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Introducing PHARMAC

The Pharmaceutical Management Agency (PHARMAC) makes decisions that help control Government spending on pharmaceuticals. This includes community pharmaceuticals, hospital pharmaceuticals, vaccines and increasingly, hospital medical devices. PHARMAC negotiates prices, sets subsidy levels and conditions, and makes decisions on changes to the subsidised list. The funding for pharmaceuticals comes from District Health Boards.

PHARMAC's role:

"Secure for eligible people in need of pharmaceuticals, the best health outcomes that can reasonably be achieved, and from within the amount of funding provided."

New Zealand Public Health and Disability Act 2000

To ensure our decisions are as fair and robust as possible we use a decision-making process that incorporates clinical, economic and commercial issues. We also seek the views of users and the wider community through consultation. The processes we generally use are outlined in our Operating Policies and Procedures.

Further information about PHARMAC and the way we make funding decisions can be found on the PHARMAC website at https://www.pharmac.govt.nz/about.

Glossary

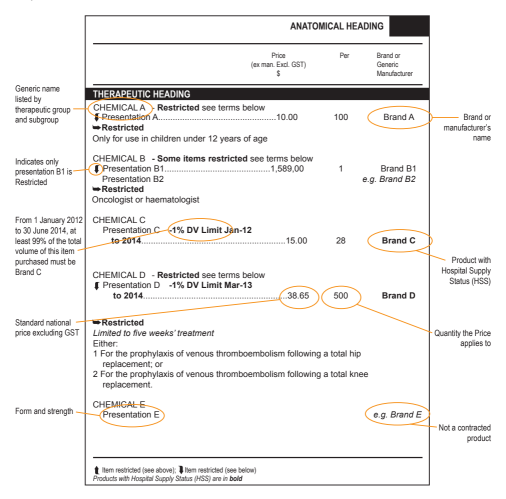
Units of Measure

| gramg kilogramkg international unitiu | microgrammcg milligrammg millilitreml | |
|---|--|--------------------------------|
| Abbreviations | | |
| applicationapp capsulecap creamcrm dispersibledisp effervescenteff emulsioneff | enteric coatedEC granulesgrans injectioninj liquidliq lotionlotn ointmentoint | suppositorysuppos tablettab |

HSS Hospital Supply Status

Guide to Section H listings

Example



General Rules for Section H of the Pharmaceutical Schedule are included in Section A.

Read the General Rules : https://www.pharmac.govt.nz/section-a.

PART II: ALIMENTARY TRACT AND METABOLISM

| | Price (ex man. excl. (\$ | GST) Per | Brand or Generic Manufacturer |
|--|---------------------------------|----------------|-------------------------------------|
| Antacids and Antiflatulents | | | |
| Antacids and Reflux Barrier Agents | | | |
| ALUMINIUM HYDROXIDE WITH MAGNESIUM HYDROXIDE AND S Tab 200 mg with magnesium hydroxide 200 mg and simeticone 2 Oral liq 400 mg with magnesium hydroxide 400 mg and simeticon 30 mg per 5 ml | 20 mg | | e.g. Mylanta e.g. Mylanta Double |
| SIMETICONE Oral drops 100 mg per ml Oral drops 20 mg per 0.3 ml Oral drops 40 mg per ml | | | Strength |
| SODIUM ALGINATE WITH MAGNESIUM ALGINATE Powder for oral soln 225 mg with magnesium alginate 87.5 mg, s SODIUM ALGINATE WITH SODIUM BICARBONATE AND CALCIUN Tab 500 mg with sodium bicarbonate 267 mg and calcium carbon | I CARBONATE | | e.g. Gaviscon Infant |
| 160 mg | hanata | | e.g. Gaviscon Double Strength |
| Oral liq 500 mg with sodium bicarbonate 267 mg and calcium ca 160 mg per 10 ml SODIUM CITRATE Oral liq 8.8% (300 mmol/l) | | 500 ml | Acidex |
| Phosphate Binding Agents | | | |
| ALUMINIUM HYDROXIDE Tab 600 mg | | | |
| CALCIUM CARBONATE – Restricted see terms below ↓ Oral liq 250 mg per ml (100 mg elemental per ml) | | 500 ml | Roxane |
| Initiation Only when prescribed for patients unable to swallow calcium carbona inappropriate | te tablets or where | e calcium carb | onate tablets are |
| Antidiarrhoeals and Intestinal Anti-Inflammatory A | gents | | |
| Antipropulsives | | | |
| DIPHENOXYLATE HYDROCHLORIDE WITH ATROPINE SULPHAT Tab 2.5 mg with atropine sulphate 25 mcg LOPERAMIDE HYDROCHLORIDE | E | | |
| Tab 2 mg Cap 2 mg – 1% DV Oct-19 to 2022 | | 400 400 | Nodia Diamide Relief |
| Rectal and Colonic Anti-Inflammatories | | | |
| BUDESONIDE – Restricted see terms on the next page Cap 3 mg | | | |

| Pi | rice | | Brand or |
|----------|-----------|-----|--------------|
| (ex man. | excl. GST | | Generic |
| | \$ | Per | Manufacturer |

→ Restricted (RS1723)

Initiation - Crohn's disease

Both:

- 1 Mild to moderate ileal, ileocaecal or proximal Crohn's disease; and
- 2 Any of the following:
 - 2.1 Diabetes; or
 - 2.2 Cushingoid habitus; or
 - 2.3 Osteoporosis where there is significant risk of fracture; or
 - 2.4 Severe acne following treatment with conventional corticosteroid therapy; or
 - 2.5 History of severe psychiatric problems associated with corticosteroid treatment; or
 - 2.6 History of major mental illness (such as bipolar affective disorder) where the risk of conventional corticosteroid treatment causing relapse is considered to be high; or
 - 2.7 Relapse during pregnancy (where conventional corticosteroids are considered to be contraindicated).

Initiation - Collagenous and lymphocytic colitis (microscopic colitis)

Patient has a diagnosis of microscopic colitis (collagenous or lymphocytic colitis) by colonoscopy with biopsies.

Initiation - Gut Graft versus Host disease

Patient has gut Graft versus Host disease following allogenic bone marrow transplantation.

Initiation - non-cirrhotic autoimmune hepatitis

Re-assessment required after 6 months

All of the following:

- 1 Patient has autoimmune hepatitis*; and
- 2 Patient does not have cirrhosis; and
- 3 Any of the following:
 - 3.1 Diabetes; or
 - 3.2 Cushingoid habitus; or
 - 3.3 Osteoporosis where there is significant risk of fracture; or
 - 3.4 Severe acne following treatment with conventional corticosteroid therapy; or
 - 3.5 History of severe psychiatric problems associated with corticosteroid treatment; or
 - 3.6 History of major mental illness (such as bipolar affective disorder) where the risk of conventional corticosteroid treatment causing relapse is considered to be high; or

7

Pentasa

- 3.7 Relapse during pregnancy (where conventional corticosteroids are considered to be contraindicated); or
- 3.8 Adolescents with poor linear growth (where conventional corticosteroid use may limit further growth).

Note: Indications marked with * are unapproved indications.

Continuation - non-cirrhotic autoimmune hepatitis

Re-assessment required after 6 months

Treatment remains appropriate and the patient is benefitting from the treatment.

HYDROCORTISONE ACETATE

| Rectal foam 10%, CFC free (14 applications) | 26.55 | 21.1 g | Colifoam |
|---|-------|--------|----------|
| HYDROCORTISONE ACETATE WITH PRAMOXINE HYDROCHLORIDE Topical Aerosol foam, 1% with pramoxine hydrochloride 1% | | | |
| MESALAZINE | | | |
| Tab EC 400 mg | 49.50 | 100 | Asacol |
| Tab EC 500 mg | 49.50 | 100 | Asamax |
| Tab long-acting 500 mg - 1% DV Jul-20 to 2023 | 56.10 | 100 | Pentasa |
| Tab 800 mg | 85.50 | 90 | Asacol |
| Modified release granules 1 g | | 120 g | Pentasa |
| Suppos 500 mg | 22.80 | 20 | Asacol |
| Suppos 1 g | 54.60 | 30 | Pentasa |

e.g. Brand indicates brand example only. It is not a contracted product.

| | | Price excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|-----------|---------------------------|------------------|--------------------------------------|
| OLSALAZINE Tab 500 mg Cap 250 mg | | | 100 100 | Dipentum Dipentum |
| SODIUM CROMOGLICATE Cap 100 mg | | | | 1 · · · |
| SULFASALAZINE Tab 500 mg Tab EC 500 mg – 1% DV Dec-19 to 2022 | | . 14.00 . 15.53 | 100 100 | Salazopyrin Salazopyrin EN |
| Local Preparations for Anal and Rectal Disorders | | | | |
| Antihaemorrhoidal Preparations | | | | |
| CINCHOCAINE HYDROCHLORIDE WITH HYDROCORTISONE Oint 5 mg with hydrocortisone 5 mg per g Suppos 5 mg with hydrocortisone 5 mg per g FLUOCORTOLONE CAPROATE WITH FLUOCORTOLONE PIVAL Oint 950 mcg with fluocortolone pivalate 920 mcg and cinchoca | ATE AND C | 9.90 | 30 g 12 IE | Proctosedyl Proctosedyl |
| hydrochloride 5 mg per g Suppos 630 mcg with fluocortolone pivalate 610 mcg and cinch hydrochloride 1 mg | ocaine | | 30 g 12 | Ultraproct |
| Management of Anal Fissures | | | | |
| GLYCERYL TRINITRATE Oint 0.2% | | .22.00 | 30 g | Rectogesic |
| Rectal Sclerosants | | | | |
| OILY PHENOL [PHENOL OILY] Inj 5%, 5 ml vial | | | | |
| Antispasmodics and Other Agents Altering Gut M | lotility | | | |
| GLYCOPYRRONIUM BROMIDE Inj 200 mcg per ml, 1 ml ampoule | | . 17.14 | 10 | Max Health |
| HYOSCINE BUTYLBROMIDE Tab 10 mg – 1% DV Oct-20 to 2023 Inj 20 mg, 1 ml ampoule – 1% DV Jul-20 to 2023 | | | 100 5 | Buscopan Buscopan |
| MEBEVERINE HYDROCHLORIDE Tab 135 mg – 1% DV Jul-20 to 2023 | | 9.20 | 90 | Colofac |
| Antiulcerants | | | | |
| Antisecretory and Cytoprotective | | | | |
| | | | | |

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|---|-----------------------------------|---|--|
| H2 Antagonists | | | |
| CIMETIDINE Tab 200 mg Tab 400 mg FAMOTIDINE Tab 20 mg Tab 40 mg Inj 10 mg per ml, 2 ml vial Inj 10 mg per ml, 4 ml vial RANITIDINE – Restricted see terms below I Tab 150 mg I Tab 300 mg I Tab 300 mg I Tab 300 mg I Inj 25 mg per ml, 2 ml ampoule (<i>Ranitidine Relief Tab 150 mg to be delisted 1 October 2020)</i> (<i>Ranitidine Relief Tab 150 mg to be delisted 1 October 2020)</i> (<i>Ranitidine Relief Tab 300 mg to be delisted 1 October 2020)</i> (<i>Zantac Inj 25 mg per ml, 2 ml ampoule to be delisted 1 March 2021)</i> → Restricted (RS1703) Initiation Either: 1 For continuation use; or 2 Routine prevention of allergic reactions | 18.21 5.14 | 500 500 300 ml 5 | Ranitidine Relief Ranitidine Relief Peptisoothe Zantac |
| Proton Pump Inhibitors | | | |
| LANSOPRAZOLE Cap 15 mg - 1% DV Sep-18 to 2021 Cap 30 mg - 1% DV Sep-18 to 2021 OMEPRAZOLE ↓ Tab dispersible 20 mg → Restricted (RS1027) Initiation Only for use in tube-fed patients. | | 100 100 | Lanzol Relief Lanzol Relief |
| Cap 10 mg Cap 20 mg Cap 40 mg Powder for oral liq Inj 40 mg ampoule with diluent – 1% DV Oct-19 to 2022 Inj 40 mg vial – 1% DV Oct-19 to 2022. PANTOPRAZOLE Tab EC 20 mg – 1% DV Oct-19 to 2022. Tab EC 40 mg – 1% DV Oct-19 to 2022. Inj 40 mg vial | | 90 90 5 g 5 5 100 100 | Omeprazole actavis 10 Omeprazole actavis 20 Omeprazole actavis 40 Midwest Dr Reddy's Omeprazole Omezol IV Panzop Relief Panzop Relief |
| Site Protective Agents | | | |
| COLLOIDAL BISMUTH SUBCITRATE Tab 120 mg SUCRALFATE Tab 1 g | 14.51 | 50 | Gastrodenol |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|---------------------|--|
| Bile and Liver Therapy | | | |
| L-ORNITHINE L-ASPARTATE – Restricted see terms below ↓ Grans for oral liquid 3 g → Restricted (RS1261) | | | |
| Initiation For patients with chronic hepatic encephalopathy who have not response where lactulose is contraindicated. | onded to treatment with | h, or are in | tolerant to lactulose, or |
| RIFAXIMIN – Restricted see terms below ↓ Tab 550 mg → Restricted (RS1416) Initiation | 625.00 | 56 | Xifaxan |
| For patients with hepatic encephalopathy despite an adequate trial of | maximum tolerated d | oses of lac | tulose. |
| Diabetes | | | |
| Alpha Glucosidase Inhibitors | | | |
| ACARBOSE Tab 50 mg – 1% DV Sep-18 to 2021 Tab 100 mg – 1% DV Sep-18 to 2021 | | 90 90 | Glucobay Glucobay |
| Hyperglycaemic Agents | | | |
| DIAZOXIDE - Restricted see terms below ↓ Cap 25 mg ↓ Cap 100 mg ↓ Oral liq 50 mg per ml | | 100 100 30 ml | Proglicem Proglycem Glucagen Hypokit |
| Insulin - Intermediate-Acting Preparations | | | |
| INSULIN ASPART WITH INSULIN ASPART PROTAMINE Inj insulin aspart 30% with insulin aspart protamine 70%, 100 u p 3 ml prefilled pen INSULIN ISOPHANE Inj insulin human 100 u per ml, 10 ml vial Inj insulin human 100 u per ml, 3 ml cartridge | | 5 | NovoMix 30 FlexPen |

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|--|-----------------------------------|----------|-------------------------------------|
| INSULIN LISPRO WITH INSULIN LISPRO PROTAMINE | Ψ | | Manufacturer |
| Inj insulin lispro 25% with insulin lispro protamine 75%, 100 u pe 3 ml cartridge | | 5 | Humalog Mix 25 |
| Inj insulin lispro 50% with insulin lispro protamine 50%, 100 u pe 3 ml cartridge | er ml, | 5 | Humalog Mix 50 |
| INSULIN NEUTRAL WITH INSULIN ISOPHANE | | | - |
| Inj insulin neutral 30% with insulin isophane 70%, 100 u per ml, vial | 10 ml | | |
| Inj insulin neutral 30% with insulin isophane 70%, 100 u per ml, cartridge | 3 ml | | |
| Inj insulin neutral 40% with insulin isophane 60%, 100 u per ml, cartridge | 3 ml | | |
| Inj insulin neutral 50% with insulin isophane 50%, 100 u per ml, cartridge | 3 ml | | |
| Insulin - Long-Acting Preparations | | | |
| NSULIN GLARGINE Inj 100 u per ml, 3 ml disposable pen | 94 50 | 5 | Lantus SoloStar |
| Inj 100 u per ml, 3 ml cartridge | | 5 | Lantus |
| lnj 100 u per ml, 10 ml vial | 63.00 | 1 | Lantus |
| Insulin - Rapid-Acting Preparations | | | |
| NSULIN ASPART Inj 100 u per ml, 10 ml vial | | | |
| Inj 100 u per ml, 3 ml cartridge Inj 100 u per ml, 3 ml syringe | 51.19 | 5 | NovoRapid FlexPen |
| NSULIN GLULISINE Inj 100 u per ml, 10 ml vial | | 1 | Apidra |
| Inj 100 u per ml, 3 ml cartridge | | 5 | Apidra |
| Inj 100 u per ml, 3 ml disposable pen | | 5 | Apidra Solostar |
| NSULIN LISPRO | | | |
| Inj 100 u per ml, 10 ml vial Inj 100 u per ml, 3 ml cartridge | | | |
| Insulin - Short-Acting Preparations | | | |
| NSULIN NEUTRAL | | | |
| Inj human 100 u per ml, 10 ml vial Inj human 100 u per ml, 3 ml cartridge | | | |
| Oral Hypoglycaemic Agents | | | |
| | 0.00 | 100 | Deenil |
| Tab 5 mg – 1% DV Oct-18 to 2021 GLICLAZIDE | 6.00 | 100 | Daonil |
| Tab 80 mg - 1% DV Nov-20 to 2023 | | 500 | Glizide |
| GLIPIZIDE Tab 5 mg – 1% DV Dec-18 to 2021 | | 100 | Minidiab |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | | Brand or |
|--|--------------------------|----------|-------------------------|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer |
| | Ψ | 1.01 | Manufacturer |
| METFORMIN HYDROCHLORIDE | 0.00 | 4 000 | A |
| Tab immediate-release 500 mg - 1% DV Feb-19 to 2021 | | 1,000 | Apotex |
| Tab immediate-release 850 mg - 1% DV Feb-19 to 2021 | 7.04 | 500 | Apotex |
| PIOGLITAZONE | | | |
| Tab 15 mg - 1% DV Oct-18 to 2021 | | 90 | Vexazone |
| Tab 30 mg - 1% DV Oct-18 to 2021 | 5.06 | 90 | Vexazone |
| Tab 45 mg - 1% DV Oct-18 to 2021 | 7.10 | 90 | Vexazone |
| VILDAGLIPTIN | | | |
| Tab 50 mg | | 60 | Galvus |
| VILDAGLIPTIN WITH METFORMIN HYDROCHLORIDE | | | |
| Tab 50 mg with 1,000 mg metformin hydrochloride | 40.00 | 60 | Galvumet |
| Tab 50 mg with 850 mg metformin hydrochloride | | 60 | Galvumet |
| | | 00 | Gaivaniet |
| Digestives Including Enzymes | | | |
| | | | |
| PANCREATIC ENZYME | | | |
| Cap pancreatin (175 mg (25,000 U lipase, 22,500 U amylase, 1,250 | U | | |
| protease)) | | | |
| Cap pancreatin 150 mg (amylase 8,000 Ph Eur U, lipase 10,000 Ph | Eur | | |
| U, total protease 600 Ph Eur U) - 1% DV Sep-18 to 2021 | | 100 | Creon 10000 |
| Cap pancreatin 300 mg (amylase 18,000 Ph Eur U, lipase 25,000 P | | | |
| Eur U, total protease 1,000 Ph Eur U) - 1% DV Sep-18 to 202 | | 100 | Creon 25000 |
| Modified release granules pancreatin 60.12 mg (amylase 3,600 Ph | | | |
| U, lipase 5,000 Ph Eur U, protease 200 Ph Eur U) | | 20 g | Creon Micro |
| Powder pancreatin 60.12 mg (3,600 Ph. Eur. u/amylase, 5,000 Ph | | | |
| Eur. u/lipase and 200 Ph. Eur. u/protease) | | | |
| LIRSODEOXYCHOLIC ACID - Restricted see terms below | | | |

