) |
|------------|---|-----------|------------------------|

SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL

Applicant/License No: EVIVE BIOTECHNOLOGY SINGAPORE PTE LTD / 2248**Trade Name:** RYZNEUTA**Proper Name:** EFBEMALENOGRASTIM ALFA-VUXW

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|------------|-----------------------|
| 761134 / 0 | 1 | 11/16/2023 | 20 MG/ML (20 MG/ML) |
|------------|---|------------|-----------------------|

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Applicant/License No: EVOLUS INC / 2070**Trade Name:** JEUVEAU**Proper Name:** PRABOTULINUMTOXINA-XVFS

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|----------|------------------------------|
| 761085 / 0 | 1 | 2/1/2019 | 100 UNITS (100 UNITS/VIAL) |
|------------|---|----------|------------------------------|

POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL

Applicant/License No: FERRING PHARMACEUTICALS INC / 2112**Trade Name:** MENOPUR**Proper Name:**

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021663 / 0 | 1 | 10/29/2004 | 75 IU/VIAL (75 IU/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: NOVAREL**Proper Name:** GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 017016 / 0 | 6 | 1/15/1974 | 5000 UNITS/VIAL (5000 UNITS/VIAL) POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL |

Trade Name: ZOMACTON**Proper Name:** SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 019774 / 0 | 2 | 1/4/2002 | 5 MG/VIAL (5 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL |
| 019774 / 0 | 3 | 3/7/2012 | 10 MG/VIAL (10 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL |

Applicant/License No: FRESENIUS KABI USA LLC / 2146**Trade Name:** CHORIONIC GONADOTROPIN**Proper Name:** GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 017067 / 0 | 2 | 3/5/1973 | 10000 UNITS/VIAL (10000 UNITS/VIAL) POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL |

Applicant/License No: GALDERMA LABORATORIES LP / 2289**Trade Name:** NEMLUVIO**Proper Name:** NEMOLIZUMAB-ILTO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761390 / 0 | 1 | 8/12/2024 | 30 MG (30 MG) POWDER / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: GENENTECH INC / 1048**Trade Name:** ACTEMRA**Proper Name:** TOCILIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125276 / 0 | 1 | 1/8/2010 | 80 MG/4 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125276 / 0 | 2 | 1/8/2010 | 200 MG/10 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125276 / 0 | 3 | 1/8/2010 | 400 MG/20 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |

Applicant/License No: GENENTECH INC / 1048

| | | | |
|-------------|---|------------|--|
| 125472 / 1 | 1 | 10/21/2013 | 162 MG/0.9 ML (162 MG/0.9 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125472 / 29 | 2 | 11/19/2018 | 162 MG/0.9 ML (162 MG/0.9 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: ACTIVASE**Proper Name:** ALTEPLASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103172 / 0 | 2 | 11/13/1987 | 50 MG (50 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: AVASTIN**Proper Name:** BEVACIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125085 / 0 | 1 | 2/26/2004 | 100 MG/4 ML (25 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125085 / 0 | 2 | 2/26/2004 | 400 MG/16 ML (25 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: CATHFLO ACTIVASE**Proper Name:** ALTEPLASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103172 / 0 | 1 | 9/4/2001 | 2 MG (2 MG/VIAL) |
| | | | POWDER / INTRACATHETER / SINGLE-DOSE VIAL |

Trade Name: COLUMVI**Proper Name:** GLOFITAMAB-GXBM

Applicant/License No: GENENTECH INC / 1048

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761309 / 0 | 1 | 6/15/2023 | 2.5 MG/2.5 ML (1 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761309 / 0 | 2 | 6/15/2023 | 10 MG/10 ML (1 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ENSPRYNG**Proper Name:** SATRALIZUMAB-MWGE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761149 / 0 | 1 | 8/14/2020 | 120 MG/ML (120 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: GAZYVA**Proper Name:** OBINUYUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125486 / 0 | 1 | 11/1/2013 | 1000 MG/40 ML (25 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: HEMLIBRA**Proper Name:** EMICIZUMAB-KXWH

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761083 / 0 | 1 | 11/16/2017 | 30 MG/ML (30 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761083 / 0 | 2 | 11/16/2017 | 60 MG/0.4 ML (60 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: GENENTECH INC / 1048

| | | | |
|------------|---|------------|--|
| 761083 / 0 | 3 | 11/16/2017 | 105 MG/0.7 ML (105 MG/0.7 ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761083 / 0 | 4 | 11/16/2017 | 150 MG/ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761083 / 0 | 5 | 3/16/2023 | 300 MG/2 ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761083 / 0 | 6 | 1/31/2024 | 12 MG/0.4 ML (12 MG/0.4 ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: HERCEPTIN**Proper Name:** TRASTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---------------|----------|---------------|--|
| 103792 / 5336 | 2 | 2/10/2017 | 150 MG (150 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: HERCEPTIN HYLECTA**Proper Name:** TRASTUZUMAB AND HYALURONIDASE-OYSK

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761106 / 0 | 1 | 2/28/2019 | 600 MG AND 10000 UNITS/5ML (120 MG AND 2000 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: KADCYLA**Proper Name:** ADO-TRASTUZUMAB EMTANSINE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
| | | | |

Applicant/License No: GENENTECH INC / 1048

| | | | |
|------------|---|-----------|---|
| 125427 / 0 | 1 | 2/22/2013 | 100 MG (100 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

| | | | |
|------------|---|-----------|---|
| 125427 / 0 | 2 | 2/22/2013 | 160 MG (160 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: LUCENTIS**Proper Name:** RANIBIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--------------|----------|---------------|--|
| 125156 / 0 | 1 | 6/30/2006 | 0.5 MG/0.05 ML (0.5 MG/0.05 ML) |
| | | | SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL & PREFILLED SYRINGE |
| 125156 / 076 | 2 | 8/10/2012 | 0.3 MG/0.05 ML (0.3 MG/0.05 ML) |
| | | | SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL & PREFILLED SYRINGE |

Trade Name: LUNSUMIO**Proper Name:** MOSUNETUZUMAB-AXGB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761263 / 0 | 1 | 12/22/2022 | 1 MG/ML (1 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761263 / 0 | 2 | 12/22/2022 | 30 MG/30 ML (1 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: LUNSUMIO VELO**Proper Name:** MOSUNETUZUMAB-AXGB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
| | | | |

Applicant/License No: GENENTECH INC / 1048

| | | | |
|------------|---|------------|--|
| 761263 / 0 | 3 | 12/19/2025 | 5 MG/0.5 ML (5 MG/0.5 ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

| | | | |
|------------|---|------------|--|
| 761263 / 0 | 4 | 12/19/2025 | 45 MG/ML (45 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: OCREVUS**Proper Name:** OCRELIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761053 / 0 | 1 | 3/28/2017 | 300 MG/10 ML (30 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: OCREVUS ZUNOVO**Proper Name:** OCRELIZUMAB AND HYALURONIDASE-OCSQ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761371 / 0 | 1 | 9/13/2024 | 920 MG AND 23,000 UNITS/23 ML (40 MG AND 1,000 UNITS/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: PERJETA**Proper Name:** PERTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125409 / 0 | 1 | 6/8/2012 | 420 MG/14 ML (30 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: PHESGO**Proper Name:** PERTUZUMAB, TRASTUZUMAB, AND HYALURONIDASE-ZZXF

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
| | | | |

Applicant/License No: GENENTECH INC / 1048

| | | | |
|------------|---|-----------|---|
| 761170 / 0 | 1 | 6/29/2020 | 600 MG, 600 MG, 20000 UNITS/10 ML (60 MG, 60 MG, 2000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
|------------|---|-----------|---|

| | | | |
|------------|---|-----------|--|
| 761170 / 0 | 2 | 6/29/2020 | 1200 MG, 600 MG, 30000 UNITS/15 ML (80 MG, 40 MG, 2000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
|------------|---|-----------|--|

Trade Name: PIASKY**Proper Name:** CROVALIMAB-AKKZ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761388 / 0 | 1 | 6/20/2024 | 340 MG/2 ML (170 MG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: POLIVY**Proper Name:** POLATUZUMAB VEDOTIN-PIIQ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761121 / 0 | 1 | 6/10/2019 | 140 MG (140 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761121 / 0 | 2 | 9/18/2020 | 30 MG (30 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: PULMOZYME**Proper Name:** DORNASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103532 / 0 | 1 | 12/30/1993 | 2.5 MG/2.5 ML (1 MG/ML) SOLUTION / INHALATION / AMPULE |

Trade Name: RITUXAN**Proper Name:** RITUXIMAB

Applicant/License No: GENENTECH INC / 1048

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103705 / 0 | 1 | 11/26/1997 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 103705 / 0 | 2 | 11/26/1997 | 500 MG/50 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: RITUXAN HYCELA**Proper Name:** RITUXIMAB AND HYALURONIDASE HUMAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761064 / 0 | 1 | 6/22/2017 | 1400 MG AND 23400 U/11.7 ML (120 MG AND 2000 U/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761064 / 0 | 2 | 6/22/2017 | 1600 MG AND 26800 U/13.4 ML (120 MG AND 2000 U/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: SUSVIMO**Proper Name:** RANIBIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761197 / 0 | 1 | 10/22/2021 | 100 MG/ML (100 MG/ML) SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL |

Trade Name: TECENTRIQ**Proper Name:** ATEZOLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761034 / 0 | 1 | 5/18/2016 | 1200 MG/20 ML (60 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: GENENTECH INC / 1048

761034 / 18 2 3/8/2019 840 MG/14 ML (60 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: TECENTRIQ HYBREZA**Proper Name:** ATEZOLIZUMAB AND HYALURONIDASE-TQJS

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761347 / 0 | 1 | 9/12/2024 | 1,875 MG ATEZOLIZUMAB AND 30,000 UNITS HYALURONIDASE PER 15 ML (125 MG/2,000 UNITS PER ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: TNKASE**Proper Name:** TENECTEPLASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103909 / 0 | 1 | 6/2/2000 | 50 MG (50 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 103909 / 0 | 2 | 2/28/2025 | 25 MG (25 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: VABYSMO**Proper Name:** FARICIMAB-SVOA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761235 / 0 | 1 | 1/28/2022 | 120 MG/ML (120 MG/ML) SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL & PREFILLED SYRINGE |

Trade Name: XOLAIR**Proper Name:** OMALIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: GENENTECH INC / 1048

| | | | |
|---------------|---|-----------|---|
| 103976 / 0 | 1 | 6/20/2003 | 150 MG/VIAL (150 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103976 / 5231 | 3 | 9/28/2018 | 75 MG/0.5 ML (75 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 103976 / 5231 | 4 | 9/28/2018 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 103976 / 0 | 5 | 8/17/2023 | 300MG/2ML (300MG/2ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 103976 / 0 | 6 | 8/17/2023 | 75 MG/0.5 ML (75 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 103976 / 0 | 7 | 8/17/2023 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 103976 / 0 | 8 | 8/17/2023 | 300 MG/2 ML (300 MG/2 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: GENMAB US INC / 2293**Trade Name:** EPKINLY**Proper Name:** EPCORITAMAB-BYSP

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761324 / 0 | 1 | 5/19/2023 | 4 MG/0.8 ML (4 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761324 / 0 | 2 | 5/19/2023 | 48 MG/0.8 ML (48 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: GENZYME CORP / 1596

Trade Name: CAMPATH

Proper Name: ALEMTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103948 / 0 | 1 | 5/7/2001 | 30 MG/ML (30 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: CEREZYME

Proper Name: IMIGLUCERASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020367 / 0 | 2 | 9/22/1999 | 400 UNITS/VIAL (400 UNITS/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: FABRAZYME

Proper Name: AGALSIDASE BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103979 / 0 | 1 | 4/24/2003 | 5 MG (5 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 103979 / 0 | 2 | 4/24/2003 | 35 MG (35 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: LEMTRADA

Proper Name: ALEMTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---------------|----------|---------------|---|
| 103948 / 5139 | 2 | 11/14/2014 | 12 MG/1.2 ML (10MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: LUMIZYME

Applicant/License No: GENZYME CORP / 1596**Proper Name:** ALGLUCOSIDASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125291 / 0 | 1 | 5/24/2010 | 50 MG (50 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: NEXVIAZYME**Proper Name:** AVALGLUCOSIDASE ALFA-NGPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761194 / 0 | 1 | 8/6/2021 | 100 MG/VIAL (100 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: THYROGEN**Proper Name:** THYROTROPIN ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020898 / 0 | 1 | 11/30/1998 | 0.9 MG/VIAL (0.9 MG/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: GILEAD SCIENCES INC / 2258**Trade Name:** HEPCLUDEX**Proper Name:** BULEVIRTIDE-GMOD

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761468 / 0 | 1 | 5/22/2026 | 8.5 MG (8.5 MG) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: TRODELVY**Proper Name:** SACITUZUMAB GOVITECAN-HZIY

Applicant/License No: GILEAD SCIENCES INC / 2258

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761115 / 0 | 1 | 4/22/2020 | 180 MG (180 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: GLAXOSMITHKLINE LLC / 1727**Trade Name:** BENLYSTA**Proper Name:** BELIMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125370 / 0 | 1 | 3/9/2011 | 120 MG (120 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

| | | | |
|------------|---|----------|---|
| 125370 / 0 | 2 | 3/9/2011 | 400 MG (400 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
|------------|---|----------|---|

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761043 / 0 | 1 | 7/20/2017 | 200 MG/ML (200 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

| | | | |
|------------|---|-----------|---|
| 761043 / 0 | 2 | 7/20/2017 | 200 MG/ML (200 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
|------------|---|-----------|---|