URSODEOXYCHOLIC ACID - Restricted see terms below

Initiation – Alagille syndrome or progressive familial intrahepatic cholestasis Either:

1 Patient has been diagnosed with Alagille syndrome; or

2 Patient has progressive familial intrahepatic cholestasis.

Initiation - Chronic severe drug induced cholestatic liver injury

All of the following:

1 Patient has chronic severe drug induced cholestatic liver injury; and

2 Cholestatic liver injury not due to Total Parenteral Nutrition (TPN) use in adults; and

3 Treatment with ursodeoxycholic acid may prevent hospital admission or reduce duration of stay.

Initiation - Primary biliary cholangitis

Both:

- 1 Primary biliary cholangitis confirmed by antimitochondrial antibody titre (AMA) > 1:80, and raised cholestatic liver enzymes with or without raised serum IgM or, if AMA is negative by liver biopsy; and
- 2 Patient not requiring a liver transplant (bilirubin > 100 umol/l; decompensated cirrhosis.

Initiation - Pregnancy

Patient diagnosed with cholestasis of pregnancy.

Initiation - Haematological transplant

Both:

1 Patient at risk of veno-occlusive disease or has hepatic impairment and is undergoing conditioning treatment prior to

continued...

| Price (ex man. excl. GST) | | Brand or Generic |
|------------------------------|-----|---------------------|
| \$ | Per | Manufacturer |

continued...

allogenic stem cell or bone marrow transplantation; and

2 Treatment for up to 13 weeks.

Initiation - Total parenteral nutrition induced cholestasis

Both:

- 1 Paediatric patient has developed abnormal liver function as indicated on testing which is likely to be induced by TPN; and
- 2 Liver function has not improved with modifying the TPN composition.

Laxatives

| CITRIC ACID WITH MAGNESIUM OXIDE AND SODIUM PICOSULFATE Powder for oral soln 12 g with magnesium oxide 3.5 g and sodium picosulfate 10 mg per sachet MACROGOL 3350 WITH ASCORBIC ACID, POTASSIUM CHLORIDE AND S Powder for oral soln 755.68 mg with ascorbic acid 85.16 mg, potassium chloride 10.55 mg, sodium chloride 37.33 mg and sodium sulphate 80.62 mg per g, 210 g sachet Powder for oral soln 755 68 mg with ascorbic acid 85 16 mg, potassium | SODIUM CH | LORIDE | e.g. PicoPrep e.g. Glycoprep-C |
|--|-----------|------------|-----------------------------------|
| Powder for oral soln 755.68 mg with ascorbic acid 85.16 mg, potassium chloride 10.55 mg, sodium chloride 37.33 mg and sodium sulphate 80.62 mg per g, 70 g sachet MACROGOL 3350 WITH POTASSIUM CHLORIDE, SODIUM BICARBONATI Powder for oral soln 59 g with potassium chloride 0.7425 g, sodium bicarbonate 1.685 g, sodium chloride 1.465 g and sodium sulphate | | | |
| 5.685 g per sachet – 1% DV Aug-19 to 2022 | 14.31 | 4 | Klean Prep |
| Bulk-Forming Agents | | | |
| ISPAGHULA (PSYLLIUM) HUSK Powder for oral soln – 1% DV Nov-20 to 2023 STERCULIA WITH FRANGULA – Restricted: For continuation only → Powder for oral soln | 12.20 | 500 g | Konsyl-D |
| Faecal Softeners | | | |
| DOCUSATE SODIUM Tab 50 mg – 1% DV Oct-20 to 2023 Tab 120 mg – 1% DV Oct-20 to 2023 DOCUSATE SODIUM WITH SENNOSIDES | 3.13 | 100 100 | Coloxyl Coloxyl |
| Tab 50 mg with sennosides 8 mg – 1% DV Jun-18 to 2021 PARAFFIN Oral liquid 1 mg per ml Enema 133 ml | 3.10 | 200 | Laxsol |
| POLOXAMER Oral drops 10% – 1% DV Nov-20 to 2023 | 3.98 | 30 ml | Coloxyl |
| Opioid Receptor Antagonists - Peripheral | | | |
| METHYLNALTREXONE BROMIDE – Restricted see terms on the next page Inj 12 mg per 0.6 ml vial | | 1 7 | Relistor Relistor |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| | exci. \$ | GST) | Per | Generic Manufacturer |
|------------------------------|---|---|--|--|
| | | | | |
| | | | erated. | |
| | | | | |
| | | | | |
| | 9.2 | 5 | 20 | PSM |
| | 3.3 | 3 | 500 ml | Laevolac |
| odium sodium DV | | | | Molaxole |
| nl – 1% | 0.7 | 0 | 00 | Moldxole |
| | .29.98 | В | 50 | Micolette |
| | 2.50 | 0 | 1 | Fleet Phosphate Enema |
| | | | | |
| | | | 200 10 | Lax-Tabs Lax-Suppositories |
| | | | | |
| 1, [.] | 142.60 | D | 1 | Myozyme |
| | rion are una RBONATE / podium sodium DV | tion are ineffectiv tion are unable to | tion are ineffective; or tion are unable to be tole 9.25 | tion are ineffective; or tion are unable to be tolerated. |

1 The patient is aged up to 24 months at the time of initial application and has been diagnosed with infantile Pompe disease; and

continued...

| Price | | Brand or |
|-------------------|-----|--------------|
| (ex man. excl. GS | | Generic |
| \$ | Per | Manufacturer |

continued...

- 2 Any of the following:
 - 2.1 Diagnosis confirmed by documented deficiency of acid alpha-glucosidase by prenatal diagnosis using chorionic villus biopsies and/or cultured amniotic cells; or
 - 2.2 Documented deficiency of acid alpha-glucosidase, and urinary tetrasaccharide testing indicating a diagnostic elevation of glucose tetrasaccharides; or
 - 2.3 Documented deficiency of acid alpha-glucosidase, and documented molecular genetic testing indicating a disease-causing mutation in the acid alpha-glucosidase gene (GAA gene); or
 - 2.4 Documented urinary tetrasaccharide testing indicating a diagnostic elevation of glucose tetrasaccharides, and molecular genetic testing indicating a disease-causing mutation in the GAA gene; and
- 3 Patient has not required long-term invasive ventilation for respiratory failure prior to starting enzyme replacement therapy (ERT); and
- 4 Patient does not have another life-threatening or severe disease where the prognosis is unlikely to be influenced by ERT or might be reasonably expected to compromise a response to ERT; and
- 5 Alglucosidase alfa to be administered at doses no greater than 20 mg/kg every 2 weeks.

Continuation

Re-assessment required after 12 months

All of the following:

- 1 The treatment remains appropriate for the patient and the patient is benefiting from treatment; and
- 2 Alglucosidase alfa to be administered at doses no greater than 20 mg/kg every 2 weeks; and
- 3 Patient has not had severe infusion-related adverse reactions which were not preventable by appropriate pre-medication and/or adjustment of infusion rates; and
- 4 Patient has not developed another life threatening or severe disease where the long term prognosis is unlikely to be influenced by ERT; and
- 5 Patient has not developed another medical condition that might reasonably be expected to compromise a response to ERT; and
- 6 There is no evidence of life threatening progression of respiratory disease as evidenced by the needed for > 14 days of invasive ventilation; and
- 7 There is no evidence of new or progressive cardiomyopathy.

ARGININE

Powder Inj 600 mg per ml, 25 ml vial

BETAINE - Restricted see terms below

→ Restricted (RS1751)

Initiation

Metabolic physician

Re-assessment required after 12 months

All of the following:

- 1 The patient has a confirmed diagnosis of homocystinuria; and
- 2 Any of the following:
 - 2.1 A cystathionine beta-synthase (CBS) deficiency; or
 - 2.2 A 5,10-methylene-tetrahydrofolate reductase (MTHFR) deficiency; or
 - 2.3 A disorder of intracellular cobalamin metabolism; and
- 3 An appropriate homocysteine level has not been achieved despite a sufficient trial of appropriate vitamin supplementation.

Continuation

Re-assessment required after 12 months

The treatment remains appropriate and the patient is benefiting from treatment.

14

| | | Price | | Brand or |
|--|-----------------|------------------|------------|----------------------------------|
| | (ex man. | excl. GST) \$ | Per | Generic Manufacturer |
| BIOTIN – Restricted see terms below | | | | |
| ↓ Cap 50 mg | | | | |
| Cap 100 mg | | | | |
| Inj 10 mg per ml, 5 ml vial | | | | |
| → Restricted (RS1330) | | | | |
| Metabolic physician or metabolic disorders dietitian | | | | |
| GALSULFASE – Restricted see terms below | | | | |
| Inj 1 mg per ml, 5 ml vial | 2,2 | 234.00 | 1 | Naglazyme |
| → Restricted (RS1752) | | | | |
| Initiation | | | | |
| Metabolic physician | | | | |
| Re-assessment required after 12 months | | | | |
| Both: | | | | |
| 1 The patient has been diagnosed with mucopolysaccharidosis | s VI; and | | | |
| 2 Either: | | | | |
| 2.1 Diagnosis confirmed by demonstration of N-acetyl-ga | | | (arylsulfa | tase B) deficiency confirme |
| by either enzyme activity assay in leukocytes or skin | | | | |
| 2.2 Detection of two disease causing mutations and patie | ent has a sibl | ling who is k | nown to h | ave mucopolysaccharidosi |
| VI. | | | | |
| Continuation | | | | |
| Re-assessment required after 12 months | | | | |
| All of the following: | | | | |
| 1 The treatment remains appropriate for the patient and the pa | | | | |
| 2 Patient has not had severe infusion-related adverse reaction | s which were | e not preven | table by a | ppropriate pre-medication |
| and/or adjustment of infusion rates; and | | | | |
| 3 Patient has not developed another life threatening or severe | disease whe | ere the long | term prog | nosis is unlikely to be |
| influenced by Enzyme Replacement Therapy (ERT); and | | | | |
| 4 Patient has not developed another medical condition that mig EDT | gnt reasonat | by be expec | ted to con | npromise a response to |
| ERT. | | | | |
| HAEM ARGINATE | | | | |
| Inj 25 mg per ml, 10 ml ampoule | | | | |
| IDURSULFASE – Restricted see terms below | | | | |
| Inj 2 mg per ml, 3 ml vial | 4,6 | 608.30 | 1 | Elaprase |
| → Restricted (RS1546) | | | | |
| Initiation | | | | |
| Metabolic physician | | | | |
| Limited to 24 weeks treatment | | | | |
| All of the following: | | | | |
| The patient has been diagnosed with Hunter Syndrome (muc 2 Either: | copolysaccha | ardosis II); a | ina | |
| | | <i></i> | | d a a lla la craithe an annuna a |
| 2.1 Diagnosis confirmed by demonstration of iduronate 2- | -suitatase de | eticlency in v | | a cells by either enzyme |
| assay in cultured skin fibroblasts; or | ata 0 aulfata | | d | |
| 2.2 Detection of a disease causing mutation in the iduron | | 0 | | months and treatment will |
| 3 Patient is going to proceed with a haematopoietic stem cell to | ranspiant (H | SCI) Within | me next 3 | months and treatment with |
| idursulfase would be bridging treatment to transplant; and | niraton (fail) | iro prior to o | tartina En | zuma Danlagament Theres |
| 4 Patient has not required long-term invasive ventilation for res (ERT); and | spiratory falle | | larling En | zyme rieplacement Therap |
| (ERT), and E. Iduraulface to be administered for a total of 04 weeks (aguing | | | | |

5 Idursulfase to be administered for a total of 24 weeks (equivalent to 12 weeks pre- and 12 weeks post-HSCT) at doses no greater than 0.5 mg/kg every week.

| | | Price | | | Brand or |
|---|---|--------------------------|-----------------------------|----------------------|---------------------------|
| | (ex man | | GST) | Per | Generic Manufacturer |
| LARONIDASE – Restricted see terms below ↓ Inj 100 U per ml, 5 ml vial | 1, | 335.1 | 6 | 1 | Aldurazyme |
| Initiation Metabolic physician <i>Limited to 24 weeks</i> treatment All of the following: 1 The patient has been diagnosed with Hurler Syndrome (mucopo 2 Either: | olysaccha | ardosi | s I-H); | and | |
| 2.1 Diagnosis confirmed by demonstration of alpha-L-iduron assay in cultured skin fibroblasts; or 2.2 Detection of two disease causing mutations in the alphato to have Hurler syndrome; and | | | | | |
| 3 Patient is going to proceed with a haematopoietic stem cell tran laronidase would be bridging treatment to transplant; and 4 Patient has not required long-term invasive ventilation for respir (ERT); and 5 Laronidase to be administered for a total of 24 weeks (equivalent than 100 units/kg every week. | atory fail | ure pr | ior to s | tarting E | nzyme Replacement Therapy |
| LEVOCARNITINE - Restricted see terms below ↓ Cap 500 mg ↓ Oral soln 1,000 mg per 10 ml ↓ Oral soln 1,100 mg per 15 ml ↓ Inj 200 mg per ml, 5 ml vial → Restricted (RS1035) Neurologist, metabolic physician or metabolic disorders dietitian PYRIDOXAL-5-PHOSPHATE - Restricted see terms below ↓ Tab 50 mg → Restricted (RS1331) Neurologist, metabolic physician or metabolic disorders dietitian SAPROPTERIN DIHYDROCHLORIDE - Restricted see terms below | | | | | |
| Tab soluble 100 mg Restricted (RS1753) Initiation Metabolic physician Re-assessment required after 1 month All of the following: | | 452.7 | 0 | 30 | Kuvan |
| Patient has phenylketonuria (PKU) and is pregnant or actively p Treatment with sapropterin is required to support management Sapropterin to be administered at doses no greater than a total Sapropterin to be used alone or in combination with PKU dietar Total treatment duration with sapropterin will not exceed 22 mor becoming pregnant) and treatment will be stopped after delivery | of PKU d daily dos y managenths for e | uring e of 2 ement | pregna 20 mg/k t; and | ancy; and (g; and | 1 |
| Continuation Re-assessment required after 12 months All of the following: | | | | | |

1 Either:

| | l (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|------------------------------|-----------------------------|--------------------|----------------------|---|
| continued | | | | | |
| 1.1 Following the initial one-month approval, the patient h of sapropterin with a clinically appropriate reduction in pregnancy; or 1.2 On subsequent renewal applications, the patient has a sapropterin and maintained adequate phenylalanine here. | ı phenylalan previously d | ine le [.] emon | vels to strated | support I respons | management of PKU during se to treatment with |
| 2 Any of the following: | | | | | 5 F - 5 , , |
| 2.1 Patient continues to be pregnant and treatment with s2.2 Patient is actively planning a pregnancy and this is the2.3 Treatment with sapropterin is required for a second or during pregnancy; and | r subsequen | al for It preg | treatm nancy | ent with to suppo | sapropterin; or |
| 3 Sapropterin to be administered at doses no greater than a toi 4 Sapropterin to be used alone or in combination with PKU diet 5 Total treatment duration with sapropterin will not exceed 22 r becoming pregnant) and treatment will be stopped after deliv | tary manage nonths for e | ement | ; and | 0 | des time for planning and |
| SODIUM BENZOATE Cap 500 mg Powder Soln 100 mg per ml Inj 20%, 10 ml ampoule | | | | | |
| SODIUM PHENYLBUTYRATE - Some items restricted see terms | below | | | | |
| Tab 500 mg Grans 483 mg per g Oral liq 250 mg per ml liq 200 mg per ml | 1,9 | 920.00 | 0 | 174 g | Pheburane |
| Inj 200 mg per ml, 10 ml ampoule → Restricted (RS1754) Initiation Metabolic physician | | | | | |
| Re-assessment required after 12 months For the chronic management of a urea cycle disorder involving a def transcarbamylase or argininosuccinate synthetase. Continuation | ficiency of ca | arbam | lylphos | phate sy | nthetase, ornithine |
| Re-assessment required after 12 months The treatment remains appropriate and the patient is benefiting from TALIGLUCERASE ALFA – Restricted see terms below | n treatment. | | | | |
| Inj 200 unit vial Restricted (RS1034) Initiation | 1,(| 072.00 | 0 | 1 | Elelyso |
| Only for use in patients with approval by the Gaucher Treatment Par TRIENTINE DIHYDROCHLORIDE Cap 300 mg | nel. | | | | |
| Minerals | | | | | |
| Calcium | | | | | |
| CALCIUM CARBONATE Tab 1.25 g (500 mg elemental) Tab eff 1.25 g (500 mg elemental) Tab eff 1.75 g (1 g elemental) | | 7.52 | 2 | 250 | Arrow-Calcium |

| | l (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------------|----------------------|------|--------|-------------------------------------|
| Fluoride | | | | | |
| SODIUM FLUORIDE Tab 1.1 mg (0.5 mg elemental) | | | | | |
| lodine | | | | | |
| POTASSIUM IODATE Tab 253 mcg (150 mcg elemental iodine) – 1% DV Oct-20 to 202 POTASSIUM IODATE WITH IODINE Oral liq 10% with iodine 5% | 3 | 4.58 | 3 | 90 | NeuroTabs |
| Iron | | | | | |
| ERRIC CARBOXYMALTOSE – Restricted see terms below Inj 50 mg per ml, 10 ml vial → Restricted (RS1417) nitiation | | 150.00 |) | 1 | Ferinject |
| Freatment with oral iron has proven ineffective or is clinically inappropr FERROUS FUMARATE Tab 200 mg (65 mg elemental) – 1% DV Jan-19 to 2021 FERROUS FUMARATE WITH FOLIC ACID | | 3.09 | Э | 100 | Ferro-tab |
| Tab 310 mg (100 mg elemental) with folic acid 350 mcg – 1% DV Jun-18 to 2021 | | 4.68 | 3 | 60 | Ferro-F-Tabs |
| ERROUS SULFATE Oral liq 30 mg (6 mg elemental) per ml – 1% DV Nov-19 to 2022. ERROUS SULPHATE | | .12.08 | 3 | 500 ml | Ferodan |
| Tab long-acting 325 mg (105 mg elemental) – 1% DV Jun-18 to 2 ERROUS SULPHATE WITH ASCORBIC ACID | 2021 | 2.00 | 6 | 30 | Ferrograd |
| Tab long-acting 325 mg (105 mg elemental) with ascorbic acid 500 RON POLYMALTOSE |) mg | | | | |
| Inj 50 mg per ml, 2 ml ampoule | | .34.50 | 0 | 5 | Ferrosig |
| RON SUCROSE Inj 20 mg per ml, 5 ml ampoule | | 100.00 | D | 5 | Venofer |

Magnesium

| MAGNESIUM AMINO ACID CHELATE Cap 750 mg (150 mg elemental) |
|---|
| MAGNESIUM CHLORIDE |
| lnj 1 mmol per 1 ml, 100 ml bag |
| MAGNESIUM HYDROXIDE Tab 311 mg (130 mg elemental) |
| MAGNESIUM OXIDE |
| Cap 663 mg (400 mg elemental) |
| Cap 696 mg (420 mg elemental) |
| |

18

| | Price (ex man. excl. GS \$ | T) Per | Brand or Generic Manufacturer |
|--|----------------------------------|-----------|-------------------------------------|
| MAGNESIUM OXIDE WITH MAGNESIUM ASPARTATE, MAGNESIL Cap 500 mg with magnesium aspartate 100 mg, magnesium ami chelate 100 mg and magnesium citrate 100 mg (360 mg eler magnesium) | no acid | IELATE AN | D MAGNESIUM CITRATE |
| MAGNESIUM SULPHATE Inj 0.4 mmol per ml, 250 ml bag Inj 2 mmol per ml, 5 ml ampoule Inj 100 mg per ml, 50 ml bag | | 10 | DBL |
| Zinc | | | |
| ZINC Oral liq 5 mg per 5 drops ZINC CHLORIDE Inj 5.3 mg per ml (5.1 mg per ml elemental), 2 ml ampoule ZINC SULPHATE Cap 137.4 mg (50 mg elemental) – 1% DV Dec-19 to 2022 | | 100 | Zincaps |
| Mouth and Throat | | | |
| Agents Used in Mouth Ulceration | | | |
| BENZYDAMINE HYDROCHLORIDE Soln 0.15% Spray 0.15% Spray 0.3% | | | |
| BENZYDAMINE HYDROCHLORIDE WITH CETYLPYRIDINIUM CHL Lozenge 3 mg with cetylpyridinium chloride | ORIDE | | |
| CARBOXYMETHYLCELLULOSE Oral spray | | | |
| CARMELLOSE SODIUM WITH PECTIN AND GELATINE Paste Powder | | | |
| CHLORHEXIDINE GLUCONATE Mouthwash 0.2% | 2.57 | 200 ml | healthE |
| CHOLINE SALICYLATE WITH CETALKONIUM CHLORIDE Adhesive gel 8.7% with cetalkonium chloride 0.01% | | | |
| DICHLOROBENZYL ALCOHOL WITH AMYLMETACRESOL Lozenge 1.2 mg with amylmetacresol 0.6 mg | | | |
| TRIAMCINOLONE ACETONIDE Paste 0.1% - 1% DV Nov-20 to 2023 | 5.33 | 5 g | Kenalog in Orabase |
| Oropharyngeal Anti-Infectives | | | |
| AMPHOTERICIN B Lozenge 10 mg | 5.96 | 20 | Fungilin |
| MICONAZOLE | | | Ū |
| Oral gel 20 mg per g – 1% DV Sep-18 to 2021 | 4.74 | 40 g | Decozol |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|---|-----------------------------------|----------|-------------------------------------|
| NYSTATIN Oral liquid 100,000 u per ml – 1% DV Oct-20 to 2023 | | 24 ml | Nilstat |
| Other Oral Agents | | | |
| HYALURONIC ACID WITH LIDOCAINE [LIGNOCAINE] Inj 20 mg per ml | | | |
| SODIUM HYALURONATE [HYALURONIC ACID] - Restricted see ↓ Inj 20 mg per ml, 1 ml syringe → Restricted (RS1175) Otolaryngologist | e terms below | | |
| THYMOL GLYCERIN Compound, BPC | 9.15 | 500 ml | PSM |
| Vitamins | | | |
| Multivitamin Preparations | | | |
| MULTIVITAMIN AND MINERAL SUPPLEMENT - Restricted see t | | 180 | Clinicians Multivit & |
| → Restricted (RS1498) Initiation Limited to 3 months treatment Both: Patient was admitted to hospital with burns; and Any of the following: | | | Mineral Boost |
| 2.3 Nutritional status prior to admission or dietary intake i MULTIVITAMIN RENAL – Restricted see terms below | | | |
| ↓ Cap → Restricted (RS1499) Initiation Fither | 6.49 | 30 | Clinicians Renal Vit |

Either:

1 The patient has chronic kidney disease and is receiving either peritoneal dialysis or haemodialysis; or

2 The patient has chronic kidney disease grade 5, defined as patient with an estimated glomerular filtration rate of < 15 ml/min/1.73m² body surface area (BSA).

| | | Price excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|--------|---------------------------|-------|-------------------------------------|
| MULTIVITAMINS | | | | |
| Tab (BPC cap strength) - 1% DV Mar-20 to 2022 | | . 11.45 | 1,000 | Mvite |
| cap vitamin A 2500 u, betacarotene 3 mg, cholecalciferol 11 mcg, a tocopherol 150 u, phytomenadione 150 mcg, folic acid 0.2 mg, ascorbic acid 100 mg, thiamine 1.5 mg, pantothenic acid 12 m riboflavin 1.7 mg, niacin 20 mg, pyridoxine hydrochloride 1.9 m cyanocobalamin 3 mcg, zinc 7.5 mg and biotin 100 mcg → Restricted (RS1620) | g, | | | e.g. Vitabdeck |
| Initiation | | | | |
| Any of the following: | | | | |
| Patient has cystic fibrosis with pancreatic insufficiency; or Patient is an infant or child with liver disease or short gut syndro Patient has severe malabsorption syndrome. | me; or | | | |
| I Powder vitamin A 4200 mcg with vitamin D 155.5 mcg, vitamin E 21.4 mg, vitamin C 400 mg, vitamin K1 166 mcg thiamine 3.2 m riboflavin 4.4 mg, niacin 35 mg, vitamin B6 3.4 mg, folic acid 303 mcg, vitamin B12 8.6 mcg, biotin 214 mcg, pantothenic ac 17 mg, choline 350 mg and inositol 700 mg → Restricted (RS1178) | 0 | | | e.g. Paediatric Seravit |
| Initiation | | | | |
| Patient has inborn errors of metabolism. Inj thiamine hydrochloride 250 mg with riboflavin 4 mg and pyridoxi hydrochloride 50 mg, 5 ml ampoule (1) and inj ascorbic acid 50 | | | | |
| with nicotinamide 160 mg and glucose 1000 mg, 5 ml ampoule Inj thiamine hydrochloride 250 mg with riboflavin 4 mg and pyridoxi hydrochloride 50 mg, 5 ml ampoule (1) and inj ascorbic acid 50 | ne | | | e.g. Pabrinex IV |
| with nicotinamide 160 mg, 2 ml ampoule (1) Inj thiamine hydrochloride 500 mg with riboflavin 8 mg and pyridoxi hydrochloride 100 mg, 10 ml ampoule (1) and inj ascorbic acid 1000 mg with nicotinamide 320 mg and glucose 2000 mg, 10 r | ne | | | e.g. Pabrinex IM |
| ampoule (1) | | | | e.g. Pabrinex IV |
| Vitamin A | | | | |

RETINOL

Tab 10,000 iu Cap 25,000 iu Oral liq 150,000 iu per ml Oral liq 666.7 mcg per 2 drops, 10 ml Oral liq 5,000 iu per drop, 30 ml

Vitamin B

| HYDROXOCOBALAMIN | | |
|--|-----|----------------|
| Inj 1 mg per ml, 1 ml ampoule - 1% DV Sep-18 to 20211.89 | 3 | Neo-B12 |
| PYRIDOXINE HYDROCHLORIDE | | |
| Tab 25 mg - 1% DV Oct-20 to 20232.70 | 90 | Vitamin B6 25 |
| Tab 50 mg 13.63 | 500 | Apo-Pyridoxine |
| Inj 100 mg per ml, 2 ml vial | | |
| Inj 100 mg per ml, 1 ml ampoule | | |
| Inj 100 mg per ml, 30 ml vial | | |

| Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|---------------------|-------------------------------------|
| THIAMINE HYDROCHLORIDE Tab 50 mg | 100 | Max Health e.g. Benerva |
| Inj 100 mg per ml, 2 ml vial VITAMIN B COMPLEX Tab strong, BPC7.15 | 500 | Bplex |
| Vitamin C | | ' |
| ASCORBIC ACID Tab 100 mg – 1% DV Mar-20 to 2022 9.90 Tab chewable 250 mg | 500 | Cvite |
| Vitamin D | | |
| ALFACALCIDOL Cap 0.25 mcg | 100 100 20 ml | One-Alpha One-Alpha One-Alpha |
| CALCITRIOL Cap 0.25 mcg – 1% DV Oct-19 to 2022 | 100 100 | Calcitriol-AFT Calcitriol-AFT |
| COLECALCIFEROL Cap 1.25 mg (50,000 iu) | 12 4.8 ml | Vit.D3 Puria |

Vitamin E

ALPHA TOCOPHERYL - Restricted see terms below

I Oral liq 156 u per ml

→ Restricted (RS1632)

Initiation – Cystic fibrosis

Both:

- 1 Cystic fibrosis patient; and
- 2 Either:
 - Patient has tried and failed the other available funded fat soluble vitamin A,D,E,K supplement (Vitabdeck); or
 The other available funded fat soluble vitamin A,D,E,K supplement (Vitabdeck) is contraindicated or clinically inappropriate for the patient.

Initiation – Osteoradionecrosis

For the treatment of osteoradionecrosis.

Initiation - Other indications

All of the following:

- 1 Infant or child with liver disease or short gut syndrome; and
- 2 Requires vitamin supplementation; and
- 3 Either:
 - 3.1 Patient has tried and failed the other available funded fat soluble vitamin A,D,E,K supplements (Vitabdeck); or
 - 3.2 The other available funded fat soluble vitamin A,D,E,K supplement (Vitabdeck) is contraindicated or clinically inappropriate for patient.

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST |) | Generic |
| \$ | Per | Manufacturer |

ALPHA TOCOPHERYL ACETATE - Restricted see terms below

- € Cap 500 u

↓ Oral liq 156 u per ml

→ Restricted (RS1176)

Initiation - Cystic fibrosis

Both:

- 1 Cystic fibrosis patient; and
- 2 Either:
 - 2.1 Patient has tried and failed the other available funded fat soluble vitamin A,D,E,K supplement (Vitabdeck); or
 - 2.2 The other available funded fat soluble vitamin A,D,E,K supplement (Vitabdeck) is contraindicated or clinically inappropriate for the patient.

Initiation – Osteoradionecrosis

For the treatment of osteoradionecrosis.

Initiation – Other indications

All of the following:

- 1 Infant or child with liver disease or short gut syndrome; and
- 2 Requires vitamin supplementation; and
- 3 Either:
 - 3.1 Patient has tried and failed the other available funded fat soluble vitamin A,D,E,K supplements (Vitabdeck); or
 - 3.2 The other available funded fat soluble vitamin A,D,E,K supplement (Vitabdeck) is contraindicated or clinically inappropriate for patient.

| | Price (ex man. excl. GST) | | Brand or Generic |
|---|---|---------------------------------|--|
| | \$ | Per | Manufacturer |
| Antianaemics | | | |
| Hypoplastic and Haemolytic | | | |
| EPOETIN ALFA - Restricted see terms below Inj 1,000 iu in 0.5 ml syringe - 1% DV Apr-19 to 2022 | 100.00 150.00 96.50 125.00 145.00 175.00 197.50 250.00 | 6 6 6 6 6 6 1 | Binocrit Binocrit Binocrit Binocrit Binocrit Binocrit Binocrit Binocrit |
| 4 Patient is on haemodialysis or peritoneal dialysis. | | | |
| Initiation – myelodysplasia* <i>Re-assessment required after 2 months</i> All of the following: 1 Patient has a confirmed diagnosis of myelodysplasia (MDS); a | and | | |
| 2 Has had symptomatic anaemia with haemoglobin < 100g/L an | | -depende | ent; and |

- 3 Patient has very low, low or intermediate risk MDS based on the WHO classification-based prognostic scoring system for myelodysplastic syndrome (WPSS); and
- 4 Other causes of anaemia such as B12 and folate deficiency have been excluded; and
- 5 Patient has a serum epoetin level of < 500 IU/L; and
- 6 The minimum necessary dose of epoetin would be used and will not exceed 80,000 iu per week.

Continuation - myelodysplasia*

Re-assessment required after 12 months

All of the following:

- 1 The patient's transfusion requirement continues to be reduced with epoetin treatment; and
- 2 Transformation to acute myeloid leukaemia has not occurred; and
- 3 The minimum necessary dose of epoetin would be used and will not exceed 80,000 iu per week.

Initiation - all other indications

Haematologist

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For use in patients where blood transfusion is not a viable treatment alternative.

Note: Indications marked with * are unapproved indications

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST |) | Generic |
| \$ | Per | Manufacturer |

EPOETIN BETA - Restricted see terms below

Note: Epoetin beta is considered a Discretionary Variance Pharmaceutical for epoetin alfa.

- Inj 2,000 iu in 0.3 ml syringe
- Inj 3,000 iu in 0.3 ml syringe
- Inj 4,000 iu in 0.3 ml syringe
- Inj 5,000 iu in 0.3 ml syringe
- Inj 6,000 iu in 0.3 ml syringe
- Inj 10,000 iu in 0.6 ml syringe

➡ Restricted (RS1661)

Initiation - chronic renal failure

All of the following:

- 1 Patient in chronic renal failure; and
- 2 Haemoglobin is less than or equal to 100g/L; and
- 3 Either:
 - 3.1 Both:
 - 3.1.1 Patient does not have diabetes mellitus; and
 - 3.1.2 Glomerular filtration rate is less than or equal to 30ml/min; or
 - 3.2 Both:
 - 3.2.1 Patient has diabetes mellitus; and
 - 3.2.2 Glomerular filtration rate is less than or equal to 45ml/min; and
- 4 Patient is on haemodialysis or peritoneal dialysis.

Initiation - myelodysplasia*

Re-assessment required after 12 months

All of the following:

- 1 Patient has a confirmed diagnosis of myelodysplasia (MDS); and
- 2 Has had symptomatic anaemia with haemoglobin < 100g/L and is red cell transfusion-dependent; and
- 3 Patient has very low, low or intermediate risk MDS based on the WHO classification-based prognostic scoring system for myelodysplastic syndrome (WPSS); and
- 4 Other causes of anaemia such as B12 and folate deficiency have been excluded; and
- 5 Patient has a serum epoetin level of < 500 IU/L; and
- 6 The minimum necessary dose of epoetin would be used and will not exceed 80,000 iu per week.