Trade Name: BLENREP**Proper Name:** BELANTAMAB MAFODOTIN-BLMF

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761440 / 0 | 1 | 10/23/2025 | 70 MG/VIAL (70 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: EXDENSUR

Applicant/License No: GLAXOSMITHKLINE LLC / 1727**Proper Name:** DEPEMOKIMAB-ULAA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761458 / 0 | 1 | 12/17/2025 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761458 / 0 | 2 | 12/17/2025 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: JEMPERLI**Proper Name:** DOSTARLIMAB-GXLY

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761174 / 0 | 1 | 4/22/2021 | 500 MG/10 ML (50 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: NUCALA**Proper Name:** MEPOLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125526 / 0 | 1 | 11/14/2015 | 100 MG (100 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761122 / 0 | 1 | 6/6/2019 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761122 / 0 | 2 | 6/6/2019 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: GLAXOSMITHKLINE LLC / 1727

761122 / 0 3 1/22/2022 40 MG/0.4 ML (40 MG/0.4 ML)

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Applicant/License No: HALOZYME THERAPEUTICS INC / 2187**Trade Name:** HYLENEX RECOMBINANT**Proper Name:** HYALURONIDASE RECOMBINANT HUMAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|---------------------------------|
| 021859 / 0 | 1 | 12/2/2005 | 150 UNITS/VIAL (150 UNITS/ML) |
|------------|---|-----------|---------------------------------|

SOLUTION / INTERSTITIAL, INTRAMUSCULAR, INTRAOCULAR,
PERIBULBAR, RETROBULBAR, SOFT TISSUE, SUBCUTANEOUS / SINGLE-DOSE VIAL**Applicant/License No: HORIZON THERAPEUTICS IRELAND DAC / 2022****Trade Name:** ACTIMMUNE**Proper Name:** INTERFERON GAMMA-1B

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|-----------------------------------|
| 103836 / 0 | 1 | 2/25/1999 | 100 MCG/0.5 ML (100 MCG/0.5 ML) |
|------------|---|-----------|-----------------------------------|

SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL

Trade Name: KRYSTEXXA**Proper Name:** PEGLOTICASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|-------------------|
| 125293 / 0 | 1 | 9/14/2010 | 8MG/ML (8MG/ML) |
|------------|---|-----------|-------------------|

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: TEPEZZA**Proper Name:** TEPROTUMUMAB-TRBW

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: HORIZON THERAPEUTICS IRELAND DAC / 2022

| | | | |
|---|---|-----------|------------------------|
| 761143 / 0 | 1 | 1/21/2020 | 500 MG (500 MG/VIAL) |
| POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

Trade Name: UPLIZNA**Proper Name:** INEBILIZUMAB-CDON

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---|----------|---------------|--|
| 761142 / 0 | 1 | 6/11/2020 | 100 MG/10 ML (10 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

Applicant/License No: HUGEL INC / 2237**Trade Name:** LETYBO**Proper Name:** LETIBOTULINUMTOXINA-WLBG

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---|----------|---------------|--|
| 761225 / 0 | 1 | 2/29/2024 | 50 UNITS/VIAL (50 UNITS/VIAL) |
| POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | | | |
| 761225 / 0 | 2 | 2/29/2024 | 100 UNITS/VIAL (100 UNITS/VIAL) |
| POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | | | |

Applicant/License No: IMCLONE LLC / 1827**Trade Name:** ERBITUX**Proper Name:** CETUXIMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---|----------|---------------|--|
| 125084 / 0 | 1 | 2/12/2004 | 100 MG/50 ML (2 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

Applicant/License No: IMCLONE LLC / 1827

125084 / 0 2 2/12/2004 200 MG/100 ML (2 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: IMMEDICA PHARMA AB / 2342**Trade Name:** LOARGYS**Proper Name:** PEGZILARGINASE-NBLN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761211 / 0 | 1 | 2/23/2026 | 2 MG/0.4 ML (2 MG/0.4 ML) |
| 761211 / 0 | 2 | 2/23/2026 | 5 MG/ML (5 MG/ML) |

SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL

SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL

Applicant/License No: IMMUNEX CORP / 1132**Trade Name:** ENBREL**Proper Name:** ETANERCEPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------------|----------|---------------|--|
| 103795 / 0 | 1 | 11/2/1998 | 25 MG (25 MG/VIAL) |
| 103795 / 5184 | 2 | 9/27/2004 | 50 MG/ML (50 MG/ML) |
| 103795 / 0 | 3 | 9/27/2004 | 50 MG/ML (50 MG/ML) |
| 103795 / 5184/ | 4 | 9/27/2004 | 25 MG/0.5 ML (25 MG/0.5 ML) |

POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL & PREFILLED SYRINGE

Applicant/License No: IMMUNEX CORP / 1132

103795 / 5556 5 9/14/2017 50 MG/ML (50 MG/ML)

SOLUTION / SUBCUTANEOUS / CARTRIDGE

Applicant/License No: IMMUNOCORE LIMITED / 2239**Trade Name:** KIMMTRAK**Proper Name:** TEBENTAFUSP-TEBN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761228 / 0 1 1/25/2022 100 MCG/0.5 ML (100 MCG/0.5 ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: IMMUNOGEN INC / 2288**Trade Name:** ELAHERE**Proper Name:** MIRVETUXIMAB SORAVTANSINE-GYNX

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761310 / 0 1 11/14/2022 100 MG/20 ML (5 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: INCYTE CORPORATION / 2228**Trade Name:** MONJUVI**Proper Name:** TAFASITAMAB-CXIX

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761163 / 0 1 7/31/2020 200 MG (200 MG/VIAL)

POWDER / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: NIKTIMVO**Proper Name:** AXATILIMAB-CSFR

Applicant/License No: INCYTE CORPORATION / 2228

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761411 / 0 | 1 | 8/14/2024 | 50 MG/ML (50 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761411 / 0 | 2 | 1/14/2025 | 9 MG/0.18 ML (9 MG/0.18 ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761411 / 0 | 3 | 1/14/2025 | 22 MG/0.44 ML (22 MG/0.44 ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ZYNYZ**Proper Name:** RETIFANLIMAB-DLWR

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761334 / 0 | 1 | 3/22/2023 | 500 MG/20ML (25 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: IOVANCE BIOTHERAPEUTICS MANUFACTURING LLC / 2353**Trade Name:** PROLEUKIN**Proper Name:** ALDESLEUKIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 103293 / 0 | 1 | 5/5/1992 | 22 MIU (22 MIU/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: IPSEN BIOPHARM LIMITED / 1787**Trade Name:** DYSPORT**Proper Name:** ABOBOTULINUMTOXINA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125274 / 0 | 1 | 4/29/2009 | 300 UNITS (300 UNITS/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |
| 125274 / 0 | 2 | 4/29/2009 | 500 UNITS (500 UNITS/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: ISO TEX DIAGNOSTICS INC / 2189**Trade Name:** MEGATOPE**Proper Name:** ALBUMIN IODINATED I-131 SERUM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 017837 / 0 | 2 | 2/23/1976 | 1mCi/VIAL (1mCi/VIAL) SOLUTION / INTRAVENOUS / MULTI-DOSE VIAL |

Applicant/License No: JANSSEN BIOTECH INC / 1864**Trade Name:** DARZALEX**Proper Name:** DARATUMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761036 / 0 | 1 | 11/16/2015 | 100 MG/5 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761036 / 0 | 2 | 11/16/2015 | 400 MG/20 ML (20 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: DARZALEX FASPRO**Proper Name:** DARATUMUMAB AND HYALURONIDASE-FIHJ

Applicant/License No: JANSSEN BIOTECH INC / 1864

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761145 / 0 | 1 | 5/1/2020 | 1800 MG/30000 UNITS/15 ML (120 MG/2000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: IMAAVY**Proper Name:** NIPOCALIMAB-AAHU

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761430 / 0 | 1 | 4/29/2025 | 300 MG/1.62 ML (185 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

| | | | |
|------------|---|-----------|---|
| 761430 / 0 | 2 | 4/29/2025 | 1200 MG/6.5 ML (185 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
|------------|---|-----------|---|

Trade Name: REMICADE**Proper Name:** INFLIXIMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 103772 / 0 | 1 | 8/24/1998 | 100 MG (100 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: RYBREVANT**Proper Name:** AMIVANTAMAB-VMJW

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761210 / 0 | 1 | 5/21/2021 | 350 MG/7 ML (50 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: RYBREVANT FASPRO**Proper Name:** AMIVANTAMAB AND HYALURONIDASE-LPUJ

Applicant/License No: JANSSEN BIOTECH INC / 1864

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761433 / 0 | 1 | 12/17/2025 | 1,600 MG AND 20,000 UNITS/10 ML (160 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761433 / 0 | 2 | 12/17/2025 | 2,240 MG AND 28,000 UNITS/14 ML (160 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761433 / 0 | 3 | 2/13/2026 | 2,400 MG AND 30,000 UNITS/15ML (160 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761433 / 0 | 4 | 2/13/2026 | 3,520 MG AND 44,000 UNITS/22ML (160 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: SIMPONI**Proper Name:** GOLIMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125289 / 0 | 1 | 4/24/2009 | 50 MG/0.5 ML (50 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125289 / 0 | 2 | 4/24/2009 | 50 MG/0.5 ML (50 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 125289 / 0 | 3 | 4/24/2009 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125289 / 0 | 4 | 4/24/2009 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: SIMPONI ARIA**Proper Name:** GOLIMUMAB

Applicant/License No: JANSSEN BIOTECH INC / 1864

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125433 / 0 | 1 | 7/18/2013 | 50 MG/4 ML (12.5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: STELARA**Proper Name:** USTEKINUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125261 / 0 | 1 | 9/25/2009 | 90 MG/ML (90 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125261 / 0 | 2 | 9/25/2009 | 45 MG/0.5 ML (45 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL & PREFILLED SYRINGE |

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761044 / 0 | 1 | 9/23/2016 | 130 MG/26 ML (5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: TALVEY**Proper Name:** TALQUETAMAB-TGVS

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761342 / 0 | 1 | 8/9/2023 | 3 MG/1.5 ML (2 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761342 / 0 | 2 | 8/9/2023 | 40 MG/ML (40 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: TECVAYLI**Proper Name:** TECLISTAMAB-CQYV

Applicant/License No: JANSSEN BIOTECH INC / 1864

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761291 / 0 | 1 | 10/25/2022 | 30 MG/3 ML (10 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761291 / 0 | 2 | 10/25/2022 | 153 MG/1.7ML (90 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: TREMFYA**Proper Name:** GUSELKUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761061 / 0 | 1 | 7/13/2017 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761061 / 0 | 4 | 9/11/2024 | 200 MG/2 ML (200 MG/2 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761061 / 0 | 5 | 9/11/2024 | 200 MG/20 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: TREMFYA PEN**Proper Name:** GUSELKUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761061 / 0 | 2 | 9/11/2024 | 100 MG/ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761061 / 0 | 3 | 9/11/2024 | 200 MG/2 ML (200 MG/2 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: JAZZ PHARMACEUTICALS IRELAND LIMITED / 2167**Trade Name:** RYLAZE**Proper Name:** ASPARAGINASE ERWINIA CHRYSANTHEMI (RECOMBINANT)-RYWN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761179 / 0 | 1 | 6/30/2021 | 10 MG/0.5 ML (10 MG/0.5 ML) SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Trade Name: ZIIHERA**Proper Name:** ZANIDATAMAB-HRII

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761416 / 0 | 1 | 11/20/2024 | 300 MG (300 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: JUBILANT DRAXIMAGE INC / 2186**Trade Name:** DRAXIMAGE MAA**Proper Name:** TECHNETIUM TC 99M ALBUMIN AGGREGATED KIT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 017881 / 0 | 1 | 12/30/1987 | 2.5 MG/VIAL (2.5 MG/VIAL) POWDER / INTRAVENOUS / MULTI-DOSE VIAL |

Applicant/License No: KINIKSA PHARMACEUTICALS UK LTD / 2236**Trade Name:** ARCALYST**Proper Name:** RILONACEPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125249 / 0 | 1 | 2/27/2008 | 220MG (220MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: KYOWA KIRIN INC / 2077**Trade Name:** CRYSVITA**Proper Name:** BUROSUMAB-TWZA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761068 / 0 | 1 | 4/17/2018 | 10 MG/ML (10 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761068 / 0 | 2 | 4/17/2018 | 20 MG/ML (20 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761068 / 0 | 3 | 4/17/2018 | 30 MG/ML (30 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: POTELIGEO**Proper Name:** MOGAMULIZUMAB-KPKC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761051 / 0 | 1 | 8/8/2018 | 20 MG/5 ML (4 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: LEO PHARMA AS / 2169**Trade Name:** ADBRY**Proper Name:** TRALOKINUMAB-LDRM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761180 / 0 | 1 | 12/27/2021 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761180 / 0 | 2 | 6/12/2024 | 300 MG/2 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: LEO PHARMA AS / 2169**Trade Name:** SPEVIGO**Proper Name:** SPESOLIMAB-SBZO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761244 / 0 | 1 | 9/1/2022 | 450 MG/7.5 ML (60 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761244 / 0 | 2 | 3/18/2024 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761244 / 0 | 3 | 5/28/2025 | 300 MG/2 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: LIB THERAPEUTICS INC / 2352**Trade Name:** LEROCHOL**Proper Name:** LERODALCIBEP-LIGA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761427 / 0 | 1 | 12/12/2025 | 300 MG/1.2 ML (250 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: LUNDBECK SEATTLE BIOPHARMACEUTICALS INC / 2097**Trade Name:** VYEPTI**Proper Name:** EPTINEZUMAB-JJMR