Continuation – myelodysplasia*

Re-assessment required after 2 months

All of the following:

- 1 The patient's transfusion requirement continues to be reduced with epoetin treatment; and
- 2 Transformation to acute myeloid leukaemia has not occurred; and
- 3 The minimum necessary dose of epoetin would be used and will not exceed 80,000 iu per week.

Initiation - all other indications

Haematologist.

For use in patients where blood transfusion is not a viable treatment alternative. *Note: Indications marked with * are unapproved indications.

Megaloblastic

FOLIC ACID

| Tab 0.8 mg - 1% DV Oct-18 to 2021 | 21.84 | 1,000 | Apo-Folic Acid |
|-----------------------------------|-------|-------|----------------|
| Tab 5 mg - 1% DV Oct-18 to 2021 | | 500 | Apo-Folic Acid |
| Oral lig 50 mcg per ml | | 25 ml | Biomed |
| Inj 5 mg per ml, 10 ml vial | | | |

| | Dries | | Drand ar |
|---|------------------------------|-----------|-------------------------------|
| | Price (ex man. excl. GST) | | Brand or Generic |
| | \$ | Per | Manufacturer |
| Antifibrinolytics, Haemostatics and Local Scleros | ants | | |
| ALUMINIUM CHLORIDE – Restricted see terms below | | | |
| | | | e.g. Driclor |
| → Restricted (RS1500) | | | |
| Initiation | | | |
| For use as a haemostatis agent. | | | |
| APROTININ – Restricted see terms below | | | |
| Inj 10,000 kIU per ml (equivalent to 200 mg per ml), 50 ml vial → Restricted (RS1332) | | | |
| Initiation | | | |
| Cardiac anaesthetist | | | |
| Either: | | | |
| Paediatric patient undergoing cardiopulmonary bypass proce Adult patient undergoing cardiac surgical procedure where th adverse effects of the drug. | | sive blee | ding outweighs the potential |
| ELTROMBOPAG – Restricted see terms below | | | |
| Tab 25 mg | 1,550.00 | 28 | Revolade |
| ↓ Tab 50 mg | 3,100.00 | 28 | Revolade |
| → Restricted (RS1648) | | | |
| Initiation – idiopathic thrombocytopenic purpura - post-splenec | tomy | | |
| Haematologist Re-assessment required after 6 weeks | | | |
| All of the following: | | | |
| 1 Patient has had a splenectomy; and | | | |
| 2 Two immunosuppressive therapies have been trialled and fai | led after therapy of 3 m | onths eac | h (or 1 month for rituximab): |
| and | | | |
| 3 Any of the following: | | | |
| 3.1 Patient has a platelet count of 20,000 to 30,000 platel | ets per microlitre and ha | as eviden | ce of significant |
| mucocutaneous bleeding; or | | | • |
| 3.2 Patient has a platelet count of less than or equal to 20 | ,000 platelets per micro | litre and | has evidence of active |
| bleeding; or | | | |
| 3.3 Patient has a platelet count of less than or equal to 10 | | litre. | |
| Initiation – idiopathic thrombocytopenic purpura - preparation f | or splenectomy | | |
| Haematologist | | | |
| Limited to 6 weeks treatment | a atamu / | | |
| The patient requires eltrombopag treatment as preparation for splen Continuation – idiopathic thrombocytopenic purpura - post-sple | | | |
| Haematologist | enectomy | | |
| Re-assessment required after 12 months | | | |
| The patient has obtained a response (see Note) from treatment durin | ng the initial approval or | subseau | ent renewal periods and |
| further treatment is required. | ig ale illusi approval el | ousooqu | ent fononal ponodo and |
| Note: Response to treatment is defined as a platelet count of > 30,0 | 000 platelets per microlit | re | |
| Initiation - idiopathic thrombocytopenic purpura contraindicate | | | |
| Haematologist | • | | |
| Re-assessment required after 3 months | | | |
| All of the following: | | | |

All of the following:

1 Patient has a significant and well-documented contraindication to splenectomy for clinical reasons; and

| Price | | Brand or | |
|--------------------|-----|--------------|--|
| (ex man. excl. GST | | Generic | |
| \$ | Per | Manufacturer | |

continued...

- 2 Two immunosuppressive therapies have been trialled and failed after therapy of 3 months each (or 1 month for rituximab); and
- 3 Either:
 - 3.1 Patient has immune thrombocytopenic purpura* with a platelet count of less than or equal to 20,000 platelets per microliter; or
 - 3.2 Patient has immune thrombocytopenic purpura* with a platelet count of 20,000 to 30,000 platelets per microlitre and significant muccoutaneous bleeding.

Continuation - idiopathic thrombocytopenic purpura contraindicated to splenectomy

Haematologist

Re-assessment required after 12 months

All of the following:

- 1 The patient's significant contraindication to splenectomy remains; and
- 2 The patient has obtained a response from treatment during the initial approval period; and
- 3 Patient has maintained a platelet count of at least 50,000 platelets per microlitre on treatment; and
- 4 Further treatment with eltrombopag is required to maintain response.

Initiation - severe aplastic anaemia

Haematologist

Re-assessment required after 3 months

Both:

- 1 Two immunosuppressive therapies have been trialled and failed after therapy of at least 3 months duration; and
- 2 Either:
 - 2.1 Patient has severe aplastic anaemia with a platelet count of less than or equal to 20,000 platelets per microliter; or
 - 2.2 Patient has severe aplastic anaemia with a platelet count of 20,000 to 30,000 platelets per microlitre and significant mucocutaneous bleeding.

Continuation - severe aplastic anaemia

Haematologist

Re-assessment required after 12 months Both:

- 1 The patient has obtained a response from treatment of at least 20,000 platelets per microlitre above baseline during the initial approval period; and
- 2 Platelet transfusion independence for a minimum of 8 weeks during the initial approval period.

FERRIC SUBSULFATE

Gel 25.9% Soln 500 ml

POLIDOCANOL

Inj 0.5%, 30 ml vial

SODIUM TETRADECYL SULPHATE

Inj 3%, 2 ml ampoule

THROMBIN

Powder

TRANEXAMIC ACID

| Tab 500 mg - 1% DV May-20 to 2022 | 5 60 | Mercury Pharma |
|---|------|----------------|
| Inj 100 mg per ml, 5 ml ampoule - 1% DV Sep-18 to 2021 | | Tranexamic-AFT |
| Inj 100 mg per ml, 10 ml ampoule - 1% DV Sep-18 to 2021 | 5 5 | Tranexamic-AFT |

Anticoagulant Reversal Agents

| IDA | ARUCIZUMAB – Restricted see terms on the next page | | |
|-----|--|---|----------|
| t | Inj 50 mg per ml, 50 ml vial4,250.00 | 2 | Praxbind |

Products with Hospital Supply Status (HSS) are in **bold**

Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| Price | | Brand or |
|-------------------|-----|--------------|
| (ex man. excl. GS | | Generic |
| \$ | Per | Manufacturer |

➡ Restricted (RS1535)

Initiation

For the reversal of the anticoagulant effects of dabigatran when required in situations of life-threatening or uncontrolled bleeding, or for emergency surgery or urgent procedures.

Blood Factors

| EF | TRENONACOG ALFA [RECOMBINANT FACTOR IX] - Restricted see terms below | N | |
|----|--|---|----------|
| t | Inj 250 iu vial | 1 | Alprolix |
| | Inj 500 iu vial | | Alprolix |
| t | Inj 1,000 iu vial2,450.00 | 1 | Alprolix |
| t | Inj 2,000 iu vial4,900.00 | 1 | Alprolix |
| t | | 1 | Alprolix |
| | | | |

Restricted (RS1684)

Initiation

For patients with haemophilia B receiving prophylaxis treatment. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group

EPTACOG ALFA [RECOMBINANT FACTOR VIIA] - Restricted see terms below

| t | Inj 1 mg syringe | 1,178.30 | 1 | NovoSeven RT |
|---|------------------|----------|---|--------------|
| t | Inj 2 mg syringe | 2,356.60 | 1 | NovoSeven RT |
| | Inj 5 mg syringe | | 1 | NovoSeven RT |
| | Inj 8 mg syringe | | 1 | NovoSeven RT |
| | | | | |

➡ Restricted (RS1704)

Initiation

For patients with haemophilia. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group. Rare Clinical Circumstances Brand of bypassing agent for > 14 days predicted use. Access to funded treatment for > 14 days predicted use is by named patient application to the Haemophilia Treaters Group, subject to access criteria

FACTOR EIGHT INHIBITOR BYPASSING FRACTION – Restricted see terms below

| t | Inj 500 U | 1 | FEIBA NF |
|---|---------------------|---|----------|
| | Inj 1,000 U2,630.00 | 1 | FEIBA NF |
| - | Inj 2,500 U6,575.00 | 1 | FEIBA NF |

Restricted (RS1705)

Initiation

For patients with haemophilia. Preferred Brand of bypassing agent for > 14 days predicted use. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group

MOROCTOCOG ALFA [RECOMBINANT FACTOR VIII] - Restricted see terms below

| Inj 250 iu prefilled syringe | 1 | Xyntha |
|--|---|--------|
| Inj 500 iu prefilled syringe | 1 | Xyntha |
| Inj 1,000 iu prefilled syringe | 1 | Xyntha |
| Inj 2,000 iu prefilled syringe2,300.00 | 1 | Xyntha |
| | 1 | Xyntha |
| Destricted (D01700) | | • |

→ Restricted (RS1706)

Initiation

For patients with haemophilia. Rare Clinical Circumstances Brand of short half-life recombinant factor VIII. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group, subject to criteria

| NO | NACOG GAMMA, [RECOMBINANT FACTOR IX] – Restricted see terms on the r | iext page | |
|----|--|-----------|---------|
| t | Inj 500 iu vial |) 1 | RIXUBIS |
| t | Inj 1,000 iu vial |) 1 | RIXUBIS |
| t | Inj 2,000 iu vial |) 1 | RIXUBIS |
| t | Inj 3,000 iu vial |) 1 | RIXUBIS |
| | | | |

t Item restricted (see \rightarrow above); **f** Item restricted (see \rightarrow below)

e.g. Brand indicates brand example only. It is not a contracted product.

| Pri | се | | Brand or |
|------------|----------|-----|--------------|
| (ex man. e | excl. GS | | Generic |
| \$ | 6 | Per | Manufacturer |

➡ Restricted (RS1679)

Initiation

For patients with haemophilia. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group

OCTOCOG ALFA [RECOMBINANT FACTOR VIII] (ADVATE) - Restricted see terms below

| t | Inj 250 iu vial | 210.00 | 1 | Advate |
|---|-------------------|----------|---|--------|
| t | Inj 500 iu vial | | 1 | Advate |
| | Inj 1,000 iu vial | | 1 | Advate |
| t | Inj 1,500 iu vial | | 1 | Advate |
| | Inj 2,000 iu vial | | 1 | Advate |
| t | Inj 3,000 iu vial | 2,520.00 | 1 | Advate |

→ Restricted (RS1707)

Initiation

For patients with haemophilia. Preferred Brand of short half-life recombinant factor VIII. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group

OCTOCOG ALFA [RECOMBINANT FACTOR VIII] (KOGENATE FS) - Restricted see terms below

| t | Inj 250 iu vial | 7.50 1 | Kogenate FS |
|---|------------------------|--------|-------------|
| t | Inj 500 iu vial | 5.00 1 | Kogenate FS |
| t | Inj 1,000 iu vial | 0.00 1 | Kogenate FS |
| t | Inj 2,000 iu vial | 0.00 1 | Kogenate FS |
| t | Inj 3,000 iu vial2,850 | 0.00 1 | Kogenate FS |

→ Restricted (RS1708)

Initiation

For patients with haemophilia. Rare Clinical Circumstances Brand of short half-life recombinant factor VIII. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group, subject to criteria

RURIOCTOCOG ALFA PEGOL [RECOMBINANT FACTOR VIII] - Restricted see terms below

| Inj 250 iu vial | | 1 | Adynovate |
|-----------------------|--------|---|-----------|
| Ini 500 iu vial | 600.00 | 1 | Adynovate |
| Inj 1,000 iu vial | | 1 | Advnovate |
| Inj 2,000 iu vial | | 1 | Adynovate |
| → Restricted (RS1682) | , | | , |

Initiation

For patients with haemophilia A receiving prophylaxis treatment. Access to funded treatment is managed by the Haemophilia Treaters Group in conjunction with the National Haemophilia Management Group

Vitamin K

| PHYTOMENADIONE | | | |
|--------------------------------|------|---|-------------|
| Inj 2 mg in 0.2 ml ampoule | 8.00 | 5 | Konakion MM |
| Inj 10 mg per ml, 1 ml ampoule | 9.21 | 5 | Konakion MM |

Antithrombotics

Anticoagulants

BIVALIRUDIN - Restricted see terms below

Inj 250 mg vial

```
→ Restricted (RS1181)
Initiation
Fither:
```

continued...

| | Price (ex man. excl. GST \$ | Per | Brand or Generic Manufacturer |
|--|-----------------------------------|------------|-------------------------------------|
| continued | | | |
| For use in heparin-induced thrombocytopaenia, heparin resista For use in patients undergoing endovascular procedures. | ance or heparin intole | rance; or | |
| CITRATE SODIUM | | | |
| Inj 4% (200 mg per 5 ml), 5 ml ampoule | | | |
| Inj 46.7% (1.4 g per 3 ml), 3 ml syringe Inj 46.7% (2.36 g per 5 ml), 5 ml ampoule | | | |
| DABIGATRAN | | | |
| Cap 75 mg | 76 36 | 60 | Pradaxa |
| Cap 110 mg | | 60 | Pradaxa |
| Cap 150 mg | | 60 | Pradaxa |
| DANAPAROID – Restricted see terms below | | | |
| Inj 750 u in 0.6 ml ampoule | | | |
| → Restricted (RS1182) | | | |
| nitiation | | | |
| or use in heparin-induced thrombocytopaenia, heparin resistance or | heparin intolerance. | | |
| DEFIBROTIDE - Restricted see terms below | | | |
| Inj 80 mg per ml, 2.5 ml ampoule | | | |
| → Restricted (RS1183) | | | |
| nitiation | | | |
| laematologist ?atient has moderate or severe sinusoidal obstruction syndrome as a | result of chemothera | ny or regi | mon-related toxicities |
| DEXTROSE WITH SODIUM CITRATE AND CITRIC ACID [ACID CIT | | | |
| Inj 24.5 mg with sodium citrate 22 mg and citric acid 7.3 mg per n | | 1 | |
| 100 ml bag | п, | | |
| | | | |
| Inj 20 mg in 0.2 ml syringe | | 10 | Clexane |
| Inj 40 mg in 0.4 ml ampoule | | | |
| Inj 40 mg in 0.4 ml syringe | | 10 | Clexane |
| Inj 60 mg in 0.6 ml syringe | | 10 | Clexane |
| Inj 80 mg in 0.8 ml syringe | | 10 | Clexane |
| Inj 100 mg in 1 ml syringe | | 10 | Clexane |
| Inj 120 mg in 0.8 ml syringe | | 10 | Clexane Clexane Forte |
| Inj 150 mg in 1 ml syringe | 133.20 | 10 | Clexane |
| , | | | Clexane Forte |
| Clexane Inj 120 mg in 0.8 ml syringe to be delisted 1 January 2021) | | | |
| Clexane Inj 150 mg in 1 ml syringe to be delisted 1 January 2021) | | | |
| ONDAPARINUX SODIUM - Restricted see terms below | | | |
| | | | |

- Inj 2.5 mg in 0.5 ml syringe
- Inj 7.5 mg in 0.6 ml syringe
- ➡ Restricted (RS1184)

Initiation

For use in heparin-induced thrombocytopaenia, heparin resistance or heparin intolerance.

| | Price | | Brand or |
|--|--------------------|----------|-------------------------|
| | (ex man. excl. GST |) Per | Generic Manufacturer |
| | \$ | Fei | Manulaciulei |
| HEPARIN SODIUM | | | |
| Inj 100 iu per ml, 250 ml bag Inj 1,000 iu per ml, 1 ml ampoule | 107.06 | 50 | Hospira |
| Inj 1,000 iu per ml, 5 ml ampoule – 1% DV Nov-18 to 2021 | | 50 | Pfizer |
| Inj 5,000 iu in 0.2 ml ampoule | | 50 | F 11201 |
| Inj 5,000 iu per ml, 1 ml ampoule | | 5 | Hospira |
| Inj 5,000 iu per ml, 5 ml ampoule – 1% DV Nov-18 to 2021 | | 50 | Pfizer |
| HEPARINISED SALINE | | | |
| Inj 10 iu per ml, 5 ml ampoule | | 50 | Pfizer |
| Inj 100 iu per ml, 2 ml ampoule | | | |
| Inj 100 iu per ml, 5 ml ampoule | | | |
| PHENINDIONE | | | |
| Tab 10 mg | | | |
| Tab 25 mg | | | |
| Tab 50 mg | | | |
| PROTAMINE SULPHATE | | | |
| Inj 10 mg per ml, 5 ml ampoule | | | |
| RIVAROXABAN | | | |
| Tab 10 mg | 83.10 | 30 | Xarelto |
| Tab 15 mg | | 28 | Xarelto |
| Tab 20 mg | | 28 | Xarelto |
| SODIUM CITRATE WITH SODIUM CHLORIDE AND POTASSIUM C | HI ORIDE | | |
| Inj 4.2 mg with sodium chloride 5.7 mg and potassium chloride 74 | - | | |
| per ml, 5,000 ml bag | no mog | | |
| WARFARIN SODIUM | | | |
| Tab 1 mg | | 100 | Marevan |
| Tab 2 mg | | | |
| Tab 3 mg | | 100 | Marevan |
| Tab 5 mg | 11.48 | 100 | Marevan |
| Antiplatelets | | | |
| ASPIRIN | | | |
| Tab 100 mg - 10% DV Nov-19 to 2022 | 1.95 | 90 | Ethics Aspirin EC |
| | 10.80 | 990 | Ethics Aspirin EC |
| Suppos 300 mg | | | |
| CLOPIDOGREL | | | |
| Tab 75 mg - 1% DV May-20 to 2022 | | 84 | Clopidogrel Multichem |
| DIPYRIDAMOLE | | • | |
| Tab 25 mg | | | |
| Tab long-acting 150 mg – 1% DV Oct-19 to 2022 | 10.90 | 60 | Pytazen SR |
| Inj 5 mg per ml, 2 ml ampoule | | | |
| EPTIFIBATIDE – Restricted see terms below | | | |
| Inj 2 mg per ml, 10 ml vial – 1% DV Nov-18 to 2021 | 138.75 | 1 | Integrilin |
| Inj 250 mcg per ml, 100 ml vial − 1% DV Nov-18 to 2021 | | 1 | Integrilin |
| → Restricted (RS1759) | | - | - J |
| Initiation | | | |
| Any of the following: | | | |
| | | | |

continued...

| | | Price excl. GS \$ | ST) | Per | Brand or Generic Manufacturer |
|---|---------------|-------------------------|---------|------------|-------------------------------------|
| continued | | | | | |
| For use in patients with acute coronary syndromes undergoin For use in patients with definite or strongly suspected intra-co For use in patients undergoing intra-cranial intervention. | | | | | |
| LYSINE ACETYLSALICYLATE [LYSINE ASPRIN] - Restricted see | e terms belo | w | | | |
| ↓ Inj 500 mg | | | | | e.g. Aspegic |
| ➡ Restricted (RS1689) | | | | | |
| Initiation | | | | | |
| Both: | | | | | |
| For use when an immediate antiplatelet effect is required pric cardiology procedure; and Administration of oral aspirin would delay the procedure. | or to an urge | ent interve | ention | al neu | iro-radiology or interventiona |
| PRASUGREL – Restricted see terms below | | | | | |
| ↓ Tab 5 mg | | 108.00 | | 28 | Effient |
| ↓ Tab 10 mg | | | | 28 | Effient |
| ➡ Restricted (RS1187) | | | | | |
| Initiation – Bare metal stents | | | | | |
| Limited to 6 months treatment | | | | | |
| Patient has undergone coronary angioplasty in the previous 4 weeks | s and is clop | idogrel-a | llergi |) . | |
| nitiation – Drug-eluting stents | | | | | |
| Limited to 12 months treatment | 4 | | | 1 - 11 | |
| Patient has had a drug-eluting cardiac stent inserted in the previous nitiation – Stent thrombosis | 4 weeks an | a is ciopi | aogre | aller | gic. |
| Patient has experienced cardiac stent thrombosis whilst on clopidog | rol | | | | |
| Initiation – Myocardial infarction | | | | | |
| Limited to 1 week treatment | | | | | |
| For short term use while in hospital following ST-elevated myocardia | I infarction. | | | | |
| Note: Clopidogrel allergy is defined as a history of anaphylaxis, urtic | caria, genera | alised ras | sh or a | asthma | a (in non-asthmatic patients |
| developing soon after clopidogrel is started and is considered unlikel | ly to be cau | sed by ar | ny oth | er trea | atment |
| TICAGRELOR – Restricted see terms below | | | | | |
| | | .90.00 | | 56 | Brilinta |
| → Restricted (RS1724) | | | | | |
| nitiation | | | | | |
| Restricted to treatment of acute coronary syndromes specifically for | | | | | |
| diagnosed with an ST-elevation or a non-ST-elevation acute coronar | y syndrome | e, and in v | whom | fibrine | olytic therapy has not been |
| given in the last 24 hours and is not planned. | | | | | |
| nitiation – thrombosis prevention post neurological stenting Re-assessment required after 12 months | | | | | |
| Both: | | | | | |
| Patient has had a neurological stenting procedure* in the last | 60 dave: a | hd | | | |
| 2 Either: | . 00 uays, ai | iu ii | | | |
| 2.1 Patient has demonstrated clopidogrel resistance using | the P2Y12 | (VerifyN | low) a | issav a | and requires antiplatelet |
| treatment with ticagrelor; or | , | | | seag (| |
| 2.2 Clopidogrel resistance has been demonstrated by the | occurrence | of a new | / cere | bral is | chemic event. |
| Continuation – thrombosis prevention post neurological stentin | | | | | |
| Re-assessment required after 12 months | - | | | | |
| Both: | | | | | |
| 1 Patient is continuing to benefit from treatment; and | | | | | |
| 2 Treatment continues to be clinically appropriate. | | | | | |

2 Treatment continues to be clinically appropriate.

Note: Indications marked with * are unapproved indications.

e.g. Brand indicates brand example only. It is not a contracted product.

| Price | | | Brand or |
|----------|-----------|----------|--------------|
| (ex man. | excl. GST |) Per | Generic |
| | \$ | Per | Manufacturer |

TICLOPIDINE

Tab 250 mg

Fibrinolytic Agents

ALTEPLASE

Inj 2 mg vial Inj 10 mg vial Inj 50 mg vial

TENECTEPLASE

lnj 50 mg vial

UROKINASE

Inj 5,000 iu vial Inj 10,000 iu vial Inj 50,000 iu vial Inj 100,000 iu vial Inj 500,000 iu vial

Colony-Stimulating Factors

Drugs Used to Mobilise Stem Cells

| PLERIXAFOR - Restricted see terms below Inj 20 mg per ml, 1.2 ml vial |
|--|
| Initiation – Autologous stem cell transplant |
| Haematologist |
| Limited to 3 days treatment |
| All of the following: |
| 5 |
| 1 Patient is to undergo stem cell transplantation; and |
| 2 Patient has not had a previous unsuccessful mobilisation attempt with plerixafor; and 2 Any of the following: |
| 3 Any of the following: |
| 3.1 Both: |
| 3.1.1 Patient is undergoing G-CSF mobilisation; and |
| 3.1.2 Either: |
| 3.1.2.1 Has a suboptimal peripheral blood CD34 count of less than or equal to 10×10^6 /L on day 5 after 4 days of G-CSF treatment; or |
| 3.1.2.2 Efforts to collect > 1 \times 10 ⁶ CD34 cells/kg have failed after one apheresis procedure; or |
| 3.2 Both: |
| 3.2.1 Patient is undergoing chemotherapy and G-CSF mobilisation; and 3.2.2 Any of the following: |
| 3.2.2.1 Both: |
| 3.2.2.1.1 Has rising white blood cell counts of > 5 $\times 10^9$ /L; and |
| 3.2.2.1.1 Has using while blood cer counts of > 3 x 10 /L, and 3.2.2.1.2 Has a suboptimal peripheral blood CD34 count of less than or equal to 10×10^6 /L; or |
| |
| 3.2.2.2 Efforts to collect > 1 × 10^6 CD34 cells/kg have failed after one apheresis procedure; or |
| 3.2.2.3 The peripheral blood CD34 cell counts are decreasing before the target has been received; or |
| 3.3 A previous mobilisation attempt with G-CSF or G-CSF plus chemotherapy has failed. |
| |

| Granulocyte Colony-Stimulating Factors | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|---------------|-------------------------------------|
| FILGRASTIM – Restricted see terms below ↓ Inj 300 mcg in 0.5 ml prefilled syringe – 1% DV May-19 to 2021 ↓ Inj 300 mcg in 1 ml vial ↓ Inj 480 mcg in 0.5 ml prefilled syringe – 1% DV Mar-19 to 2021 → Restricted (RS1188) Haematologist or oncologist | | 10 4 10 | Nivestim Neupogen Nivestim |
| PEGFILGRASTIM – Restricted see terms below ↓ Inj 6 mg per 0.6 ml syringe | 1,080.00 | 1 | Neulastim |

Initiation

For prevention of neutropenia in patients undergoing high risk chemotherapy for cancer (febrile neutropenia risk greater than or equal to 5%*).

Note: *Febrile neutropenia risk greater than or equal to 5% after taking into account other risk factors as defined by the European Organisation for Research and Treatment of Cancer (EORTC) guidelines

Fluids and Electrolytes

Intravenous Administration

| CALCIUM CHLORIDE | | |
|--|------|---------------------------------|
| Inj 100 mg per ml, 10 ml vial | | |
| Inj 100 mg per ml, 50 ml syringe | | e.g. Baxter |
| CALCIUM GLUCONATE | | |
| Inj 10%, 10 ml ampoule | | e.g. Max Health |
| COMPOUND ELECTROLYTES | | |
| Inj sodium 140 mmol/l, potassium 5 mmol/l, magnesium 1.5 mmol/l, chloride 98 mmol/l, acetate 27 mmol/l, gluconate 23 mmol/l, 500 ml | | |
| bag - 1% DV Jun-18 to 2021 | 0 18 | Plasma-Lyte 148 |
| chloride 98 mmol/l, acetate 27 mmol/l, gluconate 23 mmol/l, | | |
| 1,000 ml bag – 1% DV Jun-18 to 202127.24 | 4 12 | Plasma-Lyte 148 |
| COMPOUND ELECTROLYTES WITH GLUCOSE [DEXTROSE] | | |
| Inj sodium 140 mmol/l, 5 mmol/l potassium, 1.5 mmol/l magnesium, | | |
| 98 mmol/l chloride, 27 mmol/l acetate and 23 mmol/l gluconate, | | |
| glucose 23 mmol/l (5%), 1,000 ml bag - 1% DV Jun-18 to 2021 211.9 | 2 12 | Plasma-Lyte 148 & 5% Glucose |
| COMPOUND SODIUM LACTATE [HARTMANN'S SOLUTION] | | |
| Inj sodium 131 mmol/l with potassium 5 mmol/l, calcium 2 mmol/l, | | |
| bicarbonate 29 mmol/l, chloride 111 mmol/l, 500 ml bag – 1% DV | | |
| Jun-18 to 2021 | 0 18 | Baxter |
| Inj sodium 131 mmol/l with potassium 5 mmol/l, calcium 2 mmol/l, | | |
| bicarbonate 29 mmol/l, chloride 111 mmol/l, 1,000 ml bag – 1% DV Jun-18 to 2021 | 2 12 | Baxter |
| Juli-10 10 2021 | Z 12 | Daxlei |

| | Price | ~ | Brand or |
|--|--------------------------|----------|-------------------------|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer |
| UCOSE [DEXTROSE] | | | |
| Inj 5%, 1,000 ml bag – 1% DV Aug-18 to 2021 | 16.80 | 10 | Fresenius Kabi |
| lnj 5%, 100 ml bag – 1% DV Aug-18 to 2021 | | 50 | Fresenius Kabi |
| Inj 5%, 250 ml bag – 1% DV Aug-18 to 2021 | | 30 | Fresenius Kabi |
| lnj 5%, 50 ml bag – 1% DV Jun-18 to 2021 | | 60 | Baxter Glucose 5% |
| Inj 5%, 500 ml bag - 1% DV Aug-18 to 2021 | | 20 | Fresenius Kabi |
| Inj 10%, 1,000 ml bag - 1% DV Jun-18 to 2021 | | 12 | Baxter Glucose 10% |
| Inj 10%, 500 ml bag - 1% DV Jun-18 to 2021 | | 18 | Baxter Glucose 10% |
| Inj 50%, 10 ml ampoule - 1% DV Nov-20 to 2023 | | 5 | Biomed |
| Inj 50%, 500 ml bag - 1% DV Jun-18 to 2021 | | 18 | Baxter Glucose 50% |
| Inj 50%, 90 ml bottle - 1% DV Nov-20 to 2023 | | 1 | Biomed |
| UCOSE WITH POTASSIUM CHLORIDE | | | |
| Inj 10% glucose with 20 mmol/l potassium chloride, 500 ml ba | ig | | |
| UCOSE WITH POTASSIUM CHLORIDE AND SODIUM CHLO | RIDE | | |
| Inj 2.5% glucose with potassium chloride 20 mmol/l and sodiu 0.45%, 3,000 ml bag | ım chloride | | |
| Inj 10% glucose with potassium chloride 10 mmol/l and sodiul 15 mmol/l, 500 ml bag | m chloride | | |
| Inj 4% glucose with potassium chloride 20 mmol/l and sodium 0.18%, 1,000 ml bag - 1% DV Jun-18 to 2021 | | 12 | Baxter |
| Inj 5% glucose with potassium chloride 20 mmol/l and sodium | | | |
| 0.45%, 1,000 ml bag - 1% DV Jun-18 to 2021 | | 12 | Baxter |
| Inj 5% glucose with potassium chloride 20 mmol/l and sodium 0.9%, 1,000 ml bag – 1% DV Jun-18 to 2021 | | 12 | Baxter |
| UCOSE WITH SODIUM CHLORIDE | | | |
| Inj glucose 2.5% with sodium chloride 0.45%, 500 ml bag | | | |
| Inj 4% glucose and sodium chloride 0.18%, 1,000 ml bag - 1 | % DV | | |
| Jun-18 to 2021 | | 12 | Baxter |
| Inj 5% glucose and sodium chloride 0.45%, 1,000 ml bag -1 | | | |
| Jun-18 to 2021 | | 12 | Baxter |
| Inj 5% glucose and sodium chloride 0.9%, 1,000 ml bag – 1% Jun-18 to 2021 | | 12 | Baxter |
| TASSIUM CHLORIDE | | 12 | Daxiel |
| Inj 75 mg (1 mmol) per ml, 10 ml ampoule | | | |
| Inj 225 mg (3 mmol) per ml, 20 ml ampoule | | | |
| | | | |
| TASSIUM CHLORIDE WITH SODIUM CHLORIDE | 100 ml h a n | | |
| Inj 10 mmol potassium chloride with 0.29% sodium chloride, 1 | | 40 | Dautar |
| – 1% DV Jun-18 to 2021 Inj 20 mmol potassium chloride with 0.9% sodium chloride, 1, | | 48 | Baxter |
| – 1% DV Jun-18 to 2021 | | 12 | Baxter |
| Inj 40 mmol potassium chloride with 0.9% sodium chloride, 1, | 000 ml bag | | Buxton |
| - 1% DV Jun-18 to 2021 | | 12 | Baxter |
| Inj 40 mmol potassium chloride with 0.9% sodium chloride, 10 | | | |
| – 1% DV Jun-18 to 2021 | | 48 | Baxter |
| TASSIUM DIHYDROGEN PHOSPHATE | | | |
| Inj 1 mmol per ml, 10 ml ampoule | 151.80 | 10 | Hospira |
| NGER'S SOLUTION | | | |
| Inj sodium 147 mmol/l with potassium 4 mmol/l, calcium 2.2 m chloride 156 mmol/l, 1,000 ml bag | nmol/l, | | |
| DIUM ACETATE | | | |
| | | | |

| | Price (ex man. excl. GST) | | Brand or Generic | |
|--|------------------------------|----------|-------------------------|--|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer | |
| ODIUM BICARBONATE | | | | |
| Inj 8.4%, 10 ml vial | | | | |
| Inj 8.4%, 50 ml vial | | 1 | Biomed | |
| Inj 8.4%, 100 ml vial | 20.50 | 1 | Biomed | |
| ODIUM CHLORIDE | | | | |
| Inj 0.9%, 5 ml ampoule – 1% DV Dec-19 to 2022 | 2.80 | 20 | Fresenius Kabi | |
| Inj 0.9%, 10 ml ampoule - 1% DV Dec-19 to 2022 | | 50 | Fresenius Kabi | |
| Inj 0.9%, 3 ml syringe, non-sterile pack - 1% DV Sep-18 to 20 | 21 160.90 | 480 | BD PosiFlush | |
| → Restricted (RS1297) | | | | |
| nitiation | | | | |
| or use in flushing of in-situ vascular access devices only. | | | | |
| Inj 0.9%, 5 ml syringe, non-sterile pack – 1% DV Sep-18 to 20 | 21 162.91 | 480 | BD PosiFlush | |
| → Restricted (RS1297) | | | | |
| nitiation | | | | |
| or use in flushing of in-situ vascular access devices only. | | 100 | | |
| Inj 0.9%, 10 ml syringe, non-sterile pack – 1% DV Sep-18 to 2 | 021 170.35 | 480 | BD PosiFlush | |
| ► Restricted (RS1297) | | | | |
| nitiation or use in flushing of in-situ vascular access devices only. | | | | |
| с , | 5.00 | 00 | Francisco Kabi | |
| Inj 0.9%, 20 ml ampoule – 1% DV Dec-19 to 2022 | | 20 | Fresenius Kabi | |
| Inj 23.4% (4 mmol/ml), 20 ml ampoule | | 5 18 | Biomed Baxter | |
| Inj 0.45%, 500 ml bag Inj 3%, 1,000 ml bag | | 10 | Baxter | |
| Inj 0.9%, 50 ml bag | | 60 | Baxter | |
| Inj 0.9%, 100 ml bag | | 48 | Baxter | |
| Inj 0.9%, 250 ml bag | | 24 | Baxter | |
| Inj 0.9%, 500 ml bag | | 18 | Baxter | |
| Inj 0.9%, 1,000 ml bag | | 12 | Baxter | |
| Inj 1.8%, 500 ml bottle | | | | |
| ODIUM DIHYDROGEN PHOSPHATE [SODIUM ACID PHOSPHA | (TE) | | | |
| Inj 1 mmol per ml, 20 ml ampoule - 1% DV Oct-18 to 2021 | • | 5 | Biomed | |
| VATER | | | | |
| Inj 5 ml ampoule | 7.00 | 50 | InterPharma | |
| Inj 10 ml ampoule | | 50 | Pfizer | |
| Inj 20 ml ampoule | | 20 | Fresenius Kabi | |
| | 7.50 | 30 | InterPharma | |
| | 5.00 | 20 | Multichem | |
| Inj 250 ml bag | | | | |
| Inj 500 ml bag | | | | |
| Inj, 1,000 ml bag | 19.08 | 12 | Baxter | |
| Oral Administration | | | | |
| ALCIUM POLYSTYRENE SULPHONATE | | | | |
| Powder | | 300 g | Calcium Resonium | |
| COMPOUND ELECTROLYTES | | | | |
| Powder for oral soln - 1% DV Apr-20 to 2022 | 9.77 | 50 | Electral | |
| COMPOUND ELECTROLYTES WITH GLUCOSE [DEXTROSE] | | | | |
| Soln with electrolytes $(2 \times 500 \text{ ml}) - 1\% \text{ DV Nov-18 to 2021} \dots$ | 6.55 | 1,000 ml | Pedialyte - Bubblegun | |
| PHOSPHORUS | | | , | |
| | | | | |

t Item restricted (see → above); ↓ Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