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761119 / 0 | 1 | 2/21/2020 | 100 MG/ML (100 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: MANNKIND CORP / 2190**Trade Name:** AFREZZA**Proper Name:** INSULIN RECOMBINANT HUMAN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 022472 / 0 | 1 | 6/27/2014 | 4 UNITS/INH (4 UNITS/INH) POWDER / INHALATION / CARTRIDGE |
| 022472 / 0 | 2 | 6/27/2014 | 8 UNITS/INH (8 UNITS/INH) POWDER / INHALATION / CARTRIDGE |
| 022472 / 0 | 3 | 4/17/2015 | 12 UNITS/INH (12 UNITS/INH) POWDER / INHALATION / CARTRIDGE |

Applicant/License No: MERCK AND CO INC DIV MERCK SHARP AND DOHME / 0002**Trade Name:** ENFLONSIA**Proper Name:** CLESROVIMAB-CFOR

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761432 / 0 | 1 | 6/9/2025 | 105 MG/0.7 ML (105 MG/0.7 ML) SOLUTION / INTRAMUSCULAR / PREFILLED SYRINGE |

Trade Name: KEYTRUDA**Proper Name:** PEMBROLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125514 / 1 | 2 | 1/15/2015 | 100 MG/4 ML (25 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: KEYTRUDA QLEX**Proper Name:** PEMBROLIZUMAB AND BERAHYALURONIDASE ALFA-PMPH

Applicant/License No: MERCK AND CO INC DIV MERCK SHARP AND DOHME / 0002

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761467 / 0 | 1 | 9/19/2025 | 395 MG AND 4,800 UNITS/2.4 ML (165 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761467 / 0 | 2 | 9/19/2025 | 790 MG AND 9,600 UNITS/4.8 ML (165 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: WINREVAIR**Proper Name:** SOTATERCEPT-CSRK

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761363 / 0 | 1 | 3/26/2024 | 45 MG/VIAL (45 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761363 / 0 | 2 | 3/26/2024 | 60 MG/VIAL (60 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: MERZ PHARMACEUTICALS GMBH / 1830**Trade Name:** XEOMIN**Proper Name:** INCOBOTULINUMTOXINA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125360 / 0 | 1 | 7/30/2010 | 50 U (50 U/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |
| 125360 / 0 | 2 | 7/30/2010 | 100 U (100 U/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |
| 125360 / 3 | 3 | 11/20/2015 | 200 U (200 U/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: NOVARTIS PHARMACEUTICALS CORP / 1244**Trade Name:** ADAKVEO**Proper Name:** CRIZANLIZUMAB-TMCA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761128 / 0 | 1 | 11/15/2019 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ARZERRA**Proper Name:** OFATUMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125326 / 0 | 1 | 10/26/2009 | 100 MG/5 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125326 / 0 | 2 | 10/26/2009 | 1000 MG/50 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: BEOVU**Proper Name:** BROLUCIZUMAB-DBLL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761125 / 0 | 1 | 10/7/2019 | 6 MG/0.05 ML (6 MG/0.05 ML) SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL |

Trade Name: COSENTYX**Proper Name:** SECUKINUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125504 / 0 | 2 | 1/21/2015 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: NOVARTIS PHARMACEUTICALS CORP / 1244

| | | | |
|------------|---|-----------|--|
| 125504 / 0 | 3 | 1/21/2015 | 150 MG/ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

| | | | |
|------------|---|-----------|---|
| 125504 / 0 | 4 | 5/28/2021 | 75 MG/0.5 ML (75 MG/0.5 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

| | | | |
|------------|---|-----------|---|
| 125504 / 0 | 5 | 5/11/2023 | 300 MG/2 ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

| | | | |
|------------|---|-----------|--|
| 125504 / 0 | 6 | 5/11/2023 | 300 MG/2 ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|---|
| 761349 / 0 | 1 | 10/6/2023 | 125 MG/5 ML (25 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: EXTAVIA**Proper Name:** INTERFERON BETA-1B

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|--|
| 125290 / 0 | 1 | 8/14/2009 | 0.3 MG (0.3 MG/VIAL) |
| | | | POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: ILARIS**Proper Name:** CANAKINUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|------------|--|
| 125319 / 0 | 3 | 12/22/2016 | 150 MG/ML (150 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: KESIMPTA

Applicant/License No: NOVARTIS PHARMACEUTICALS CORP / 1244**Proper Name:** OFATUMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|-------------|----------|---------------|---|
| 125326 / 70 | 3 | 8/20/2020 | 20 MG/0.4 ML (20 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125326 / 70 | 4 | 8/20/2020 | 20 MG/0.4 ML (20 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: SIMULECT**Proper Name:** BASILIXIMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103764 / 0 | 1 | 5/12/1998 | 10 MG (10 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 103764 / 0 | 2 | 5/12/1998 | 20 MG (20 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: NOVO NORDISK INC / 1261**Trade Name:** ALHEMO**Proper Name:** CONCIZUMAB-MTCI

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761315 / 0 | 1 | 12/20/2024 | 60 MG/1.5 ML (40 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761315 / 0 | 2 | 12/20/2024 | 150 MG/1.5 ML (100 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: NOVO NORDISK INC / 1261

761315 / 0 3 12/20/2024 300 MG/3 ML (100 MG/ML)

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

Trade Name: FIASP**Proper Name:** INSULIN ASPART

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

208751 / 0 1 9/27/2017 1000 UNITS/10 ML (100 UNITS/ML)

SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL

Trade Name: FIASP FLEXTOUCH**Proper Name:** INSULIN ASPART

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

208751 / 0 2 9/27/2017 300 UNITS/3 ML (100 UNITS/ML)

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

Trade Name: FIASP PENFILL**Proper Name:** INSULIN ASPART

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

208751 / 0 3 9/24/2018 300 UNITS/3 ML (100 UNITS/ML)

SOLUTION / SUBCUTANEOUS / CARTRIDGE

Trade Name: FIASP PUMPCART**Proper Name:** INSULIN ASPART

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

208751 / 0 4 6/21/2023 160 UNITS/1.6 ML (100 UNITS/ML)

SOLUTION / SUBCUTANEOUS / CARTRIDGE

Trade Name: NORDITROPIN FLEXP

Applicant/License No: NOVO NORDISK INC / 1261

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021148 / 0 | 8 | 3/1/2010 | 5MG/1.5ML (3.33MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 021148 / 0 | 9 | 3/1/2010 | 10 MG/1.5ML (6.67 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 021148 / 0 | 10 | 3/1/2010 | 15MG/1.5ML (10MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 021148 / 0 | 11 | 1/23/2015 | 30MG/3ML (10MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: NOVOLOG

Proper Name: INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020986 / 0 | 1 | 6/7/2000 | 1000 UNITS/10 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: NOVOLOG FLEXPEN

Proper Name: INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 020986 / 0 | 3 | 1/19/2001 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: NOVOLOG MIX 70/30

Proper Name: INSULIN ASPART PROTAMINE RECOMBINANT; INSULIN ASPART RECOMBINANT

Applicant/License No: NOVO NORDISK INC / 1261

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021172 / 0 | 1 | 11/1/2001 | 700 UNITS/10ML; 300 UNITS/10 ML (70 UNITS/ML; 30 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: NOVOLOG MIX 70/30 FLEXPEN**Proper Name:** INSULIN ASPART PROTAMINE RECOMBINANT; INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021172 / 0 | 4 | 5/3/2002 | 210 UNITS/3ML; 90 UNITS/3 ML (70 UNITS/ML; 30 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: NOVOLOG PENFILL**Proper Name:** INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020986 / 0 | 2 | 6/7/2000 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE |

Trade Name: SOGROYA**Proper Name:** SOMAPACITAN-BECO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761156 / 0 | 1 | 8/28/2020 | 10 MG/1.5 ML (6.7 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761156 / 0 | 2 | 10/1/2021 | 5 MG/1.5 ML (3.3 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761156 / 0 | 3 | 4/28/2023 | 15 MG/1.5ML (10 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: NOVO NORDISK INC / 1261**Trade Name:** TRESIBA**Proper Name:** INSULIN DEGLUDEC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 203314 / 0 | 1 | 9/25/2015 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 203314 / 0 | 2 | 9/25/2015 | 600 UNITS/3 ML (200 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 203314 / 0 | 3 | 11/21/2018 | 1000 UNITS/10 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: XULTOPHY 100/3.6**Proper Name:** INSULIN DEGLUDEC; LIRAGLUTIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 208583 / 0 | 1 | 11/21/2016 | 300 UNITS/3 ML; 10.8MG/3 ML (100 UNITS/ML; 3.6MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: NPS PHARMACEUTICALS INC / 1908**Trade Name:** NATPARA**Proper Name:** PARATHYROID HORMONE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125511 / 1 | 1 | 1/23/2015 | 0.4 MG/CARTRIDGE (25 MCG/DOSE) POWDER / SUBCUTANEOUS / CARTRIDGE |
| 125511 / 1 | 2 | 1/23/2015 | 0.8 MG/CARTRIDGE (50 MCG/DOSE) POWDER / SUBCUTANEOUS / CARTRIDGE |

Applicant/License No: NPS PHARMACEUTICALS INC / 1908

| | | | |
|------------|---|-----------|-----------------------------------|
| 125511 / 1 | 3 | 1/23/2015 | 1.21 MG/CARTRIDGE (75 MCG/DOSE) |
| | | | POWDER / SUBCUTANEOUS / CARTRIDGE |

| | | | |
|------------|---|-----------|------------------------------------|
| 125511 / 1 | 4 | 1/23/2015 | 1.61 MG/CARTRIDGE (100 MCG/DOSE) |
| | | | POWDER / SUBCUTANEOUS / CARTRIDGE |

Applicant/License No: OMEROS CORPORATION / 2141**Trade Name:** YARTEMLEA**Proper Name:** NARSOPLIMAB-WUUG

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761152 / 0 | 1 | 12/23/2025 | 370 MG/2 ML (185 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ONY INC / 2192**Trade Name:** INFASURF PRESERVATIVE FREE**Proper Name:** CALFACTANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020521 / 0 | 1 | 7/1/1998 | 105 MG/ 3 ML & 210 MG/ 6 ML (35 MG/ML) |
| | | | SUSPENSION / INTRATRACHEAL / SINGLE-DOSE VIAL |

Applicant/License No: ORGANON USA LLC A SUBSIDIARY OF ORGANON AND CO / 2331**Trade Name:** FOLLISTIM AQ**Proper Name:** FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021211 / 0 | 1 | 3/23/2004 | 300 IU/0.36 ML (300 IU/0.36 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: ORGANON USA LLC A SUBSIDIARY OF ORGANON AND CO / 2331

| | | | |
|------------|---|-----------|--|
| 021211 / 0 | 2 | 3/23/2004 | 600 IU/0.72 ML (600 IU/0.72 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

| | | | |
|------------|---|-----------|--|
| 021211 / 0 | 4 | 2/11/2005 | 900 IU/1.08 ML (900 IU/1.08 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: PREGNYL**Proper Name:** GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 017692 / 0 | 1 | 10/20/1976 | 10000 UNITS/VIAL (10000 UNITS/VIAL) |
| | | | POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL |

Applicant/License No: OTSUKA PHARMACEUTICAL COMPANY LTD / 2387**Trade Name:** VOYXACT**Proper Name:** SIBEPRENIMAB-SZSI

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761434 / 0 | 1 | 11/25/2025 | 400 MG/2 ML (200 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: PARTNER THERAPEUTICS INC / 2087**Trade Name:** BIZENGRI**Proper Name:** ZENOCUTUZUMAB-ZBCO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761352 / 0 | 1 | 12/4/2024 | 375 MG/18.75 ML (20 MG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: LEUKINE**Proper Name:** SARGRAMOSTIM

Applicant/License No: PARTNER THERAPEUTICS INC / 2087

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 103362 / 0 | 1 | 3/5/1991 | 250 MCG (250 MCG/VIAL) POWDER / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: PFIZER INC / 2001**Trade Name:** ELELYSO**Proper Name:** TALIGLUCERASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 022458 / 0 | 1 | 5/1/2012 | 200 UNITS/VIAL (200 UNITS/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ELREXFIO**Proper Name:** ELRANATAMAB-BCMM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761345 / 0 | 1 | 8/14/2023 | 76 MG/1.9 ML (40 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761345 / 0 | 2 | 8/14/2023 | 44 MG/1.1 ML (40 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Trade Name: HYMPAVZI**Proper Name:** MARSTACIMAB-HNCQ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761369 / 0 | 2 | 10/11/2024 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: PFIZER IRELAND PHARMACEUTICALS / 2060**Trade Name:** NGENLA**Proper Name:** SOMATROGON-GHLA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761184 / 0 | 1 | 6/29/2023 | 24 MG/1.2 ML (20 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761184 / 0 | 2 | 6/29/2023 | 60 MG/1.2 ML (50 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: PHARMAAND GMBH / 2291**Trade Name:** PEGASYS**Proper Name:** PEGINTERFERON ALFA 2A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103964 / 0 | 1 | 10/16/2002 | 180 MCG/ML (180 MCG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 103964 / 0 | 2 | 10/16/2002 | 180 MCG/0.5 ML (180 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: PHARMACIA AND UPJOHN CO / 1216**Trade Name:** GENOTROPIN**Proper Name:** SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020280 / 0 | 6 | 8/24/1995 | 5.8 MG/VIAL (5.8 MG/VIAL) POWDER / SUBCUTANEOUS / CARTRIDGE |

Applicant/License No: PHARMACIA AND UPJOHN CO / 1216

020280 / 0 7 10/23/1996 13.8 MG/VIAL (13.8 MG/VIAL)

POWDER / SUBCUTANEOUS / CARTRIDGE

Trade Name: GENOTROPIN PRESERVATIVE FREE**Proper Name:** SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020280 / 0 | 1 | 1/27/1998 | 0.2 MG/VIAL (0.2 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 2 | 1/27/1998 | 0.4 MG/VIAL (0.4 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 3 | 1/27/1998 | 0.6 MG/VIAL (0.6 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 5 | 1/27/1998 | 0.8 MG/VIAL (0.8 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 8 | 1/27/1998 | 1 MG/VIAL (1 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 9 | 1/27/1998 | 1.2 MG/VIAL (1.2 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 10 | 1/27/1998 | 1.4 MG/VIAL (1.4 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |
| 020280 / 0 | 11 | 1/27/1998 | 1.6 MG/VIAL (1.6 MG/VIAL) POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: PHARMACIA AND UPJOHN CO / 1216

| | | | |
|------------|----|-----------|---|
| 020280 / 0 | 12 | 1/27/1998 | 1.8 MG/VIAL (1.8 MG/VIAL) |
| | | | POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |

| | | | |
|------------|----|-----------|---|
| 020280 / 0 | 13 | 1/27/1998 | 2 MG/VIAL (2 MG/VIAL) |
| | | | POWDER / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: SOMAVERT**Proper Name:** PEGVISOMANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021106 / 0 | 1 | 3/25/2003 | 10MG/VIAL (10MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 021106 / 0 | 2 | 3/25/2003 | 15MG/VIAL (15MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 021106 / 0 | 3 | 3/25/2003 | 20MG/VIAL (20MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 021106 / 0 | 4 | 7/31/2014 | 25MG/VIAL (25MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 021106 / 0 | 5 | 7/31/2014 | 30MG/VIAL (30MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: PHARMAESSENTIA CORPORATION / 2155**Trade Name:** BESREMI**Proper Name:** ROPEGINTERFERON ALFA-2B-NJFT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761166 / 0 | 1 | 11/12/2021 | 500 MCG/ML (500 MCG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: PROVENTION BIO INC / 2170**Trade Name:** TZIELD**Proper Name:** TEPLIZUMAB-MZWV

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761183 / 0 | 1 | 11/17/2022 | 2 MG/2 ML (1 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: QOL MEDICAL LLC / 2195**Trade Name:** SUCRAID**Proper Name:** SACROSIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 020772 / 0 | 2 | 5/25/2022 | 17,000 IU/2 ML (8500 IU/ML) SOLUTION / ORAL / SINGLE DOSE CONTAINER |

Applicant/License No: RECORDATI RARE DISEASES INC / 1899**Trade Name:** ENJAYMO**Proper Name:** SUTIMLIMAB-JOME

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761164 / 0 | 1 | 2/4/2022 | 1,100 MG/22 ML (50 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: SYLVANT**Proper Name:** SILTUXIMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125496 / 0 | 1 | 4/23/2014 | 100 MG (100 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125496 / 0 | 2 | 4/23/2014 | 400 MG (400 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: REGENERON PHARMACEUTICALS INC / 1760**Trade Name:** DUPIXENT**Proper Name:** DUPILUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761055 / 0 | 1 | 3/28/2017 | 300 MG/2 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761055 / 7 | 2 | 10/19/2018 | 200 MG/1.14 ML (175 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761055 / 0 | 3 | 6/18/2020 | 300 MG/2 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: REGENERON PHARMACEUTICALS INC / 1760

761055 / 0 4 6/14/2021 200 MG/1.14 ML (175 MG/ML)

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

Trade Name: EVKEEZA**Proper Name:** EVINACUMAB-DGNB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761181 / 0 1 2/11/2021 345 MG/2.3ML (150 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

761181 / 0 2 2/11/2021 1200 MG/8 ML (150 MG/ML)

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Trade Name: EYLEA**Proper Name:** AFLIBERCEPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

125387 / 0 1 11/18/2011 2 MG/0.05 ML (2 MG/0.05 ML)

SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL & PREFILLED SYRINGE

Trade Name: EYLEA HD**Proper Name:** AFLIBERCEPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761355 / 0 1 8/18/2023 8 MG (0.07 ML OF 114.3 MG/ML) (8 MG (0.07 ML OF 114.3 MG/ML))

SOLUTION / INTRAVITREAL / SINGLE-DOSE VIAL

Trade Name: INMAZEB**Proper Name:** ATOLTIVIMAB, MAFTIVIMAB, AND ODESIVIMAB-EBGN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: REGENERON PHARMACEUTICALS INC / 1760

| | | | |
|---|---|------------|--|
| 761169 / 0 | 1 | 10/14/2020 | 241.7 MG/241.7 MG/241.7 MG/14.5 ML (16.67 MG/16.67 MG/16.67 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

| | | | |
|---|---|-----------|--|
| 761169 / 0 | 2 | 9/29/2021 | 483.3 MG/483.3 MG/483.3 MG/14.5 ML (33.33 MG/33.33 MG/33.33 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

Trade Name: LIBTAYO**Proper Name:** CEMIPILIMAB-RWLC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---|----------|---------------|--|
| 761097 / 0 | 1 | 9/28/2018 | 350 MG/7 ML (50 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

Trade Name: LYNOZYFIC**Proper Name:** LINVOSELTAMAB-GCPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---|----------|---------------|--|
| 761400 / 0 | 1 | 7/2/2025 | 5 MG/2.5 ML (2 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

| | | | |
|---|---|----------|---------------------------|
| 761400 / 0 | 2 | 7/2/2025 | 200 MG/10 ML (20 MG/ML) |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | |

Trade Name: PRALUENT**Proper Name:** ALIROCUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--|----------|---------------|--|
| 125559 / 0 | 3 | 7/24/2015 | 75 MG/ML (75 MG/ML) |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | |

Applicant/License No: REGENERON PHARMACEUTICALS INC / 1760

125559 / 0 4 7/24/2015 150 MG/ML (150 MG/ML)

SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN

Trade Name: VEOPOZ**Proper Name:** POZELIMAB-BBFG

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761339 / 0 1 8/18/2023 400 MG/2 ML (200 MG/ML)

SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL

Applicant/License No: REVANCE THERAPEUTICS INC / 2101**Trade Name:** DAXXIFY**Proper Name:** DAXIBOTULINUMTOXINA-LANM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

761127 / 0 2 9/7/2022 100 UNITS/VIAL (100 UNITS/VIAL)

POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL

Applicant/License No: SANDOZ INC / 2003**Trade Name:** OMNITROPE**Proper Name:** SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

021426 / 0 1 5/30/2006 5.8 MG/VIAL (5.8 MG/VIAL)

POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL

021426 / 0 3 1/16/2008 5 MG/1.5ML (3.33 MG/ML)

SOLUTION / SUBCUTANEOUS / CARTRIDGE

Applicant/License No: SANDOZ INC / 2003

021426 / 0 4 8/25/2008 10 MG/1.5 ML (6.67 MG/ML)

SOLUTION / SUBCUTANEOUS / CARTRIDGE

Applicant/License No: SANOFI AVENTIS US LLC / 1752**Trade Name:** ADMELOG**Proper Name:** INSULIN LISPRO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 209196 / 0 | 1 | 12/11/2017 | 1000 UNITS/10 ML (100 UNITS/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL |
| 209196 / 0 | 3 | 10/19/2018 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: ADMELOG SOLOSTAR**Proper Name:** INSULIN LISPRO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 209196 / 0 | 2 | 12/11/2017 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: APIDRA**Proper Name:** INSULIN GLULISINE RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021629 / 0 | 1 | 4/16/2004 | 1000 UNITS/10 ML (100 UNITS/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: APIDRA SOLOSTAR**Proper Name:** INSULIN GLULISINE RECOMBINANT

Applicant/License No: SANOFI AVENTIS US LLC / 1752

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021629 / 0 | 3 | 2/24/2009 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: ELITEK**Proper Name:** RASBURICASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|---------------|----------|---------------|---|
| 103946 / 0 | 1 | 7/12/2002 | 1.5 MG (1.5 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| 103946 / 5020 | 2 | 1/6/2006 | 7.5 MG (7.5 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: KEVZARA**Proper Name:** SARILUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761037 / 0 | 1 | 5/22/2017 | 150 MG/1.14 ML (131.58 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761037 / 0 | 2 | 5/22/2017 | 200 MG/1.14 ML (175.44 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761037 / 1 | 3 | 4/13/2018 | 150 MG/1.14 ML (131.58 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761037 / 1 | 4 | 4/13/2018 | 200 MG/1.14 ML (175.44 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: LANTUS

Applicant/License No: SANOFI AVENTIS US LLC / 1752**Proper Name:** INSULIN GLARGINE RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 021081 / 0 | 1 | 4/20/2000 | 1000 UNITS/10 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL |

Trade Name: LANTUS SOLOSTAR**Proper Name:** INSULIN GLARGINE RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 021081 / 0 | 2 | 4/27/2007 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: SARCLISA**Proper Name:** ISATUXIMAB-IRFC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761113 / 0 | 1 | 3/2/2020 | 100 MG/5 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761113 / 0 | 2 | 3/2/2020 | 500 MG/25 ML (20 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: SOLIQUA 100/33**Proper Name:** INSULIN GLARGINE; LIXISENATIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 208673 / 0 | 1 | 11/21/2016 | 300 UNITS/3 ML; 99MCG/3 ML (100 UNITS/ML; 33MCG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: TOUJEO MAX SOLOSTAR**Proper Name:** INSULIN GLARGINE RECOMBINANT

Applicant/License No: SANOFI AVENTIS US LLC / 1752

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 206538 / 0 | 2 | 3/26/2018 | 900 UNITS/3 ML (300 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: TOUJEO SOLOSTAR**Proper Name:** INSULIN GLARGINE RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 206538 / 0 | 1 | 2/25/2015 | 450 UNITS/1.5 ML (300 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: ZALTRAP**Proper Name:** ZIV-AFLIBERCEPT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125418 / 0 | 1 | 8/3/2012 | 100 MG/4 ML (25 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 125418 / 0 | 2 | 8/3/2012 | 200 MG/8 ML (25 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: SEAGEN INC / 2257**Trade Name:** ADCETRIS**Proper Name:** BRENTUXIMAB VEDOTIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125388 / 1 | 1 | 8/19/2011 | 50 MG (50 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: TIVDAK

Applicant/License No: SEAGEN INC / 2257**Proper Name:** TISOTUMAB VEDOTIN-TFTV

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761208 / 0 | 1 | 9/22/2021 | 40 MG/VIAL (40 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: SERVIER PHARMACEUTICALS LLC / 2125**Trade Name:** ASPARLAS**Proper Name:** CALASPARGASE PEGOL-MKNL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761102 / 0 | 1 | 12/20/2018 | 3750 UNITS/5 ML (750 UNITS/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: ONCASPAR**Proper Name:** PEGASPARGASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103411 / 0 | 1 | 2/1/1994 | 3750 IU/5 ML (750 IU/ML) SOLUTION / INTRAVENOUS, INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: SHIRE HUMAN GENETIC THERAPIES INC / 1593**Trade Name:** ELAPRASE**Proper Name:** IDURSULFASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125151 / 0 | 1 | 7/24/2006 | 6 MG/3 ML (2 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: VPRIV

Applicant/License No: SHIRE HUMAN GENETIC THERAPIES INC / 1593**Proper Name:** VELAGLUCERASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 022575 / 0 | 1 | 2/26/2010 | 400 UNITS/VIAL (400 UNITS/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: SMITH & NEPHEW INC / 2004**Trade Name:** REGRANEX**Proper Name:** BECAPLERMIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103691 / 0 | 1 | 12/16/1997 | 15 GM TUBE (100 UG/GM) GEL / TOPICAL / |

Trade Name: SANTYL**Proper Name:** COLLAGENASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 101995 / 0 | 1 | 6/4/1965 | 30 GM & 90 GM TUBE (250 U/GM) OINTMENT / TOPICAL / |

Applicant/License No: SOLSTICE NEUROSCIENCES LLC / 1718**Trade Name:** MYOBLOC**Proper Name:** RIMABOTULINUMTOXINB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103846 / 0 | 1 | 12/8/2002 | 2500 IU/0.5 ML (2500 IU/0.5 ML) SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: SOLSTICE NEUROSCIENCES LLC / 1718

| | | | |
|------------|---|-----------|---|
| 103846 / 0 | 2 | 12/8/2002 | 5000 IU/ML (5000 IU/ML) |
| | | | SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |

| | | | |
|------------|---|-----------|---|
| 103846 / 0 | 3 | 12/8/2002 | 10,000 IU/2 ML (5000 IU/ML) |
| | | | SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: SPECTRUM PHARMACEUTICALS INC / 2312**Trade Name:** ROLVEDON**Proper Name:** EFLAPEGRASTIM-XNST

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761148 / 0 | 1 | 9/9/2022 | 13.2 MG/0.6ML (13.2 MG/0.6 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: STEMLINE THERAPEUTICS INC / 2088**Trade Name:** ELZONRIS**Proper Name:** TAGRAXOFUSP-ERZS

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761116 / 0 | 1 | 12/21/2018 | 1000 MCG/ML (1000 MCG/ML) |
| | | | SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: SUN PHARMA GLOBAL FZE / 2092**Trade Name:** ILUMYA**Proper Name:** TILDRAKIZUMAB-ASMN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761067 / 0 | 1 | 3/20/2018 | 100 MG/ML (100 MG/ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: SWEDISH ORPHAN BIOVITRUM AB / 1859**Trade Name:** GAMIFANT**Proper Name:** EMAPALUMAB-LZSG