BLOOD AND BLOOD FORMING ORGANS

| | ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------|----------------------|------|-------|-------------------------------------|
| POTASSIUM CHLORIDE Tab eff 548 mg (14 mmol) with chloride 285 mg (8 mmol) Tab long-acting 600 mg (8 mmol) – 1% DV Oct-18 to 2021 Oral lig 2 mmol per ml | | 8.9 | 0 | 200 | Span-K |
| SODIUM BICARBONATE Cap 840 mg | | 8.5 | 2 | 100 | Sodibic |
| SODIUM CHLORIDE Tab 600 mg Oral liq 2 mmol/ml | | | | | |
| SODIUM POLYSTYRENE SULPHONATE Powder – 1% DV Sep-18 to 2021 | | .84.6 | 5 | 454 g | Resonium A |
| Plasma Volume Expanders | | | | | |
| GELATINE, SUCCINYLATED Inj 4%, 500 ml bag – 1% DV Jun-18 to 2021 | | 120.0 | 0 | 10 | Gelofusine |

| | Price (ex man. excl. GST \$ | ^T) Per | Brand or Generic Manufacturer |
|---|-----------------------------------|-----------------------|-------------------------------------|
| | | | |
| Agents Affecting the Renin-Angiotensin System | | | |
| ACE Inhibitors | | | |
| CAPTOPRIL I Oral liq 5 mg per ml | | 95 ml | Capoten |
| ➡ Restricted (RS1263) | | | |
| Initiation | | | |
| Any of the following: | | | |
| For use in children under 12 years of age; or For use in tube-fed patients; or | | | |
| 3 For management of rebound transient hypertension followin | g cardiac surgery. | | |
| CILAZAPRIL | | | |
| Tab 0.5 mg - 1% DV Sep-19 to 2022 | 2.09 | 90 | Zapril |
| Tab 2.5 mg - 1% DV Feb-20 to 2022 | | 90 | Zapril |
| Tab 5 mg – 1% DV Feb-20 to 2022 | 8.35 | 90 | Zapril |
| ENALAPRIL MALEATE | | | . . |
| Tab 5 mg - 1% DV Jun-20 to 2022 | | 100 | Acetec |
| Tab 10 mg – 1% DV Jun-20 to 2022 Tab 20 mg – 1% DV Jun-20 to 2022 | | 100 100 | Acetec Acetec |
| | | 100 | ALELEL |
| Tab 5 mg - 1% DV Dec-18 to 2021 | 2 07 | 90 | Ethics Lisinopril |
| Tab 10 mg - 1% DV Dec-18 to 2021 | | 90 | Ethics Lisinopril |
| Tab 20 mg – 1% DV Dec-18 to 2021 | | 90 | Ethics Lisinopril |
| PERINDOPRIL | | | |
| Tab 2 mg | 3.75 | 30 | Apo-Perindopril |
| Tab 4 mg | 4.80 | 30 | Apo-Perindopril |
| QUINAPRIL | | | |
| Tab 5 mg - 1% DV Nov-18 to 2021 | | 90 | Arrow-Quinapril 5 |
| Tab 10 mg - 1% DV Nov-18 to 2021 | | 90 | Arrow-Quinapril 10 |
| Tab 20 mg – 1% DV Nov-18 to 2021 | 4.89 | 90 | Arrow-Quinapril 20 |
| ACE Inhibitors with Diuretics | | | |
| CILAZAPRIL WITH HYDROCHLOROTHIAZIDE - Restricted: Fo | , | | |
| Tab 5 mg with hydrochlorothiazide 12.5 mg | 10.18 | 100 | Apo-Cilazapril/ |
| (And Cilesonvill Indrachlarathiaside Tab E manuith hydrochlarathia | -ida 10 E ma ta ba dali | atad 1 Day | Hydrochlorothiazide |
| (Apo-Cilazapril/ Hydrochlorothiazide Tab 5 mg with hydrochlorothia | zide 12.3 my lo be dell | sieu i Dec | enidel 2020) |
| QUINAPRIL WITH HYDROCHLOROTHIAZIDE Tab 10 mg with hydrochlorothiazide 12.5 mg – 1% DV Dec-18 | to 2021 2.92 | 30 | Accuretic 10 |
| Tab 20 mg with hydrochlorothiazide 12.5 mg – 1% DV Dec-18 | | 30 | Accuretic 20 |
| Angiotensin II Antagonists | | | |
| CANDESARTAN CILEXETIL | | | |
| Tab 4 mg – 1% DV Sep-18 to 2021 | 1.90 | 90 | Candestar |
| Tab 8 mg – 1% DV Sep-18 to 2021 | 2.28 | 90 | Candestar |
| Tab 16 mg - 1% DV Sep-18 to 2021 | | 90 | Candestar |
| Tab 32 mg - 1% DV Sep-18 to 2021 | 6.39 | 90 | Candestar |

1 Item restricted (see \Rightarrow above); **1** Item restricted (see \Rightarrow below)

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e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | | Brand or |
|---|-------------------------------|------------|-----------------------------|
| | (ex man. excl. GST) | | Generic |
| | (on main onon alor) \$ | Per | Manufacturer |
| | Ŧ | | |
| LOSARTAN POTASSIUM | | | |
| Tab 12.5 mg | 1.39 | 84 | Losartan Actavis |
| Tab 25 mg | | 84 | Losartan Actavis |
| 5 | | • · | |
| Tab 50 mg | | 84 | Losartan Actavis |
| Tab 100 mg | 2.31 | 84 | Losartan Actavis |
| Angiotensin II Antagonists with Diuretics | | | |
| • • | | | |
| LOSARTAN POTASSIUM WITH HYDROCHLOROTHIAZIDE | | | |
| Tab 50 mg with hydrochlorothiazide 12.5 mg - 1% DV Jan-1 | 9 to 2021 1.88 | 30 | Arrow-Losartan & |
| | | | Hydrochlorothiazide |
| | | | • |
| | | | |
| Angiotensin II Antagonists with Neprilysin Inhibit | itors | | |
| SACUBITRIL WITH VALSARTAN - Restricted see terms below | | | |
| Tab 24.3 mg with valsartan 25.7 mg | | 56 | Entresto 24/26 |
| Tab 48.6 mg with valsartan 51.4 mg | | 56 | Entresto 49/51 |
| | | | |
| Tab 97.2 mg with valsartan 102.8 mg | | 56 | Entresto 97/103 |
| ➡ Restricted (RS1738) | | | |
| Initiation | | | |
| Re-assessment required after 12 months | | | |
| • | | | |
| All of the following: | | | |
| Patient has heart failure; and | | | |
| 2 Any of the following: | | | |
| | | | |
| 2.1 Patient is in NYHA/WHO functional class II; or | | | |
| 2.2 Patient is in NYHA/WHO functional class III; or | | | |
| 2.3 Patient is in NYHA/WHO functional class IV; and | | | |
| | | | |
| 3 Either: | | | |
| 3.1 Patient has a documented left ventricular ejection fr | action (LVEF) of less than | n or equal | to 35%: or |
| 3.2 An ECHO is not reasonably practical, and in the opi | . , | | |
| | mon or the treating practi | | patient would benefit from |
| treatment; and | | | |
| 4 Patient is receiving concomitant optimal standard chronic h | eart failure treatments. | | |
| Continuation | | | |
| | | | |
| Re-assessment required after 12 months | | | |
| The treatment remains appropriate and the patient is benefiting fro | om treatment. | | |
| Note: Due to the angiotensin II receptor blocking activity of sacub | itril with valsartan it shoul | d not be c | co-administered with an ACE |
| inhibitor or another ARB. | | | |
| | | | |
| Alpha-Adrenoceptor Blockers | | | |
| | | | |
| DOXAZOSIN | | | |
| Tab 2 mg | 6 75 | 500 | Apo-Doxazosin |
| | | | • |
| Tab 4 mg | 9.09 | 500 | Apo-Doxazosin |
| PHENOXYBENZAMINE HYDROCHLORIDE | | | |
| | | | |
| Cap 10 mg | | | |
| Inj 50 mg per ml, 1 ml ampoule | | | |
| Inj 50 mg per ml, 2 ml ampoule | | | |
| | | | |
| PHENTOLAMINE MESYLATE | | | |
| Inj 5 mg per ml, 1 ml ampoule | | | |
| | | | |
| Inj 10 mg per ml, 1 ml ampoule | | | |

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|-----|---------------------------------------|
| PRAZOSIN | | | |
| Tab 1 mg | 5.53 | 100 | Apo-Prazosin |
| Tab 2 mg | 7.00 | 100 | Apo-Prazosin |
| Tab 5 mg | 11.70 | 100 | Apo-Prazosin |
| ERAZOSIN | | | |
| Tab 1 mg | 0.59 | 28 | Actavis |
| Tab 2 mg | | 500 | Apo-Terazosin |
| Tab 5 mg | | 500 | Apo-Terazosin |
| Actavis Tab 1 mg to be delisted 1 October 2020) | | | ,.po |
| Antiarrhythmics | | | |
| DENOSINE | | | |
| Inj 3 mg per ml, 2 ml vial – 1% DV Feb-20 to 2022 | 62.73 | 6 | Adenocor |
| Inj 3 mg per ml, 10 ml vial → Restricted (RS1266) | | Ū | |
| nitiation for use in cardiac catheterisation, electrophysiology and MRI. | | | |
| AJMALINE – Restricted see terms below Inj 5 mg per ml, 10 ml ampoule → Restricted (RS1001) Cardiologist | | | |
| MIODARONE HYDROCHLORIDE | | | |
| Tab 100 mg - 1% DV Dec-19 to 2022 | | 30 | Aratac |
| Tab 200 mg - 1% DV Dec-19 to 2022 | | 30 | Aratac |
| Inj 50 mg per ml, 3 ml ampoule – 1% DV Feb-20 to 2022 TROPINE SULPHATE | | 10 | Max Health |
| Inj 600 mcg per ml, 1 ml ampoule – 1% DV Oct-18 to 2021 | 12.07 | 10 | Martindale |
| | 12.07 | 10 | wai unuale |
| | 7.00 | | |
| Tab 62.5 mcg – 1% DV Nov-19 to 2022 | | 240 | Lanoxin PG |
| Tab 250 mcg – 1% DV Nov-19 to 2022 Oral liq 50 mcg per ml Inj 250 mcg per ml, 2 ml vial | 15.20 | 240 | Lanoxin |
| NSOPYRAMIDE PHOSPHATE | | | |
| Cap 100 mg | | | |
| LECAINIDE ACETATE | | | |
| Tab 50 mg - 1% DV Feb-20 to 2022 | | 60 | Flecainide BNM |
| Cap long-acting 100 mg - 1% DV Dec-19 to 2022 | | 90 | Flecainide Controlled Release Teva |
| Cap long-acting 200 mg - 1% DV Dec-19 to 2022 | 61.06 | 90 | Flecainide Controlled Release Teva |
| Inj 10 mg per ml, 15 ml ampoule | | 5 | Tambocor |
| VABRADINE - Restricted see terms below | | | |
| ↓ Tab 5 mg → Restricted (RS1566) | | | |
| nitiation | | | |
| | | | |

| (6 | P ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|--------------|----------------------|--------|-------------|-------------------------------------|
| continued | | | | | |
| Patient is indicated for computed tomography coronary angiograph Either: | ny; and | | | | |
| 2.1 Patient has a heart rate of greater than 70 beats per minute or2.2 Patient is unable to tolerate beta blockers. | e while | takin | g a ma | ximally tol | erated dose of beta blocker; |
| MEXILETINE HYDROCHLORIDE | | | | | |
| Cap 150 mg | 1 | 62.0 | D | 100 | Mexiletine Hydrochloride USP |
| Cap 250 mg | 2 | 202.0 | D | 100 | Mexiletine Hydrochloride USP |
| PROPAFENONE HYDROCHLORIDE | | | | | |

Tab 150 mg

Antihypotensives

MIDODRINE - Restricted see terms below

- ↓ Tab 2.5 mg
- → Restricted (RS1427)

Initiation

Patient has disabling orthostatic hypotension not due to drugs.

Beta-Adrenoceptor Blockers

| ATENOLOL | | |
|---|--------|-------------------|
| Tab 50 mg - 1% DV Sep-18 to 2021 | 500 | Mylan Atenolol |
| Tab 100 mg - 1% DV Sep-18 to 20217.30 | 500 | Mylan Atenolol |
| Oral liq 5 mg per ml21.25 | 300 ml | Atenolol-AFT |
| BISOPROLOL FUMARATE | | |
| Tab 2.5 mg | 90 | Bosvate |
| Tab 5 mg | 90 | Bosvate |
| Tab 10 mg | 90 | Bosvate |
| CARVEDILOL | | |
| Tab 6.25 mg | 60 | Carvedilol Sandoz |
| Tab 12.5 mg2.30 | 60 | Carvedilol Sandoz |
| Tab 25 mg | 60 | Carvedilol Sandoz |
| CELIPROLOL | | |
| Tab 200 mg | 180 | Celol |
| ESMOLOL HYDROCHLORIDE | | |
| Inj 10 mg per ml, 10 ml vial | | |
| | | |
| | | |
| Tab 50 mg | 100 | Presolol |
| Tab 100 mg - 1% DV Sep-20 to 202411.36 14.50 | 100 | Trandate |
| Tab 200 mg - 1% DV Sep-20 to 2024 | 100 | Presolol |
| 1 ab 200 mg - 1 /8 DV Sep-20 to 2024 | 100 | Trandate |
| Inj 5 mg per ml, 20 ml ampoule | | Tandate |
| (Presolol Tab 100 mg to be delisted 1 September 2020) | | |
| | | |

(Presolol Tab 200 mg to be delisted 1 September 2020)

| | Price | | Brand or |
|--|--------------------|-----|----------------------|
| | (ex man. excl. GST | | Generic |
| | \$ | Per | Manufacturer |
| METOPROLOL SUCCINATE | | | |
| Tab long-acting 23.75 mg | 1.03 | 30 | Betaloc CR |
| Tab long-acting 47.5 mg | 1.25 | 30 | Betaloc CR |
| Tab long-acting 95 mg | 1.99 | 30 | Betaloc CR |
| Tab long-acting 190 mg | 3.00 | 30 | Betaloc CR |
| METOPROLOL TARTRATE | | | |
| Tab 50 mg - 1% DV Oct-18 to 2021 | 5.66 | 100 | Apo-Metoprolol |
| Tab 100 mg - 1% DV Oct-18 to 2021 | 7.55 | 60 | Apo-Metoprolol |
| Tab long-acting 200 mg | 23.40 | 28 | Slow-Lopresor |
| Inj 1 mg per ml, 5 ml vial - 1% DV Feb-19 to 31 Jan 2022 | 29.50 | 5 | Metroprolol IV Mylan |
| NADOLOL | | | |
| Tab 40 mg - 1% DV Oct-18 to 2021 | | 100 | Apo-Nadolol |
| Tab 80 mg - 1% DV Oct-18 to 2021 | | 100 | Apo-Nadolol |
| PINDOLOL | | | |
| Tab 5 mg - 1% DV Oct-18 to 2021 | | 100 | Apo-Pindolol |
| Tab 10 mg - 1% DV Oct-18 to 2021 | | 100 | Apo-Pindolol |
| Tab 15 mg - 1% DV Oct-18 to 2021 | | 100 | Apo-Pindolol |
| PROPRANOLOL | | | - |
| Tab 10 mg - 1% DV Oct-18 to 2021 | 4.64 | 100 | Apo-Propranolol |
| Tab 40 mg - 1% DV Oct-18 to 2021 | | 100 | Apo-Propranolol |
| Cap long-acting 160 mg | | 100 | Cardinol LA |
| Oral liq 4 mg per ml | | | |
| Inj 1 mg per ml, 1 ml ampoule | | | |
| SOTALOL | | | |
| Tab 80 mg - 1% DV Oct-19 to 2022 | | 500 | Mylan |
| Tab 160 mg - 1% DV Oct-19 to 2022 | | 100 | Mylan |
| | | | - |
| | | | |