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761107 / 0 | 1 | 11/20/2018 | 10 MG/2 ML (5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761107 / 0 | 2 | 11/20/2018 | 50 MG/10 ML (5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |
| 761107 / 0 | 3 | 6/26/2020 | 100 MG/20 ML (5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: KEPIVANCE**Proper Name:** PALIFERMIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125103 / 0 | 1 | 12/15/2004 | 5.16 MG/VIAL (5.16 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: KINERET**Proper Name:** ANAKINRA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 103950 / 0 | 1 | 11/14/2001 | 100 MG/0.67 ML (100 MG/0.67 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: SYNAGIS**Proper Name:** PALIVIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

Applicant/License No: SWEDISH ORPHAN BIOVITRUM AB / 1859

| | | | |
|------------|---|-----------|---|
| 103770 / 0 | 1 | 6/19/1998 | 50 MG/0.5 ML (50 MG/0.5 ML) |
| | | | SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |
| 103770 / 0 | 2 | 6/19/1998 | 100 MG/ML (100 MG/ML) |
| | | | SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL |

Applicant/License No: TAKEDA PHARMACEUTICALS USA INC / 1898**Trade Name:** ENTYVIO**Proper Name:** VEDOLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 125476 / 0 | 1 | 5/20/2014 | 300 MG (300 MG/VIAL) |
| | | | POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |
| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
| 761133 / 0 | 2 | 9/27/2023 | 108 MG/0.68 ML (108 MG/0.68 ML) |
| | | | SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: ENTYVIO PEN**Proper Name:** VEDOLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761133 / 0 | 3 | 9/27/2023 | 108 MG/0.68 ML (108 MG/0.68 ML) |
| | | | SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: TERSERA THERAPEUTICS LLC / 2383**Trade Name:** MARGENZA**Proper Name:** MARGETUXIMAB-CMKB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761150 / 0 | 1 | 12/16/2020 | 250 MG/10 ML (25 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: TEVA PHARMACEUTICALS USA INC / 2016**Trade Name:** AJOVY**Proper Name:** FREMANEZUMAB-VFRM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761089 / 0 | 1 | 9/14/2018 | 225 MG/1.5 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761089 / 2 | 2 | 1/27/2020 | 225MG/1.5 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Applicant/License No: TEVA RESPIRATORY LLC / 2047**Trade Name:** CINQAIR**Proper Name:** RESLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761033 / 0 | 1 | 3/23/2016 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: TG THERAPEUTICS INC / 2090**Trade Name:** BRIUMVI**Proper Name:** UBLITUXIMAB-XIY

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|------------|--------------------------|
| 761238 / 0 | 1 | 12/28/2022 | 150 MG/6 ML (25 MG/ML) |
|------------|---|------------|--------------------------|

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: THERATECHNOLOGIES INC / 2091**Trade Name:** EGRIFTA SV**Proper Name:** TESAMORELIN ACETATE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|------------|---------------------------------|
| 022505 / 0 | 2 | 11/10/2010 | 2MG BASE/VIAL (2MG BASE/VIAL) |
|------------|---|------------|---------------------------------|

POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL

Trade Name: EGRIFTA WR**Proper Name:** TESAMORELIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|-------------------------------|
| 022505 / 0 | 3 | 3/25/2025 | 11.6 MG/VIAL (11.6 MG/VIAL) |
|------------|---|-----------|-------------------------------|

POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL

Trade Name: TROGARZO**Proper Name:** IBALIZUMAB-UIYK

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|----------|------------------------------|
| 761065 / 0 | 1 | 3/6/2018 | 200 MG/1.33 ML (150 MG/ML) |
|------------|---|----------|------------------------------|

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: UAB TEVA BALTICS / 1803**Trade Name:** GRANIX**Proper Name:** TBO-FILGRASTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|-------------|----------|---------------|--|
| 125294 / 0 | 1 | 8/29/2012 | 300 MCG/0.5 ML (300 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125294 / 0 | 2 | 8/29/2012 | 480 MCG/0.8 ML (480 MCG/0.8 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 125294 / 45 | 3 | 7/31/2018 | 300 MCG/ML (300 MCG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: UCB INC / 1736**Trade Name:** BIMZELX**Proper Name:** BIMEKIZUMAB-BKZX

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761151 / 0 | 1 | 10/17/2023 | 160 MG/ML (160 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |
| 761151 / 0 | 2 | 10/17/2023 | 160 MG/ML (160 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761151 / 0 | 3 | 10/11/2024 | 320 MG/2 ML (160 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |
| 761151 / 0 | 4 | 10/11/2024 | 320 MG/2 ML (160 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN |

Trade Name: CIMZIA

Applicant/License No: UCB INC / 1736**Proper Name:** CERTOLIZUMAB PEGOL

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|--------------|----------|---------------|--|
| 125160 / 0 | 1 | 4/22/2008 | 200 MG (200 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 125160 / 080 | 2 | 5/13/2009 | 200 MG/ML (200 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE |

Trade Name: RYSTIGGO**Proper Name:** ROZANOLIXIZUMAB-NOLI

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761286 / 0 | 1 | 6/26/2023 | 280 MG/2 ML (140 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761286 / 0 | 2 | 6/14/2024 | 420 MG/3 ML (140 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761286 / 0 | 3 | 6/14/2024 | 560 MG/4 ML (140 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |
| 761286 / 0 | 4 | 6/14/2024 | 840 MG/6 ML (140 MG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL |

Applicant/License No: ULTRAGENYX PHARMACEUTICAL INC / 2040**Trade Name:** MEPSEVII**Proper Name:** VESTRONIDASE ALFA-VJBK

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|------------|------------------------|
| 761047 / 0 | 1 | 11/15/2017 | 10 MG/5 ML (2 MG/ML) |
|------------|---|------------|------------------------|

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: UNITED THERAPEUTICS CORP / 1993**Trade Name:** UNITUXIN**Proper Name:** DINUTUXIMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|----------------------------|
| 125516 / 0 | 1 | 3/10/2015 | 17.5 MG/5 ML (3.5 MG/ML) |
|------------|---|-----------|----------------------------|

SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL

Applicant/License No: VALEANT PHARMACEUTICALS LUXEMBOURG SARL / 2053**Trade Name:** SILIQ**Proper Name:** BRODALUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|----------|----------|---------------|--|
|----------|----------|---------------|--|

| | | | |
|------------|---|-----------|-----------------------------|
| 761032 / 0 | 1 | 2/15/2017 | 210 MG/1.5 ML (140 MG/ML) |
|------------|---|-----------|-----------------------------|

SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE

Applicant/License No: VERICEL CORPORATION / 2010**Trade Name:** NEXOBRID**Proper Name:** ANACAULASE-BCDB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761192 / 0 | 2 | 12/28/2022 | 5 GM LYOPHILIZED POWDER/50 GM GEL (8.8% W/W (5 GM LYOPHILIZED POWDER/50 GM GEL)) GEL / TOPICAL / KIT |

Applicant/License No: VIFOR INTERNATIONAL AG / 2039**Trade Name:** MIRCERA**Proper Name:** METHOXYPOLYETHYLENE GLYCOL EPOETIN BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 125164 / 0 | 1 | 11/14/2007 | 50 MCG/0.3 ML (50 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 0 | 2 | 11/14/2007 | 75 MCG/0.3 ML (75 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 0 | 3 | 11/14/2007 | 100 MCG/0.3 ML (100 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 0 | 4 | 11/14/2007 | 150 MCG/0.3 ML (150 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 0 | 5 | 11/14/2007 | 200 MCG/0.3 ML (200 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 0 | 6 | 11/14/2007 | 250 MCG/0.3 ML (250 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: VIFOR INTERNATIONAL AG / 2039

| | | | |
|--------------|---|-----------|---|
| 125164 / 073 | 7 | 4/28/2016 | 30 MCG/0.3 ML (30 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 073 | 8 | 4/28/2016 | 120 MCG/0.3 ML (120 MCG/0.3 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |
| 125164 / 073 | 9 | 4/28/2016 | 360 MCG/0.6 ML (360 MCG/0.6 ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE |

Applicant/License No: VIOKACE LLC / 2196**Trade Name:** VIOKACE**Proper Name:** PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 022542 / 0 | 1 | 3/1/2012 | 39,150USP UNITS; 10,440USP UNITS; 39,150USP UNITS (39,150USP UNITS; 10,440USP UNITS; 39,150USP UNITS) TABLET / ORAL / |
| 022542 / 0 | 2 | 3/1/2012 | 78,300USP UNITS; 20,880USP UNITS; 78,300USP UNITS (78,300USP UNITS; 20,880USP UNITS; 78,300USP UNITS) TABLET / ORAL / |

Applicant/License No: VIVUS LLC / 2197**Trade Name:** PANCREAZE**Proper Name:** PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 022523 / 0 | 1 | 4/12/2010 | 24,600USP UNITS; 4,200USP UNITS; 14,200USP UNITS (24,600USP UNITS; 4,200USP UNITS; 14,200USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022523 / 0 | 2 | 4/12/2010 | 61,500USP UNITS; 10,500USP UNITS; 35,500USP UNITS (61,500USP UNITS; 10,500USP UNITS; 35,500USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |

Applicant/License No: VIVUS LLC / 2197

| | | | |
|------------|---|-----------|--|
| 022523 / 0 | 3 | 4/12/2010 | 98,400USP UNITS; 16,800USP UNITS; 56,800USP UNITS (98,400USP UNITS; 16,800USP UNITS; 56,800USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022523 / 0 | 4 | 4/12/2010 | 83,900USP UNITS; 21,000USP UNITS; 54,700USP UNITS (83,900USP UNITS; 21,000USP UNITS; 54,700USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022523 / 0 | 5 | 3/7/2014 | 10,850USP UNITS; 2,600USP UNITS; 6,200USP UNITS (10,850USP UNITS; 2,600USP UNITS; 6,200USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022523 / 0 | 6 | 4/26/2021 | 149,900USP UNITS; 37,000USP UNITS; 97,300USP UNITS (149,900USP UNITS; 37,000USP UNITS; 97,300USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |

Applicant/License No: WYETH PHARMACEUTICALS INC / 0003**Trade Name:** BESPONSA**Proper Name:** INOTUZUMAB OZOGAMICIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761040 / 0 | 1 | 8/17/2017 | 0.9 MG (0.9 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Trade Name: MYLOTARG**Proper Name:** GEMTUZUMAB OZOGAMICIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 761060 / 0 | 1 | 9/1/2017 | 4.5 MG (4.5 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: Y-MABS THERAPEUTICS INC / 2209**Trade Name:** DANYELZA**Proper Name:** NAXITAMAB-GQGK

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|--|
| 761171 / 0 | 1 | 11/25/2020 | 40 MG/10 ML (4 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL |

Applicant/License No: ZENPEP LLC / 2198**Trade Name:** ZENPEP**Proper Name:** PANCRELIPASE (AMYLASE;LIPASE;PROTEASE)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation |
|------------|----------|---------------|---|
| 022210 / 0 | 1 | 8/27/2009 | 24,000USP UNITS; 5,000USP UNITS; 17,000USP UNITS (24,000USP UNITS; 5,000USP UNITS; 17,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022210 / 0 | 2 | 8/27/2009 | 42,000USP UNITS; 10,000USP UNITS; 32,000USP UNITS (42,000USP UNITS; 10,000USP UNITS; 32,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022210 / 0 | 3 | 8/27/2009 | 63,000USP UNITS; 15,000USP UNITS; 47,000USP UNITS (63,000USP UNITS; 15,000USP UNITS; 47,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022210 / 0 | 4 | 8/27/2009 | 84,000USP UNITS; 20,000USP UNITS; 63,000USP UNITS (84,000USP UNITS; 20,000USP UNITS; 63,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022210 / 0 | 5 | 6/15/2011 | 14,000USP UNITS; 3,000USP UNITS; 10,000USP UNITS (14,000USP UNITS; 3,000USP UNITS; 10,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
| 022210 / 0 | 6 | 7/13/2011 | 105,000USP UNITS; 25,000USP UNITS; 79,000USP UNITS (105,000USP UNITS; 25,000USP UNITS; 79,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |

Applicant/License No: ZENPEP LLC / 2198

| | | | |
|------------|---|-----------|--|
| 022210 / 0 | 7 | 3/25/2014 | 168,000USP UNITS; 40,000USP UNITS; 126,000USP UNITS (168,000USP UNITS; 40,000USP UNITS; 126,000USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
|------------|---|-----------|--|

| | | | |
|------------|---|------------|--|
| 022210 / 0 | 8 | 10/27/2023 | 252,600USP UNITS; 60,000USP UNITS; 189,600USP UNITS (252,600USP UNITS; 60,000USP UNITS; 189,600USP UNITS) CAPSULE, DELAYED RELEASE / ORAL / |
|------------|---|------------|--|

Applicant/License No: ABBVIE INC / 1889

Trade Name: HUMIRA

Proper Name: ADALIMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 125057 / 0 | 1 | 12/31/2002 | 40 MG/0.8 ML (40 MG/0.8 ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL | 7/18/2022 |
| 125057 / 0 | 3 | 12/31/2002 | 10 MG/0.2 ML (10 MG/0.2 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 7/23/2020 |
| 125057 / 0 | 4 | 12/31/2002 | 20 MG/0.4 ML (20 MG/0.4 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 7/23/2020 |

Trade Name: SKYRIZI

Proper Name: RISANKIZUMAB-RZAA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761105 / 0 | 1 | 4/23/2019 | 75 MG/0.83ML (75 MG/0.83ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 6/7/2023 |