Tab 10 mg

Calcium Channel Blockers

Dihydropyridine Calcium Channel Blockers

AMLODIPINE

| Tab 2.5 mg 1.72 Tab 5 mg 3.33 Tab 10 mg 4.40 | 100 250 250 | Apo-Amlodipine Apo-Amlodipine Apo-Amlodipine |
|--|-------------------|--|
| FELODIPINE | | |
| Tab long-acting 2.5 mg - 1% DV Sep-18 to 2021 1.45 | 30 | Plendil ER |
| Tab long-acting 5 mg - 1% DV Dec-18 to 2021 | 90 | Felo 5 ER |
| Tab long-acting 10 mg - 1% DV Dec-18 to 2021 | 90 | Felo 10 ER |

ISRADIPINE

Tab 2.5 mg Cap 2.5 mg

NICARDIPINE HYDROCHLORIDE - Restricted see terms below

Inj 2.5 mg per ml, 10 ml vial

➡ Restricted (RS1699)

Initiation

Anaesthetist, intensivist, cardiologist or paediatric cardiologist Any of the following:

continued...

| | Price (ex man. exc \$ | l. GST) Per | Brand or Generic Manufacturer |
|--|-----------------------------|----------------|-------------------------------------|
| continued 1 Patient has hypertension requiring urgent treatment with an | | t: or | |
| Patient has excessive ventricular afterload; or Patient is awaiting or undergoing cardiac surgery using ca | Ũ | | |
| NIFEDIPINE | | | |
| Tab long-acting 10 mg | | 63 60 | Adalat 10 |
| Tab long-acting 20 mg | | | Nyefax Retard |
| Tab long-acting 30 mg | | 14 30 | Adalat Oros |
| Tab long-acting 60 mg | | | Adalat Oros |
| Cap 5 mg | | | |
| NIMODIPINE | | | |
| Tab 30 mg - 1% DV Jul-20 to 2022 | 350 (| 00 100 | Nimotop |
| Inj 200 mcg per ml, 50 ml vial – 1% DV Jul-20 to 2022 | | | Nimotop |
| | | | |
| Other Calcium Channel Blockers | | | |
| DILTIAZEM HYDROCHLORIDE | | | |
| Tab 30 mg | 4.6 | 60 100 | Dilzem |
| Tab 60 mg | 8.5 | 50 100 | Dilzem |
| Cap long-acting 120 mg - 1% DV Oct-18 to 2021 | | 42 500 | Apo-Diltiazem CD |
| Cap long-acting 180 mg - 1% DV Oct-18 to 2021 | | 05 500 | Apo-Diltiazem CD |
| Cap long-acting 240 mg – 1% DV Oct-18 to 2021 Inj 5 mg per ml, 5 ml vial | | 76 500 | Apo-Diltiazem CD |
| PERHEXILINE MALEATE | | | |
| Tab 100 mg - 1% DV Oct-19 to 2022 | | 90 100 | Pexsig |
| VERAPAMIL HYDROCHLORIDE | | | |
| Tab 40 mg | 7 (| 01 100 | Isoptin |
| Tab 80 mg | | | Isoptin |
| Tab long-acting 120 mg | | | Isoptin SR |
| Tab long-acting 240 mg | | | Isoptin SR |
| | 25.0 | | Verpamil SR |
| Inj 2.5 mg per ml, 2 ml ampoule | | | Isoptin |
| Verpamil SR Tab long-acting 240 mg to be delisted 1 September | | | loopun |
| | / | | |
| Centrally-Acting Agents | | | |
| CLONIDINE | | | |
| Patch 2.5 mg, 100 mcg per day - 1% DV Nov-20 to 2023 | | 34 4 | Mylan |
| Patch 5 mg, 200 mcg per day - 1% DV Nov-20 to 2023 | | 18 4 | Mylan |
| Patch 7.5 mg, 300 mcg per day - 1% DV Nov-20 to 2023 | | | Mylan |
| CLONIDINE HYDROCHLORIDE | | | |
| | | | . |

METHYLDOPA

Clonidine BNM

Methyldopa Mylan

Catapres

Medsurge

112

100

10

100

| | Price (ex man. excl. \$ | GST) Per | Brand or Generic Manufacturer |
|--|-------------------------------|------------------------|--|
| Diuretics | | | |
| Loop Diuretics | | | |
| 3UMETANIDE Tab 1 mg Inj 500 mcg per ml, 4 ml vial FUROSEMIDE [FRUSEMIDE] | 16.36 | 6 100 | Burinex |
| Tab 40 mg - 1% DV Dec-19 to 2021 Tab 500 mg - 1% DV Mar-19 to 2021 Oral liq 10 mg per ml - 1% DV Jan-20 to 2022 Inj 10 mg per ml, 2 ml ampoule - 1% DV Oct-19 to 2022 Inj 10 mg per ml, 25 ml ampoule - 1% DV Jan-20 to 2022 | |) 50) 30 ml 5 5 | Apo-Furosemide Urex Forte Lasix Frusemide-Claris Lasix |
| Osmotic Diuretics | | | |
| VANNITOL Inj 10%, 1,000 ml bag – 1% DV Jun-18 to 2021 Inj 20%, 500 ml bag – 1% DV Jun-18 to 2021 | | | Baxter Baxter |
| Potassium Sparing Combination Diuretics | | | |
| AMILORIDE HYDROCHLORIDE WITH FUROSEMIDE Tab 5 mg with furosemide 40 mg AMILORIDE HYDROCHLORIDE WITH HYDROCHLOROTHIAZIDE Tab 5 mg with hydrochlorothiazide 50 mg | | | |
| Potassium Sparing Diuretics | | | |
| MILORIDE HYDROCHLORIDE | | | |
| Tab 5 mg Oral liq 1 mg per ml | |) 25 ml | Biomed |
| EPLERENONE - Restricted see terms below Tab 25 mg - 1% DV Sep-18 to 2021 Tab 50 mg - 1% DV Dec-18 to 2021 → Restricted (RS1640) nitiation | | | Inspra Inspra |
| 30th: 1 Patient has heart failure with ejection fraction less than 40%; 2 Either: | | | |
| 2.1 Patient is intolerant to optimal dosing of spironolactore2.2 Patient has experienced a clinically significant adverse | | ptimal dosing | of spironolactone. |
| SPIRONOLACTONE Tab 25 mg Tab 100 mg Oral lig 5 mg per ml - 1% DV Nov-19 to 2022 | |) 100 | Spiractin Spiractin Biomed |
| Thiazide and Related Diuretics | | | 2.01104 |
| BENDROFLUMETHIAZIDE [BENDROFLUAZIDE] | | | |
| Tab 2.5 mg Tab 5 mg | | | Arrow-Bendrofluazide Arrow-Bendrofluazide |

I tem restricted (see \rightarrow above); I tem restricted (see \rightarrow below)

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e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | | Brand or |
|---|---------------------|-------|-------------------|
| | (ex man. excl. GST) | | Generic |
| | \$ | Per | Manufacturer |
| CHLOROTHIAZIDE | | | |
| Oral liq 50 mg per ml | 26.00 | 25 ml | Biomed |
| | 20100 | | Diomiou |
| | 0.50 | 50 | llumeten |
| Tab 25 mg - 1% DV Dec-19 to 2022 | | 50 | Hygroton |
| NDAPAMIDE | | | |
| Tab 2.5 mg - 1% DV Nov-20 to 2023 | 10.45 | 90 | Dapa-Tabs |
| METOLAZONE | | | |
| Tab 5 mg | | | |
| Lipid-Modifying Agents | | | |
| , | | | |
| Fibrates | | | |
| BEZAFIBRATE | | | |
| Tab 200 mg – 1% DV Dec-18 to 2021 | | 90 | Bezalip |
| Tab long-acting 400 mg - 1% DV Dec-18 to 2021 | | 30 | Bezalip Retard |
| GEMFIBROZIL – Restricted: For continuation only | | | |
| → Tab 600 mg | 19 56 | 60 | Lipazil |
| Lipazil Tab 600 mg to be delisted 1 January 2021) | | 00 | црагіі |
| | | | |
| HMG CoA Reductase Inhibitors (Statins) | | | |
| ATORVASTATIN | | | |
| Tab 10 mg - 1% DV Sep-18 to 2021 | 6.96 | 500 | Lorstat |
| Tab 20 mg - 1% DV Sep-18 to 2021 | 9.99 | 500 | Lorstat |
| Tab 40 mg - 1% DV Sep-18 to 2021 | 15.93 | 500 | Lorstat |
| Tab 80 mg - 1% DV Sep-18 to 2021 | 27.19 | 500 | Lorstat |
| PRAVASTATIN | | | |
| Tab 10 mg | | | |
| Tab 20 mg | 4.72 | 100 | Apo-Pravastatin |
| Tab 40 mg | 8.06 | 100 | Apo-Pravastatin |
| SIMVASTATIN | | | |
| Tab 10 mg – 1% DV Nov-20 to 2023 | | 90 | Simvastatin Mylan |
| Tab 20 mg – 1% DV Nov-20 to 2023 | | 90 | Simvastatin Mylan |
| Tab 40 mg – 1% DV Nov-20 to 2023 | | 90 | Simvastatin Mylan |
| Tab 80 mg - 1% DV Nov-20 to 2023 | | 90 | Simvastatin Mylan |
| Resins | | | |
| | | | |
| CHOLESTYRAMINE Rowder for oral lig 4 g | | | |
| Powder for oral liq 4 g | | | |
| COLESTIPOL HYDROCHLORIDE | | | |
| Grans for oral liq 5 g | | | |
| | | | |
| Selective Cholesterol Absorption Inhibitors | | | |
| Selective Cholesterol Absorption Inhibitors EZETIMIBE – Restricted see terms on the next page | | | |
| | | 30 | Ezetimibe Sandoz |

| | Price (ex man. excl. GS | | Brand or Generic Manufacturar |
|--|------------------------------------|----------------|-------------------------------------|
| | \$ | Per | Manufacturer |
| | | | |
| → Restricted (RS1005) nitiation | | | |
| All of the following: | | | |
| Patient has a calculated absolute risk of cardiova | coular disease of at least 15% or | or E voore: | and |
| 2 Patient's LDL cholesterol is 2.0 mmol/litre or grea | | lei 5 years, a | anu |
| 3 Any of the following: | | | |
| 3.1 The patient has rhabdomyolysis (defined a | as muscle aches and creatine kir | nasa mora th | an 10 × normal) when |
| treated with one statin: or | | | |
| 3.2 The patient is intolerant to both simvastati | n and atomastatin: or | | |
| 3.3 The patient has not reduced their LDL cho | | re with the us | se of the maximal tolerate |
| dose of atorvastatin. | | | |
| EZETIMIBE WITH SIMVASTATIN – Restricted see terr | ms below | | |
| Tab 10 mg with simvastatin 10 mg | | 30 | Zimybe |
| Tab 10 mg with simvastatin 20 mg | | 30 | Zimybe |
| Tab 10 mg with simvastatin 40 mg | | 30 | Zimybe |
| Tab 10 mg with simvastatin 80 mg | | 30 | Zimybe |
| → Restricted (RS1006) | | | , |
| Initiation | | | |
| All of the following: | | | |
| 1 Patient has a calculated absolute risk of cardiova | scular disease of at least 15% ov | /er 5 vears: a | and |
| 2 Patient's LDL cholesterol is 2.0 mmol/litre or grea | | | |
| 3 The patient has not reduced their LDL cholestero | I to less than 2.0 mmol/litre with | the use of the | e maximal tolerated dose |
| atorvastatin. | | | |
| Other Linid Medifying Agente | | | |
| Other Lipid-Modifying Agents | | | |
| ACIPIMOX | | | |
| Cap 250 mg | | | |
| NICOTINIC ACID | | | |
| Tab 50 mg | | 100 | Apo-Nicotinic Acid |
| Tab 500 mg | | 100 | Apo-Nicotinic Acid |
| - | | | |
| Nitrates | | | |
| | | | |
| | | | |
| Inj 1 mg per ml, 5 ml ampoule | | | |
| Inj 1 mg per ml, 10 ml ampoule Inj 1 mg per ml, 50 ml vial | | | |
| Inj 1 mg per mi, 50 mi viai Inj 5 mg per mi, 10 mi ampoule | 100.00 | 5 | Hospira |
| Oral pump spray, 400 mcg per dose | | 250 dose | Nitrolingual Pump Spra |
| Patch 25 mg, 5 mg per day | | 200 00se 30 | Nitroderm TTS 5 |
| Patch 50 mg, 10 mg per day | | 30 | Nitroderm TTS 10 |
| 1 aton 50 mg, 10 mg per uay | | 30 | |

ISOSORBIDE MONONITRATE

| Tab 20 mg - 1% DV Nov-20 to 2023 19.55 100 | Ismo-20 |
|--|----------------|
| Tab long-acting 40 mg - 1% DV Nov-20 to 2023 | Ismo 40 Retard |
| Tab long-acting 60 mg - 1% DV Nov-20 to 2023 | Duride |

Other Cardiac Agents

LEVOSIMENDAN - Restricted see terms on the next page

Inj 2.5 mg per ml, 5 ml vial

Inj 2.5 mg per ml, 10 ml vial

t Item restricted (see → above); t Item restricted (see → below) e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | | Brand or |
|----------|-----------|-----|--------------|
| (ex man. | excl. GST | | Generic |
| | \$ | Per | Manufacturer |

→ Restricted (RS1007)

Initiation – Heart transplant

Either:

- 1 For use as a bridge to heart transplant, in patients who have been accepted for transplant; or
- 2 For the treatment of heart failure following heart transplant.

Initiation – Heart failure

Cardiologist or intensivist

For the treatment of severe acute decompensated heart failure that is non-responsive to dobutamine.

Sympathomimetics

| ADF | RENALINE | | | |
|-----|---|----------------|---------|------------------------------------|
| | Inj 1 in 1,000, 1 ml ampoule | 4.98 10.76 | 5 | Aspen Adrenaline DBL Adrenaline |
| | Inj 1 in 1,000, 30 ml vial | | | |
| | Inj 1 in 10,000, 10 ml ampoule | 49.00 27.00 | 10 5 | Aspen Adrenaline Hospira |
| | Inj 1 in 10,000, 10 ml syringe | 27.00 | 5 | nospira |
| - | BUTAMINE | | | |
| | Inj 12.5 mg per ml, 20 ml ampoule - 1% DV Jan-19 to 2021 | 61.13 | 5 | Dobutamine-hameIn |
| | PAMINE HYDROCHLORIDE Inj 40 mg per ml, 5 ml ampoule – 1% DV Sep-18 to 2021 | 29.73 | 10 | Max Health Ltd |
| | IEDRINE | | | |
| | Inj 3 mg per ml, 10 ml syringe | 00.00 | 10 | Max Health |
| | Inj 30 mg per ml, 1 ml ampoule – 1% DV Oct-20 to 2023 | 30.63 | 10 | Max Health |
| | PRENALINE [ISOPROTERENOL] Inj 200 mcg per ml, 1 ml ampoule | | | |
| | Inj 200 mcg per ml, 5 ml ampoule | | | |
| MET | TARAMINOL | | | |
| | Inj 0.5 mg per ml, 10 ml syringe | | | |
| | Inj 0.5 mg per ml, 20 ml syringe Inj 0.5 mg per ml, 5 ml syringe | | | |
| | Inj 1 mg per ml, 1 ml ampoule | | | |
| | Inj 1 mg per ml, 10 ml syringe | | | |
| | Inj 10 mg per ml, 1 ml ampoule - 1% DV Jan-21 to 2023 | 55.20 | 10 | Torbay |
| | RADRENALINE | | | |
| | Inj 0.06 mg per ml, 100 ml bag | | | |
| | Inj 0.06 mg per ml, 50 ml syringe Inj 0.1 mg per ml, 100 ml bag | | | |
| | Inj 0.1 mg per ml, 50 ml syringe | | | |
| | Inj 0.12 mg per ml, 100 ml bag | | | |
| | Inj 0.12 mg per ml, 50 ml syringe Inj 0.16 mg per ml, 50 ml syringe | | | |
| | Inj 1 mg per ml, 100 ml bag | | | |
| | Inj 1 mg per ml, 4 ml ampoule – 1% DV Oct-19 to 2022 | 45.00 | 10 | Noradrenaline BNM |
| | ENYLEPHRINE HYDROCHLORIDE | | | |
| | Inj 10 mg per ml, 1 ml ampoule | 142.07 | 25 | Neosynephrine HCL |
| | | | | |

| | | | D |
|--|------------------------------|-----------|-----------------------------|
| | Price (ex man. excl. GST) | | Brand or Generic |
| | (ex man. excl. GST) | Per | Manufacturer |
| | | | |
| Vasodilators | | | |
| ALPROSTADIL HYDROCHLORIDE | 1 765 50 | 5 | Prostin VR |
| Inj 500 mcg per ml, 1 ml ampoule - 1% DV Dec-18 to 2021 | | 5 | FIOSUII VN |
| DIAZOXIDE Inj 15 mg per ml, 20 ml ampoule | | | |
| HYDRALAZINE HYDROCHLORIDE | | | |
| | | | |
| → Restricted (RS1008) | | | |
| nitiation | | | |
| Either: | | | |
| For the treatment of refractory hypertension; or For the treatment of heart failure, in combination with a nitra ACE inhibitors and/or angiotensin receptor blockers. | te, in patients who are int | olerant c | or have not responded to |
| Inj 20 mg ampoule | | 5 | Apresoline |
| <i>A</i> ILRINONE | | | |
| Inj 1 mg per ml, 10 ml ampoule - 1% DV Sep-18 to 2021 | | 10 | Primacor |
| /INOXIDIL | | | |
| Tab 10 mg | 70.00 | 100 | Loniten |
| IICORANDIL | | | |
| Tab 10 mg - 1% DV Dec-19 to 2022 | | 60 | lkorel |
| Tab 20 mg - 1% DV Dec-19 to 2022 | | 60 | lkorel |
| PAPAVERINE HYDROCHLORIDE | | | |
| Inj 30 mg per ml, 1 ml vial | | | |
| Inj 12 mg per ml, 10 ml ampoule | 217.90 | 5 | Hospira |
| PENTOXIFYLLINE [OXPENTIFYLLINE] | | | |
| Tab 400 mg | | | |
| SODIUM NITROPRUSSIDE | | | |
| Inj 50 mg vial | | | |
| Endothelin Receptor Antagonists | | | |
| | | | |
| MBRISENTAN – Restricted see terms below I Tab 5 mg | 4 595 00 | 30 | Volibris |
| Tab 5 mg | | 30 | Volibris |
| Restricted (RS1621) | 4,000.00 | 00 | Volibrio |
| nitiation | | | |
| lither: | | | |
| 1 For use in patients with a valid Special Authority approval fo | r ambrisentan by the Pulr | monary A | Arterial Hypertension Panel |
| or | | | |
| 2 In-hospital stabilisations in emergency situations. | | | |
| BOSENTAN - Restricted see terms below | | | |
| Tab 62.5 mg - 1% DV Dec-18 to 2021 | | 60 | Bosentan Dr Reddy's |
| Tab 125 mg – 1% DV Dec-18 to 2021 | 141.00 | 60 | Bosentan Dr Reddy's |
| Restricted (RS1622) | | | |
| nitiation – Pulmonary arterial hypertension Re-assessment required after 6 months | | | |
| Either: | | | |
| | | | continued |
| | | | continued |