Applicant/License No: AKORN INC / 2173

Trade Name: HYDASE

Proper Name: HYALURONIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021716 / 0 | 1 | 10/25/2005 | 150 UNITS/ML (150 UNITS/ML) SOLUTION / INTERSTITIAL, INTRAOCULAR, RETROBULBAR, SOFT TISSUE, SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/5/2020 |

Applicant/License No: ALEXION PHARMACEUTICALS INC / 1743

Trade Name: ULTOMIRIS

Proper Name: RAVULIZUMAB-CWVZ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761108 / 0 | 1 | 12/21/2018 | 300 MG/30 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 5/30/2025 |
| 761108 / 0 | 4 | 7/22/2022 | 245 MG/3.5ML (70 MG/ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE | 6/3/2024 |

Applicant/License No: ALLERGAN INC / 1145

Trade Name: BOTOX

Proper Name: BOTULINUM TOXIN TYPE A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---------------|----------|---------------|---|------------------------------------|
| 103000 / 5101 | 3 | 4/14/2005 | 50 U (50 U/VIAL) POWDER / INTRAMUSCULAR, INTRADERMAL, INTRADETRUSOR / SINGLE-DOSE VIAL | 8/22/2022 |

Applicant/License No: AMGEN INC / 1080

Trade Name: AIMOVIG

Proper Name: ERENUMAB-AOOE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761077 / 0 | 1 | 5/17/2018 | 70 MG (70 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 9/26/2025 |
| 761077 / 1 | 3 | 3/11/2019 | 140 MG (140 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 9/26/2025 |

Trade Name: ARANESP

Proper Name: DARBEPOETIN ALFA (ALBUMIN SOLUTION)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|----------|----------|---------------|--|------------------------------------|
|----------|----------|---------------|--|------------------------------------|

Applicant/License No: AMGEN INC / 1080

Trade Name: ARANESP

| | | | | |
|--|----|-----------|-------------------------------------|-----------|
| 103951 / 0 | 16 | 9/17/2001 | 25 MCG/ML (25 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 17 | 9/17/2001 | 25 MCG/0.42 ML (25 MCG/0.42 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 18 | 9/17/2001 | 40 MCG/ML (40 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 19 | 9/17/2001 | 40 MCG/0.4 ML (40 MCG/0.4 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 20 | 9/17/2001 | 60 MCG/ML (60 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 21 | 9/17/2001 | 60 MCG/0.3 ML (60 MCG/0.3 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 22 | 9/17/2001 | 100 MCG/ML (100 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 23 | 9/17/2001 | 100 MCG/0.5 ML (100 MCG/0.5 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 24 | 9/17/2001 | 150 MCG/0.75 ML (150 MCG/0.75 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 25 | 9/17/2001 | 150 MCG/0.3 ML (150 MCG/0.3 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 26 | 9/17/2001 | 200 MCG/ML (200 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |

Applicant/License No: AMGEN INC / 1080

Trade Name: ARANESP

| | | | | |
|--|----|-----------|-----------------------------------|-----------|
| 103951 / 0 | 27 | 9/17/2001 | 200 MCG/0.4 ML (200 MCG/0.4 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 28 | 9/17/2001 | 300 MCG/ML (300 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 29 | 9/17/2001 | 300 MCG/0.6 ML (300 MCG/0.6 ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 103951 / 0 | 30 | 9/17/2001 | 500 MCG/ML (500 MCG/ML) | 9/27/2012 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |

Proper Name: DARBEPOETIN ALFA IN POLYSORBATE SOLUTION

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 103951 / 0 | 5 | 9/17/2001 | 150 MCG/0.75 ML (150 MCG/0.75 ML) | 12/20/2018 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103951 / 0 | 7 | 9/17/2001 | 300 MCG/ML (300 MCG/ML) | 4/30/2024 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |

Trade Name: REPATHA

Proper Name: EVOLOCUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|-------------------------------|----------|---------------|--|------------------------------------|
| 125522 / 1 | 2 | 7/11/2016 | 420 MG/3.5 ML (120 MG/ML) | 3/25/2024 |
| SOLUTION / SUBCUTANEOUS / KIT | | | | |

Trade Name: VECTIBIX

Proper Name: PANITUMUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 125147 / 0 | 2 | 9/27/2006 | 200 MG/10 ML (20 MG/ML) | 6/29/2017 |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | | |

Applicant/License No: AYTU BIOSCIENCES INC / 2035

Trade Name: PROSTASCINT

Proper Name: CAPROMAB PENDETIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103608 / 0 | 1 | 10/28/1996 | 0.5 MG/ML (0.5 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 5/16/2019 |

Applicant/License No: BAUSCH AND LOMB INC / 2180

Trade Name: VITRASE

Proper Name: HYALURONIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021640 / 0 | 1 | 5/5/2004 | 6200 UNITS/ML (6200 UNITS/ML) SOLUTION / INTERSTITIAL, INTRAMUSCULAR, INTRAOCULAR, PERIBULBAR, RETROBULBAR, SOFT TISSUE, SUBCUTANEOUS / SINGLE- DOSE VIAL | 9/23/2009 |
| 021640 / 0 | 2 | 12/2/2004 | 200 UNITS/ML (200 UNITS/ML) SOLUTION / INTERSTITIAL, INTRAMUSCULAR, INTRAOCULAR, PERIBULBAR, RETROBULBAR, SOFT TISSUE, SUBCUTANEOUS / SINGLE- DOSE VIAL | 9/15/2023 |

Applicant/License No: BAYER HEALTHCARE PHARMACEUTICALS INC / 1778

Trade Name: TRASYLOL

Proper Name: APROTININ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 020304 / 0 | 1 | 12/29/1993 | 1,000,000 KIU/100 ML (10000 KIU/ ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 5/25/2012 |
| 020304 / 0 | 2 | 12/29/1993 | 2,000,000 KIU/200 ML (10000 KIU/ ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 5/25/2012 |

Applicant/License No: BEL MAR LABORATORIES INC / 2182

Trade Name: CHORIONIC GONADOTROPIN

Proper Name: GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 017054 / 0 | 1 | 3/26/1974 | 5000 UNITS/VIAL (5000 UNITS/VIAL) / / | 1/1/1990 |
| 017054 / 0 | 2 | 3/26/1974 | 10000 UNITS/VIAL (10000 UNITS/VIAL) / / | 1/1/1990 |

Applicant/License No: BEONE MEDICINES USA INC / 2232

Trade Name: TEVIMBRA

Proper Name: TISLELIZUMAB-JSGR

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761417 / 0 | 1 | 12/26/2024 | 100 MG/10 ML (10 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 12/26/2024 |

Applicant/License No: BIOGEN INC / 1697

Trade Name: AVONEX

Proper Name: INTERFERON BETA 1A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103628 / 0 | 1 | 5/17/1996 | 30 MCG (30 MCG/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 10/30/2018 |

Applicant/License No: BRACCO DIAGNOSTICS INC / 2183

Trade Name: MACROTEC

Proper Name: TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 017833 / 0 | 1 | 2/23/1976 | 1.5 MG/10 MG/0.07 MG/0.19 MG/1.8 MG/VIAL (1.5 MG/10 MG/0.07 MG/0.19 MG/1.8 MG/VIAL) POWDER / INTRAVENOUS / | 2/5/2018 |

Applicant/License No: BRISTOL MYERS SQUIBB CO / 1713

Trade Name: OPDIVO QVANTIG

Proper Name: NIVOLUMAB AND HYALURONIDASE-NVHY

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761429 / 0 | 1 | 12/27/2024 | 600 MG AND 10,000 UNITS/5 ML (120 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL | 8/21/2025 |

Applicant/License No: DISCURE MEDICAL LLC / 2185

Trade Name: CHYMODIACTIN

Proper Name: CHYMOPAPAIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 018663 / 0 | 1 | 11/10/1982 | 10000 UNITS/VIAL (10000 UNITS/VIAL) POWDER / INTRADISCAL / MULTI-DOSE VIAL | 1/1/1900 |
| 018663 / 0 | 2 | 8/21/1984 | 4000 UNITS/VIAL (4000 UNITS/VIAL) POWDER / INTRADISCAL / MULTI-DOSE VIAL | 7/16/2002 |

Applicant/License No: EISAI INC / 1862

Trade Name: ONTAK

Proper Name: DENILEUKIN DIFTITOX

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103767 / 0 | 1 | 2/5/1999 | 300 MCG/2 ML (150 MCG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 1/30/2014 |

Applicant/License No: EKR THERAPEUTICS INC / 1814

Trade Name: RETAVASE

Proper Name: RETEPLASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103786 / 0 | 1 | 5/6/1998 | 10 U (10 U/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 7/2/2024 |

Applicant/License No: ELI LILLY AND CO / 1891

Trade Name: HUMALOG MIX 50/50

Proper Name: INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021018 / 0 | 1 | 12/22/1999 | 500 UNITS/10 ML; 500 UNITS/10 ML (50 UNITS/ML; 50 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL | 6/22/2023 |

Trade Name: HUMALOG MIX 50/50 PEN

Proper Name: INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021018 / 0 | 3 | 12/22/1999 | 150 UNITS/3 ML; 150 UNITS/3 ML (50 UNITS/ML; 50 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 6/22/2012 |

Trade Name: HUMALOG MIX 75/25 PEN

Proper Name: INSULIN LISPRO PROTAMINE RECOMBINANT; INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|----------|----------|---------------|--|------------------------------------|
| | | | | |

Applicant/License No: ELI LILLY AND CO / 1891

Trade Name: HUMALOG MIX 75/25 PEN

| | | | | |
|------------|---|------------|--|-----------|
| 021017 / 0 | 3 | 12/22/1999 | 226 UNITS/3 ML; 75 UNITS/3 ML (75 UNITS/ML; 25 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 6/22/2012 |
|------------|---|------------|--|-----------|

Trade Name: HUMALOG TEMPO PEN

Proper Name: INSULIN LISPRO RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 020563 / 0 | 5 | 11/15/2019 | 300 UNITS/3ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 10/16/2025 |

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 205747 / 0 | 2 | 5/7/2025 | 600 UNITS/3 ML (200 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 9/19/2025 |

Trade Name: HUMATROPE

Proper Name: SOMATROPIN RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 019640 / 0 | 1 | 6/23/1987 | 2 MG/VIAL (2 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 1/1/1900 |

| | | | | |
|------------|---|--|---|-----------|
| 019640 / 0 | 4 | | 5 MG/VIAL (5 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/27/2022 |
|------------|---|--|---|-----------|

Trade Name: LYUMJEV TEMPO PEN

Proper Name: INSULIN LISPRO-AABC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 761109 / 0 | 4 | 6/15/2020 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 10/16/2025 |

Trade Name: PORTRAZZA

Proper Name: NECITUMUMAB

Applicant/License No: ELI LILLY AND CO / 1891

Trade Name: PORTRAZZA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 125547 / 0 | 1 | 11/24/2015 | 800 MG/50 ML (16 MG/ML) SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | 9/25/2025 |

Trade Name: TRULICITY

Proper Name: DULAGLUTIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 125469 / 0 | 1 | 9/18/2014 | 0.75 MG/0.5 ML (0.75 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 6/28/2018 |
| 125469 / 0 | 2 | 9/18/2014 | 1.5 MG/0.5 ML (1.5 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 6/28/2018 |

Applicant/License No: EMD SERONO INC / 1773

Trade Name: GONAL-F

Proper Name: FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 020378 / 0 | 1 | 9/29/1997 | 75 IU/VIAL (NF) / / | 7/8/2004 |
| 020378 / 0 | 2 | 9/29/1997 | 150 IU/VIAL (NF) / / | 7/8/2004 |
| 020378 / 0 | 3 | 9/29/1997 | 37.5 IU/VIAL (NF) / / | 6/25/2002 |
| 021765 / 0 | 1 | 3/25/2004 | 37.5 IU/VIAL (NF) / / | 6/7/2004 |

Applicant/License No: EMD SERONO INC / 1773

Trade Name: GONAL-F

| | | | | |
|------------|---|-----------|--------------------|----------|
| 021765 / 0 | 3 | 3/25/2004 | 150 IU/VIAL (NF) | 6/7/2004 |
| / / | | | | |

Trade Name: GONAL-F RFF

Proper Name: FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021765 / 0 | 2 | 3/25/2004 | 75 IU/VIAL (75 IU/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 3/28/2025 |

Trade Name: GONAL-F RFF REDI-JECT

Proper Name: FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021684 / 0 | 4 | 11/25/2019 | 150 IU/0.25 ML (150 IU/0.25 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 2/27/2020 |

Trade Name: OVIDREL

Proper Name: CHORIOGONADOTROPIN ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021149 / 0 | 1 | 9/20/2000 | 0.25 MG/VIAL (0.25 MG/VIAL) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL | 6/7/2004 |

Trade Name: PERGONAL

Proper Name: MENOTROPINS (FSH;LH)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 017646 / 0 | 1 | 8/22/1975 | 75 IU/VIAL (75 IU/VIAL) POWDER / / | 2/1/2002 |
| 017646 / 0 | 2 | 8/22/1975 | 150 IU/VIAL (150 IU/VIAL) POWDER / / | 6/7/2004 |

Applicant/License No: EMD SERONO INC / 1773

Trade Name: SEROSTIM

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 020604 / 0 | 4 | 9/6/2001 | 8.8 MG/VIAL (8.8 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/14/2008 |

Trade Name: SEROSTIM LQ

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 020604 / 0 | 5 | 2/11/2005 | 6 MG/0.5 ML (6 MG/0.5 ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE | 5/14/2008 |

Trade Name: ZORBTIVE

Proper Name: SOMATROPIN RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021597 / 0 | 1 | 12/1/2003 | 4 MG/VIAL (4 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL | 6/7/2004 |
| 021597 / 0 | 2 | 12/1/2003 | 5 MG/VIAL (5 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL | 6/7/2004 |
| 021597 / 0 | 3 | 12/1/2003 | 6 MG/VIAL (6 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL | 6/7/2004 |
| 021597 / 0 | 4 | 12/1/2003 | 8.8 MG/VIAL (8.8 MG/VIAL) POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL | 9/30/2021 |

Applicant/License No: ENDO GLOBAL AESTHETICS LTD / 2136

Trade Name: QWO

Proper Name: COLLAGENASE CLOSTRIDIUM HISTOLYTICUM-AAES

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761146 / 0 | 1 | 7/6/2020 | 0.92 MG (0.92 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 12/15/2022 |
| 761146 / 0 | 2 | 7/6/2020 | 1.84 MG (1.84 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 12/15/2022 |

Applicant/License No: FERRING PHARMACEUTICALS INC / 2112

Trade Name: A.P.L.

Proper Name: GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 017055 / 0 | 1 | 12/13/1974 | 5000 UNITS/VIAL (5000 UNITS/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 6/23/2003 |
| 017055 / 0 | 2 | 12/13/1974 | 10000 UNITS/VIAL (10000 UNITS/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 6/23/2003 |
| 017055 / 0 | 3 | 12/13/1974 | 20000 UNITS/VIAL (20000 UNITS/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 6/20/2002 |

Trade Name: ACTHREL

Proper Name: CORTICORELIN OVINE TRIFLUTATE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 020162 / 0 | 1 | 5/23/1996 | 100 MCG/VIAL (100MCG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 5/18/2021 |

Trade Name: BIO-TROPIN

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|----------|----------|---------------|--|------------------------------------|
|----------|----------|---------------|--|------------------------------------|

Applicant/License No: FERRING PHARMACEUTICALS INC / 2112

Trade Name: BIO-TROPIN

| | | | | |
|---|---|-----------|-----------------------------|-----------|
| 019774 / 0 | 1 | 5/25/1995 | 4.8 MG/VIAL (4.8 MG/VIAL) | 3/20/2003 |
| POWDER / SUBCUTANEOUS / MULTI-DOSE VIAL | | | | |

Trade Name: NOVAREL

Proper Name: GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 017016 / 0 | 4 | 1/15/1974 | 20000 UNITS/VIAL (20000 UNITS/VIAL) POWDER / / | 10/12/1994 |
| 017016 / 0 | 7 | 1/15/1974 | 10000 UNITS/VIAL (10000 UNITS/VIAL) POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL | 5/16/2023 |
| 017016 / 0 | 9 | 12/27/1984 | 2000 UNITS/VIAL (2000 UNITS/VIAL) POWDER / / | 1/1/1900 |
| 017016 / 0 | 10 | 1/15/1974 | 15000 UNITS/VIAL (15000 UNITS/VIAL) POWDER / / | 10/12/1994 |
| 017016 / 0 | 11 | 2/16/1990 | 2000 UNITS/VIAL (2000 UNITS/VIAL) POWDER / / | 6/28/2002 |

Trade Name: REPRONEX

Proper Name: MENOTROPINS (FSH;LH)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021047 / 0 | 1 | 8/27/1999 | 75 IU/VIAL (75 IU/VIAL) POWDER / INTRAMUSCULAR, SUBCUTANEOUS / SINGLE-DOSE VIAL | 2/1/2002 |
| 021047 / 0 | 2 | 5/20/1985 | 150 IU/VIAL (150 IU/VIAL) POWDER / INTRAMUSCULAR, SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/6/2003 |

Applicant/License No: FRESENIUS KABI USA LLC / 2146

Trade Name: CHORIONIC GONADOTROPIN

Proper Name: GONADOTROPIN, CHORIONIC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 017067 / 0 | 1 | 3/5/1973 | 5000 UNITS/VIAL (5000 UNITS/VIAL) POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL | 6/23/1994 |
| 017067 / 0 | 3 | 3/5/1973 | 15000 UNITS/VIAL (15000 UNITS/VIAL) POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL | 1/1/1900 |
| 017067 / 0 | 4 | 3/5/1973 | 20000 UNITS/VIAL (20000 UNITS/VIAL) POWDER / INTRAMUSCULAR / MULTI-DOSE VIAL | 6/23/1994 |

Applicant/License No: GALDERMA LABORATORIES LP / 2289

Trade Name: NEMLUVIO

Proper Name: NEMOLIZUMAB-ILTO

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761391 / 0 | 1 | 12/13/2024 | 30 MG/0.49 ML (30 MG/0.49 ML) POWDER & SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 6/26/2025 |

Applicant/License No: GENENTECH INC / 1048

Trade Name: ACTIVASE

Proper Name: ALTEPLASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103172 / 0 | 3 | 3/2/1992 | 100 MG (100 MG/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 5/14/2026 |

Trade Name: HERCEPTIN

Proper Name: TRASTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|----------|----------|---------------|--|------------------------------------|
|----------|----------|---------------|--|------------------------------------|

Applicant/License No: GENENTECH INC / 1048

Trade Name: HERCEPTIN

| | | | | |
|--|---|-----------|------------------------|----------|
| 103792 / 0 | 1 | 9/25/1998 | 420 MG (420 MG/VIAL) | 2/3/2021 |
| POWDER / INTRAVENOUS / MULTI-DOSE VIAL | | | | |

Trade Name: XOLAIR

Proper Name: OMALIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|--|----------|---------------|--|------------------------------------|
| 103976 / 0 | 2 | 6/20/2003 | 125 MG/VIAL (125 MG/VIAL) | 8/25/2016 |
| POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |

Applicant/License No: GENZYME CORP / 1596

Trade Name: CAMPATH

Proper Name: ALEMTUZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 103948 / 0 | 3 | 5/7/2001 | 10 MG/VIAL (10 MG/VIAL) | 7/6/2011 |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | | |

Trade Name: CEREZYME

Proper Name: IMIGLUCERASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 020367 / 0 | 1 | 5/23/1994 | 200 UNITS/VIAL (200 UNITS/VIAL) | 7/30/2021 |
| POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | | | | |

Applicant/License No: HEMISPHERX BIOPHARMA INC / 1703

Trade Name: ALFERON N INJECTION

Proper Name: INTERFERON ALFA-N3 (HUMAN LEUKOCYTE DERIVED)

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|--|----------|---------------|--|------------------------------------|
| 103158 / 0 | 1 | 10/10/1989 | 5 MIU/VIAL (5 MIU/VIAL) | 6/27/2013 |
| SOLUTION / INTRALESIONAL / MULTI-DOSE VIAL | | | | |

Applicant/License No: ISO TEX DIAGNOSTICS INC / 2189

Trade Name: MEGATOPE

Proper Name: ALBUMIN IODINATED I-131 SERUM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 017837 / 0 | 1 | 2/23/1976 | 0.5mCi/VIAL (0.5mCi/VIAL) SOLUTION / INTRAVENOUS / MULTI-DOSE VIAL | 9/29/2022 |
| 017837 / 0 | 3 | 2/23/1976 | 2mCi/VIAL (2mCi/VIAL) SOLUTION / INTRAVENOUS / MULTI-DOSE VIAL | 1/1/1900 |
| 017837 / 0 | 4 | 2/23/1976 | 5uCi/AMP (5uCi/AMP) SOLUTION / INTRAVENOUS / MULTI-DOSE VIAL | 1/1/1900 |
| 017837 / 0 | 5 | 2/23/1976 | 20uCi/AMP (20uCi/AMP) SOLUTION / INTRAVENOUS / MULTI-DOSE VIAL | 6/7/2002 |

Applicant/License No: JANSSEN BIOTECH INC / 1864

Trade Name: RYBREVANT FASPRO

Proper Name: AMIVANTAMAB AND HYALURONIDASE-LPUJ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761484 / 0 | 1 | 2/13/2026 | 2,400 MG AND 30,000 UNITS/15 ML (160 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/13/2026 |
| 761484 / 0 | 2 | 2/13/2026 | 3,520 MG AND 44,000 UNITS/22 ML (160 MG AND 2,000 UNITS/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/13/2026 |

Applicant/License No: JUBILANT DRAXIMAGE INC / 2186

Trade Name: PULMOLITE

Proper Name: TECHNETIUM TC-99M ALBUMIN AGGREGATED KIT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 017776 / 0 | 1 | 11/16/1976 | 1 MG/10 MG/0.24 MG/0.7 MG0.13 MG/10 MG/VIAL (1 MG/10 MG/0.24 MG/0.7 MG0.13 POWDER / INTRAVENOUS / | 6/25/2002 |

Applicant/License No: LEADIANT BIOSCIENCES INC / 2073

Trade Name: ADAGEN

Proper Name: PEGADEMASE BOVINE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 019818 / 0 | 1 | 3/21/1990 | 375 UNITS/1.5 ML (250 UNITS/ML) SOLUTION / INTRAMUSCULAR / SINGLE-DOSE VIAL | 10/31/2019 |

Applicant/License No: MERCK AND CO INC DIV MERCK SHARP AND DOHME / 0002

Trade Name: INTRON A

Proper Name: INTERFERON ALFA-2B

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103132 / 0 | 1 | 6/4/1986 | 10 MIU (10 MIU/VIAL) POWDER / INTRAMUSCULAR, SUBCUTANEOUS, INTRALESIONAL, INTRAVENOUS / SINGLE-DOSE VIAL | 3/26/2021 |
| 103132 / 0 | 2 | 6/4/1986 | 18 MIU (18 MIU/VIAL) POWDER / INTRAMUSCULAR, SUBCUTANEOUS, INTRAVENOUS / SINGLE- DOSE VIAL | 3/26/2021 |
| 103132 / 0 | 3 | 6/4/1986 | 50 MIU (50 MIU/VIAL) POWDER / INTRAMUSCULAR, SUBCUTANEOUS, INTRAVENOUS / SINGLE- DOSE VIAL | 3/26/2021 |
| 103132 / 0 | 4 | 6/4/1986 | 22.8 MIU/3.8 ML (6 MIU/ML) SOLUTION / INTRAMUSCULAR, SUBCUTANEOUS / MULTI-DOSE VIAL | 3/26/2021 |

Applicant/License No: MERCK AND CO INC DIV MERCK SHARP AND DOHME / 0002

Trade Name: INTRON A

| | | | | |
|---|---|----------|-------------------------------------|-----------|
| 103132 / 0 | 5 | 6/4/1986 | 32 MIU/3.2 ML (10 MIU/ML) | 3/26/2021 |
| SOLUTION / INTRAMUSCULAR, SUBCUTANEOUS, INTRALESIONAL / MULTI-DOSE VIAL | | | | |
| 103132 / 0 | 6 | 6/4/1986 | 10 MIU/ML (10 MIU/ML) | 9/23/2013 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 103132 / 0 | 7 | 6/4/1986 | 22.5 MIU/1.5 ML (22.5 MIU/1.5 ML) | 9/23/2013 |
| SOLUTION / SUBCUTANEOUS / PREFILLED PEN | | | | |
| 103132 / 0 | 8 | 6/4/1986 | 37.5 MIU/1.5 ML (37.5 MIU/1.5 ML) | 9/23/2013 |
| SOLUTION / SUBCUTANEOUS / PREFILLED PEN | | | | |
| 103132 / 0 | 9 | 6/4/1986 | 75 MIU/1.5 ML (75 MIU/1.5 ML) | 9/23/2013 |
| SOLUTION / SUBCUTANEOUS / PREFILLED PEN | | | | |

Trade Name: KEYTRUDA

Proper Name: PEMBROLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 125514 / 0 | 1 | 9/4/2014 | 50 MG (50 MG/VIAL) | 6/13/2018 |
| POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | | | | |

Trade Name: ZINPLAVA

Proper Name: BEZLOTOXUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 761046 / 0 | 1 | 10/21/2016 | 1000 MG/40 ML (25 MG/ML) | 8/9/2024 |
| SOLUTION / INTRAVENOUS / SINGLE-DOSE VIAL | | | | |

Applicant/License No: MICROBIX BIOSYSTEMS INC / 2191

Trade Name: KINLYTIC

Proper Name: UROKINASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021846 / 0 | 1 | 1/16/1978 | 250000 IU/VIAL (250000 IU/VIAL) POWDER / INTRAVENOUS / MULTI-DOSE VIAL | 8/25/2009 |
| 021846 / 0 | 2 | 1/16/1978 | 9000 IU/VIAL (9000 IU/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 8/25/2009 |
| 021846 / 0 | 3 | 1/16/1978 | 5000 IU/VIAL (5000 IU/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 6/30/2003 |

Applicant/License No: NOVARTIS PHARMACEUTICALS CORP / 1244

Trade Name: ILARIS

Proper Name: CANAKINUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|-------------|----------|---------------|--|------------------------------------|
| 125319 / 0 | 1 | 6/17/2009 | 180 MG/VIAL (180 MG/VIAL) / / | 5/25/2018 |
| 125319 / 88 | 2 | 12/22/2016 | 150 MG (150 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 5/25/2018 |

Applicant/License No: NOVO NORDISK INC / 1261

Trade Name: ALHEMO

Proper Name: CONCIZUMAB-MTCI

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 761428 / 0 | 1 | 7/31/2025 | 60 MG/1.5 ML (40 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 7/31/2025 |

Applicant/License No: NOVO NORDISK INC / 1261

Trade Name: ALHEMO

| | | | | |
|--|---|-----------|-----------------------------|-----------|
| 761428 / 0 | 2 | 7/31/2025 | 150 MG/1.5 ML (100 MG/ML) | 7/31/2025 |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | | |

| | | | | |
|--|---|-----------|---------------------------|-----------|
| 761428 / 0 | 3 | 7/31/2025 | 300 MG/3 ML (100 MG/ML) | 7/31/2025 |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | | |

Trade Name: LEVEMIR

Proper Name: INSULIN DETEMIR RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 021536 / 0 | 1 | 6/16/2005 | 1000 UNITS/10 ML (100 UNITS/ML) | 12/31/2024 |
| SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL | | | | |

Trade Name: LEVEMIR FLEXPEN

Proper Name: INSULIN DETEMIR RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|--|----------|---------------|--|------------------------------------|
| 021536 / 0 | 2 | 6/16/2005 | 300 UNITS/3 ML (100 UNITS/ML) | 4/1/2024 |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | | |

Trade Name: LEVEMIR FLEXTOUCH

Proper Name: INSULIN DETEMIR RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|--|----------|---------------|--|------------------------------------|
| 021536 / 0 | 5 | 10/31/2013 | 300 UNITS/3 ML (100 UNITS/ML) | 1/26/2023 |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | | |

Trade Name: LEVEMIR INNOLET

Proper Name: INSULIN DETEMIR RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|--|----------|---------------|--|------------------------------------|
| 021536 / 0 | 3 | 6/16/2005 | 300 UNITS/3 ML (100 UNITS/ML) | 12/12/2012 |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | | |

Applicant/License No: NOVO NORDISK INC / 1261

Trade Name: LEVEMIR PENFILL

Proper Name: INSULIN DETEMIR RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021536 / 0 | 4 | 6/16/2005 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE | 12/12/2012 |

Trade Name: NORDITROPIN

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021148 / 0 | 1 | 6/20/2000 | 5MG/1.5ML (3.33MG/ML) SOLUTION / SUBCUTANEOUS / | 8/10/2012 |
| 021148 / 0 | 2 | 6/20/2000 | 10MG/1.5ML (6.67MG/ML) SOLUTION / SUBCUTANEOUS / | 8/10/2012 |
| 021148 / 0 | 3 | 6/20/2000 | 15MG/1.5ML (10MG/ML) SOLUTION / SUBCUTANEOUS / | 9/25/2012 |

Trade Name: NORDITROPIN NORDIFLEX

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021148 / 0 | 4 | 10/1/2004 | 5MG/1.5ML (3.33MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 9/25/2012 |
| 021148 / 0 | 5 | 10/1/2004 | 10MG/1.5ML (6.67MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 9/25/2012 |
| 021148 / 0 | 6 | 10/1/2004 | 15MG/1.5ML (10MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 7/21/2015 |
| 021148 / 0 | 7 | 3/10/2009 | 30MG/3ML (10MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 7/21/2015 |

Applicant/License No: NOVO NORDISK INC / 1261

Trade Name: NOVOLOG FLEXTOUCH

Proper Name: INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 020986 / 0 | 5 | 10/31/2013 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 1/17/2018 |

Trade Name: NOVOLOG INNOLET

Proper Name: INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 020986 / 0 | 4 | 4/23/2004 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 12/12/2012 |

Trade Name: NOVOLOG MIX 70/30 PENFILL

Proper Name: INSULIN ASPART PROTAMINE RECOMBINANT; INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021172 / 0 | 2 | 11/1/2001 | 210 UNITS/3ML; 90 UNITS/3 ML (70 UNITS/ML; 30 UNITS/ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE | 12/12/2012 |

Trade Name: NOVOLOG MIX 70/30 PFS

Proper Name: INSULIN ASPART PROTAMINE RECOMBINANT; INSULIN ASPART RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021172 / 0 | 3 | 11/1/2001 | 210 UNITS/3 ML; 90 UNITS/3 ML (70 UNITS/ML; 30 UNITS/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 12/12/2012 |

Trade Name: RYZODEG 70/30

Proper Name: INSULIN ASPART; INSULIN DEGLUDEC

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 203313 / 0 | 1 | 9/25/2015 | 90 UNITS/3 ML; 210 UNITS/3 ML (30 UNITS/ML; 70 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 1/12/2018 |

Applicant/License No: ORGANON USA LLC A SUBSIDIARY OF ORGANON AND CO / 2331

Trade Name: FOLLISTIM AQ

Proper Name: FOLLITROPIN ALFA/BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021211 / 0 | 3 | 3/23/2004 | 150 IU/0.18 ML (150 IU/0.18 ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE | 6/27/2006 |

Applicant/License No: PARTNER THERAPEUTICS INC / 2087

Trade Name: LEUKINE

Proper Name: SARGRAMOSTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 103362 / 0 | 2 | 3/5/1991 | 500 MCG/ML (500 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / MULTI-DOSE VIAL | 6/15/2018 |

Applicant/License No: PFIZER INC / 2001

Trade Name: HYMPAVZI

Proper Name: MARSTACIMAB-HNCQ

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761369 / 0 | 1 | 10/11/2024 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 6/25/2025 |

Applicant/License No: PHARMAAND GMBH / 2291

Trade Name: PEGASYS

Proper Name: PEGINTERFERON ALFA 2A

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---------------|----------|---------------|---|------------------------------------|
| 103964 / 5204 | 3 | 9/29/2011 | 135 MCG/0.5 ML (135 MCG/0.5 ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 7/28/2020 |

Applicant/License No: PHARMAAND GMBH / 2291

Trade Name: PEGASYS

| | | | | |
|--|---|-----------|-----------------------------------|-----------|
| 103964 / 5204 | 4 | 9/29/2011 | 180 MCG/0.5 ML (180 MCG/0.5 ML) | 7/28/2020 |
| SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | | | | |

Applicant/License No: PHARMACIA AND UPJOHN CO / 1216

Trade Name: GENOTROPIN PRESERVATIVE FREE

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|---|----------|---------------|--|------------------------------------|
| 020280 / 0 | 4 | 8/24/1995 | 1.5 MG/VIAL (1.5 MG/VIAL) | 6/26/2018 |
| POWDER / SUBCUTANEOUS / PREFILLED SYRINGE | | | | |

Applicant/License No: QOL MEDICAL LLC / 2195

Trade Name: SUCRAID

Proper Name: SACROSIDASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|-----------------------------------|----------|---------------|--|------------------------------------|
| 020772 / 0 | 1 | 4/9/1998 | 1003000 IU/118 ML (8500 IU/ML) | 10/7/2024 |
| SOLUTION / ORAL / MULTI-DOSE VIAL | | | | |

Applicant/License No: RECORDATI RARE DISEASES INC / 1899

Trade Name: ELSPAR

Proper Name: ASPARAGINASE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|--|----------|---------------|--|------------------------------------|
| 101063 / 0 | 1 | 1/10/1978 | 10000 IU (10000 IU/VIAL) | 4/9/2014 |
| POWDER / INTRAVENOUS, INTRAMUSCULAR / SINGLE-DOSE VIAL | | | | |

Applicant/License No: REGENERON PHARMACEUTICALS INC / 1760

Trade Name: DUPIXENT

Proper Name: DUPILUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761055 / 0 | 5 | 10/20/2021 | 100 MG/0.67 ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 4/12/2024 |

Trade Name: PRALUENT

Proper Name: ALIROCUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 125559 / 0 | 1 | 7/24/2015 | 75 MG/ML (75 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 9/25/2020 |
| 125559 / 0 | 2 | 7/24/2015 | 150 MG/ML (150 MG/ML) SOLUTION / SUBCUTANEOUS / PREFILLED SYRINGE | 9/25/2020 |

Applicant/License No: REVANCE THERAPEUTICS INC / 2101

Trade Name: DAXXIFY

Proper Name: DAXIBOTULINUMTOXINA-LANM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761127 / 0 | 1 | 9/7/2022 | 50 UNITS/VIAL (50 UNITS/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 5/30/2025 |

Applicant/License No: SANDOZ INC / 2003

Trade Name: OMNITROPE

Proper Name: SOMATROPIN

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021426 / 0 | 2 | 5/30/2006 | 1.5 MG/VIAL (1.5 MG/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 7/19/2021 |

Applicant/License No: SANOFI AVENTIS US LLC / 1752

Trade Name: ADLYXIN

Proper Name: LIXISENATIDE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 208471 / 0 | 1 | 7/27/2016 | 0.15MG/3ML (0.05MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 2/22/2023 |
| 208471 / 0 | 2 | 7/27/2016 | 0.3MG/3ML (0.1MG/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 2/22/2023 |

Trade Name: APIDRA

Proper Name: INSULIN GLULISINE RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 021629 / 0 | 2 | 12/20/2005 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / AUTOINJECTOR-PREFILLED PEN | 9/25/2018 |

Trade Name: LANTUS

Proper Name: INSULIN GLARGINE RECOMBINANT

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 021081 / 0 | 3 | 4/20/2000 | 500 UNITS/5 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / MULTI-DOSE VIAL | 4/20/2000 |
| 021081 / 0 | 4 | 4/20/2000 | 300 UNITS/3 ML (100 UNITS/ML) SOLUTION / SUBCUTANEOUS / CARTRIDGE | 3/31/2011 |

Applicant/License No: SHIRE HUMAN GENETIC THERAPIES INC / 1593

Trade Name: VPRIV

Proper Name: VELAGLUCERASE ALFA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 022575 / 0 | 2 | 2/26/2010 | 200 UNITS/VIAL (200 UNITS/VIAL) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 10/6/2011 |

Applicant/License No: SWEDISH ORPHAN BIOVITRUM AB / 1859

Trade Name: SYNAGIS

Proper Name: PALIVIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 103770 / 0 | 3 | 6/19/1998 | 50 MG (50 MG/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 2/28/2007 |
| 103770 / 0 | 4 | 6/19/1998 | 100 MG (100 MG/VIAL) POWDER / INTRAMUSCULAR / SINGLE-DOSE VIAL | 2/28/2007 |

Applicant/License No: TAKEDA PHARMACEUTICALS USA INC / 1898

Trade Name: ENTYVIO

Proper Name: VEDOLIZUMAB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761133 / 0 | 1 | 9/27/2023 | 300 MG (300 MG) POWDER / INTRAVENOUS / SINGLE-DOSE VIAL | 5/15/2024 |

Applicant/License No: THERATECHNOLOGIES INC / 2091

Trade Name: EGRIFTA SV

Proper Name: TESAMORELIN ACETATE

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|---|------------------------------------|
| 022505 / 0 | 1 | 11/10/2010 | 1MG BASE/VIAL (1MG BASE/VIAL) POWDER / SUBCUTANEOUS / SINGLE-DOSE VIAL | 7/31/2020 |

Applicant/License No: UAB TEVA BALTICS / 1803

Trade Name: GRANIX

Proper Name: TBO-FILGRASTIM

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|-------------|----------|---------------|--|------------------------------------|
| 125294 / 45 | 4 | 7/31/2018 | 480MCG/1.6 ML (300 MCG/ML) SOLUTION / SUBCUTANEOUS / SINGLE-DOSE VIAL | 7/25/2025 |

Applicant/License No: VERICEL CORPORATION / 2010

Trade Name: NEXOBRID

Proper Name: ANACAULASE-BCDB

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 761192 / 0 | 1 | 12/28/2022 | 2 GM LYOPHILIZED POWDER/20 GM GEL (8.8% W/W (2 GM LYOPHILIZED POWDER/20 GEL / TOPICAL / KIT | 11/26/2024 |

Applicant/License No: VIFOR INTERNATIONAL AG / 2039

Trade Name: MIRCERA

Proper Name: METHOXYPOLYETHYLENE GLYCOL EPOETIN BETA

| BLA#/STN | Product# | Approval Date | Total Drug Content (Concentration) Dosage Form/Route/Presentation | Date added to Discontinued List |
|------------|----------|---------------|--|------------------------------------|
| 125164 / 0 | 10 | 11/14/2007 | 50 MCG/ML (50 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | 9/24/2008 |
| 125164 / 0 | 11 | 11/14/2007 | 100 MCG/ML (100 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | 9/24/2008 |
| 125164 / 0 | 12 | 11/14/2007 | 200 MCG/ML (200 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | 9/24/2008 |
| 125164 / 0 | 13 | 11/14/2007 | 300 MCG/ML (300 MCG/ML) SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | 9/24/2008 |

Applicant/License No: VIFOR INTERNATIONAL AG / 2039

Trade Name: MIRCERA

| | | | | |
|--|----|------------|-----------------------------------|-----------|
| 125164 / 0 | 14 | 11/14/2007 | 400 MCG/ML (400 MCG/ML) | 9/24/2008 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 125164 / 0 | 15 | 11/14/2007 | 400 MCG/0.6 ML (400 MCG/0.6 ML) | 9/24/2008 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 125164 / 0 | 16 | 11/14/2007 | 600 MCG/ML (600 MCG/ML) | 9/24/2008 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |
| 125164 / 0 | 17 | 11/14/2007 | 600 MCG/0.6 ML (600 MCG/0.6 ML) | 9/24/2008 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 125164 / 0 | 18 | 11/14/2007 | 800 MCG/0.6 ML (800 MCG/0.6 ML) | 9/24/2008 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / PREFILLED SYRINGE | | | | |
| 125164 / 0 | 19 | 11/14/2007 | 1000 MCG/ML (1000 MCG/ML) | 9/24/2008 |
| SOLUTION / INTRAVENOUS, SUBCUTANEOUS / SINGLE-DOSE VIAL | | | | |