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | | Brand or |
|----|------------------|---------|--------------|
| (e | ex man. excl. GS | Generic | |
| | \$ | Per | Manufacturer |

- 1 All of the following:
 - 1.1 Patient has pulmonary arterial hypertension (PAH); and
 - 1.2 PAH is in Group 1, 4 or 5 of the WHO (Venice) clinical classifications; and
 - 1.3 PAH is at NYHA/WHO functional class II, III, or IV; and
 - 1.4 Any of the following:
 - 1.4.1 Both:
 - 1.4.1.1 Bosentan is to be used as PAH monotherapy; and
 - 1.4.1.2 Either:
 - 1.4.1.2.1 Patient is intolerant or contraindicated to sildenafil; or
 - 1.4.1.2.2 Patient is a child with idiopathic PAH or PAH secondary to congenital heart disease; or
 - 1.4.2 Both:
 - 1.4.2.1 Bosentan is to be used as PAH dual therapy; and
 - 1.4.2.2 Either:
 - 1.4.2.2.1 Patient has tried a PAH monotherapy for at least three months and failed to respond; or
 - 1.4.2.2.2 Patient deteriorated while on a PAH monotherapy; or
 - 1.4.3 Both:
 - 1.4.3.1 Bosentan is to be used as PAH triple therapy; and
 - 1.4.3.2 Any of the following:
 - 1.4.3.2.1 Patient is on the lung transplant list; or
 - 1.4.3.2.2 Patient is presenting acutely with idiopathic pulmonary arterial hypertension (IPAH) in New York Heart Association/World Health Organization (NYHA/WHO) Functional Class IV; or
 - 1.4.3.2.3 Patient is deteriorating rapidly to NYHA/WHO Functional Class IV who may be lung transplant recipients in the future, if their disease is stabilised; or
 - 1.4.3.2.4 Patient has PAH associated with the scleroderma spectrum of diseases (APAHSSD) who have no major morbidities and are deteriorating despite combination therapy; or
- 2 In-hospital stabilisation in emergency situations.

Continuation - Pulmonary arterial hypertension

- Re-assessment required after 6 months
- Any of the following:
 - 1 Both:
 - 1.1 Bosentan is to be used as PAH monotherapy; and
 - 1.2 Patient is stable or has improved while on bosentan; or
 - 2 Both:
 - 2.1 Bosentan is to be used as PAH dual therapy; and
 - 2.2 Patient has tried a PAH monotherapy for at least three months and either failed to respond or later deteriorated; or
 - 3 Both:
 - 3.1 Bosentan is to be used as PAH triple therapy; and
 - 3.2 Any of the following:
 - 3.2.1 Patient is on the lung transplant list; or
 - 3.2.2 Patient is presenting acutely with idiopathic pulmonary arterial hypertension (IPAH) in New York Heart Association/World Health Organization (NYHA/WHO) Functional Class IV; or
 - 3.2.3 Patient is deteriorating rapidly to NYHA/WHO Functional Class IV who may be lung transplant recipients in the future, if their disease is stabilised; or
 - 3.2.4 Patient has PAH associated with the scleroderma spectrum of diseases (APAHSSD) who have no major morbidities and are deteriorating despite combination therapy.

| | | Price excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------|---------------------------|--------------|-------------------------------------|
| Phosphodiesterase Type 5 Inhibitors | | | | |
| SILDENAFIL - Restricted see terms below Image: Tab 25 mg - 1% DV Sep-18 to 2021 Image: Tab 50 mg - 1% DV Sep-18 to 2021 Image: Tab 100 mg - 1% DV Sep-18 to 2021 | | 0.64 | 4 4 12 | Vedafil Vedafil Vedafil |
| Initiation – tablets Raynaud's Phenomenon All of the following: Patient has Raynaud's phenomenon; and Patient has severe digital ischaemia (defined as severe pain ulceration; digital ulcers; or gangrene); and Patient is following lifestyle management (proper body insula avoidance of sympathomimetic drugs); and Patient has persisting severe symptoms despite treatment wi | tion, avoidar | nce of cold e | xposure, | smoking cessation support, |
| contraindicated or not tolerated). Initiation – tablets Pulmonary arterial hypertension Any of the following: | | | | |
| All of the following: 1.1 Patient has pulmonary arterial hypertension (PAH); ar 1.2 Any of the following: 1.2.1 PAH is in Group 1 of the WHO (Venice) clinica 1.2.2 PAH is in Group 4 of the WHO (Venice) clinica 1.2.3 PAH is in Group 5 of the WHO (Venice) clinica | l classification | ons; or | | |
| 1.3 Any of the following: 1.3.1 PAH is in NYHA/WHO functional class II; or 1.3.2 PAH is in NYHA/WHO functional class III; or 1.3.3 PAH is in NYHA/WHO functional class IV; and | | Silo, and | | |
| 1.4 Either: 1.4.1 All of the following: 1.4.1.1 Patient has a pulmonary capillary wedge 1.4.1.2 Either: 1.4.1.2.1 Patient has a mean pulmonary ar | | , | | |
| 1.4.1.2.2 Patient is peri Fontan repair; and 1.4.1.3 Patient has a pulmonary vascular resista 240 International Units (dyn s cm-5); or | , | | | |
| 1.4.2 Testing for PCWP, PAPm, or PVR cannot be p capacity constraints; or 2 For use in neonatal units for persistent pulmonary hypertensi 3 In-hospital stabilisation in emergency situations. | | | | ng age, or health system |
| Initiation – tablets other conditions Any of the following: 1 For use in weaning patients from inhaled nitric oxide; or 2 For perioperative use in cardiac surgery patients; or | | | | |

- 3 For use in intensive care as an alternative to nitric oxide; or
- 4 For use in the treatment of erectile dysfunction secondary to spinal cord injury in patients being treated in a spinal unit.

e.g. Brand indicates brand example only. It is not a contracted product.

| Pr | ice | | Brand or |
|------------|------------|-----|--------------|
| (ex man. e | excl. GST) | | Generic |
| e e | \$ | Per | Manufacturer |

continued...

Initiation - injection

Both:

- 1 For use in the treatment of pulmonary hypertension in infants or children being treated in paediatric intensive care units and neonatal intensive care units when the enteral route is not accessible; and
- 2 Any of the following:
 - 2.1 For perioperative use following cardiac surgery; or
 - 2.2 For use in persistent pulmonary hypertension of the newborn (PPHN); or
 - 2.3 For use in congenital diaphragmatic hernia.

Prostacyclin Analogues

| EPOPROSTENOL – Restricted see terms below | | |
|---|---|---------|
| Inj 500 mcg vial | 1 | Veletri |
| Inj 1.5 mg vial | 1 | Veletri |
| ⇒ Restricted (RS1624) | | |

➡ Restricted (RS1624) Initiation

Either:

- 1 For use in patients with a valid Special Authority approval for epoprostenol by the Pulmonary Arterial Hypertension Panel; or
- 2 In-hospital stabilisation in emergency situations.

ILOPROST

| | Inj 50 mcg in 0.5 ml ampoule - 1% DV Jan-20 to 2022 |) 5 | Clinect |
|---|---|------|------------|
| t | Nebuliser soln 10 mcg per ml, 2 ml - 1% DV Jan-20 to 2022 |) 30 |) Ventavis |
| ⇒ | Restricted (RS1625) | | |

Initiation

Any of the following:

- 1 For use in patients with a valid Special Authority approval for iloprost by the Pulmonary Arterial Hypertension Panel; or
- 2 For diagnostic use in catheter laboratories; or
- 3 For use following mitral or tricuspid valve surgery; or
- 4 In-hospital stabilisation in emergency situations.

| | Price excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|---------------------------|------------|-------------------------------------|
| Anti-Infective Preparations | | | |
| Antibacterials | | | |
| HYDROGEN PEROXIDE Crm 1% | 8.56 | 15 g | Crystaderm |
| MAFENIDE ACETATE – Restricted see terms below ↓ Powder 50 g sachet → Restricted (RS1299) | | | |
| Initiation For the treatment of burns patients. MUPIROCIN Oint 2% | | | |
| SODIUM FUSIDATE [FUSIDIC ACID] Crm 2% – 1% DV May-19 to 2021 Oint 2% – 1% DV May-19 to 2021 | | 5 g 5 g | Foban Foban |
| SULFADIAZINE SILVER Crm 1% | .10.80 | 50 g | Flamazine |
| Antifungals | | | |
| AMOROLFINE Nail soln 5% – 1% DV Oct-20 to 2023 | .14.93 | 5 ml | MycoNail |
| CICLOPIROX OLAMINE Nail soln 8% – 1% DV Sep-18 to 2021 | 5.72 | 7 ml | Apo-Ciclopirox |
| CLOTRIMAZOLE Crm 1% → Soln 1% – Restricted: For continuation only | 0.70 | 20 g | Clomazol |
| ECONAZOLE NITRATE → Crm 1% - Restricted: For continuation only Foaming soln 1% | | | |
| KETOCONAZOLE Shampoo 2% - 1% DV Nov-20 to 2023 | 3.23 | 100 ml | Sebizole |
| METRONIDAZOLE Gel 0.75% | | | |
| MICONAZOLE NITRATE Crm 2% → Lotn 2% – Restricted: For continuation only | 0.74 | 15 g | Multichem |
| Tinc 2% NYSTATIN Crm 100,000 u per g | | | |
| Antiparasitics | | | |
| DIMETHICONE Lotn 4% - 1% DV Oct-19 to 2022 | 4.98 | 200 ml | healthE Dimethicone 4% Lotion |

| | - | | | Durand au |
|---|------|-----------------------|------------------|--|
| (ex | man. | ice excl. GS \$ | ST) Per | Brand or Generic Manufacturer |
| MALATHION [MALDISON] Lotn 0.5% Shampoo 1% | | | | |
| PERMETHRIN Crm 5% – 1% DV Nov-20 to 2023 Lotn 5% – 1% DV Nov-20 to 2023 | | | 30 g 30 ml | Lyderm A-Scabies |
| PHENOTHRIN Shampoo 0.5% | | | | |
| Antiacne Preparations | | | | |
| ADAPALENE Crm 0.1% Gel 0.1% | | | | |
| BENZOYL PEROXIDE Soln 5% | | | | |
| ISOTRETINOIN Cap 5 mg - 1% DV Oct-18 to 2021 Cap 10 mg - 1% DV Oct-18 to 2021 Cap 20 mg - 1% DV Oct-18 to 2021 | 1 | 3.34 | 60 120 120 | Oratane Oratane Oratane |
| TRETINOIN Crm 0.05% – 1% DV Jun-18 to 2021 | | | 50 g | ReTrieve |
| Antipruritic Preparations | | | | |
| CALAMINE Crm, aqueous, BP – 1% DV Nov-18 to 2021 | | .1.26 | 100 g | healthE Calamine Aqueous Cream BP |
| CROTAMITON Crm 10% – 1% DV Sep-18 to 2021 | | .3.29 | 20 g | Itch-Soothe |
| Barrier Creams and Emollients | | | | |
| Barrier Creams | | | | |
| DIMETHICONE Crm 5% tube - 1% DV Oct-19 to 2022 | | .1.53 | 100 g | healthE Dimethicone |
| Crm 5% pump bottle Crm 10% pump bottle – 1% DV Sep-18 to 2021 | | | 500 ml 500 ml | 5% healthE Dimethicone 5% healthE Dimethicone 10% |
| ZINC Crm | | | | e.g. Zinc Cream (Orion-) ;Zinc Cream (PSM) |
| Oint Paste | | | | e.g. Zinc oxide (PSM) |

| | Price | | Brand or |
|---|--------------------|----------|---------------------|
| | (ex man. excl. GST |) | Generic |
| | `\$ | Per | Manufacturer |
| NC AND CASTOR OIL | | | |
| Crm | | 20 g | Orion |
| Oint | | 500 g | Boucher |
| Note: DV limit applies to the pack sizes of greater that 30 g. | | 0 | |
| Oint, BP | 1.26 | 20 g | healthE |
| Note: DV limit applies to the pack sizes of 30 g or less. | | • | |
| NC WITH WOOL FAT | | | |
| Crm zinc 15.25% with wool fat 4% | | | e.g. Sudocrem |
| | | | |
| Emollients | | | |
| QUEOUS CREAM | | | |
| Crm 100 g - 1% DV Oct-18 to 2021 | 1.05 | 100 g | Pharmacy Health |
| | | - | SLS-free |
| Note: DV limit applies to the pack sizes of 100 g or less. | | | |
| Crm 500 g - 1% DV Dec-18 to 2021 | 1.92 | 500 g | Boucher |
| Note: DV limit applies to the pack sizes of greater than 100 g. | | | |
| TOMACROGOL | | | |
| Crm BP, 500 g - 1% DV Sep-18 to 2021 | | 500 g | healthE |
| Crm BP, 100 g - 1% DV Sep-18 to 2021 | 1.42 | 1 | healthE |
| ETOMACROGOL WITH GLYCEROL | | | |
| Crm 90% with glycerol 10%, -1% DV Dec-19 to 2022 | 1.65 | 100 g | healthE |
| Note: DV limit applies to the pack sizes of 100 g or less. | | • | |
| Crm 90% with glycerol 10% - 1% DV Mar-20 to 2022 | 2.35 | 500 ml | ADE |
| | 3.10 | 1,000 ml | ADE |
| | 2.35 | 500 ml | Boucher |
| | 3.10 | 1,000 ml | Boucher |
| Note: DV limit applies to the pack sizes of greater than 100 g. | | | |
| /ULSIFYING OINTMENT | | | |
| Oint BP - 1% DV Oct-20 to 2023 | 1.84 | 100 g | Jaychem |
| Note: DV limit applies to pack sizes of less than 200 g. | | | |
| Oint BP, 500 g | 3.59 | 500 g | AFT |
| Note: DV limit applies to pack sizes of greater than 200 g. | | | |
| YCEROL WITH PARAFFIN | | | |
| Crm glycerol 10% with white soft paraffin 5% and liquid paraffin 10 | % | | e.g. QV cream |
| L IN WATER EMULSION | | | 0 |
| Crm, 500 g – 1% DV Jan-19 to 2021 | 2 19 | 500 g | O/W Fatty Emulsion |
| | | 000 g | Cream |
| Note: DV limit applies to the pack sizes of greater than 100 g. | | | |
| Crm, 100 g - 1% DV Dec-18 to 2021 | 1.44 | 1 | healthE Fatty Cream |
| ARAFFIN | | | |
| Oint liquid paraffin 50% with white soft paraffin 50% - 1% DV Jan- | ·19 | | |
| to 2021 | | 100 g | healthE |
| Note: DV limit applies to the pack sizes of 100 g or greater. | | | · |
| White soft – 1% DV Sep-18 to 2021 | 0.79 | 10 g | healthE |
| Note: DV limit applies to pack sizes of 30 g or less, and to bot | | | |
| White soft, - 1% DV Apr-20 to 2022 | | 450 g | healthE |
| Yellow soft | | - | |

| | Price (ex man. excl. GS \$ | T) Per | Brand or Generic Manufacturer |
|---|----------------------------------|----------------|--|
| PARAFFIN WITH WOOL FAT | | | |
| Lotn liquid paraffin 15.9% with wool fat 0.6% | | | e.g. AlphaKeri;BK ;DP; Hydroderm Lotn |
| Lotn liquid paraffin 91.7% with wool fat 3% | | | e.g. Alpha Keri Bath Oil |
| UREA | 4.07 | 400 | |
| Crm 10% | 1.37 | 100 g | healthE Urea Cream |
| WOOL FAT Crm | | | |
| Corticosteroids | | | |
| | | | |
| BETAMETHASONE DIPROPIONATE Crm 0.05% | | | |
| Oint 0.05% | | | |
| BETAMETHASONE VALERATE | | | |
| Crm 0.1% – 1% DV Oct-18 to 2021 | | 50 g | Beta Cream |
| Oint 0.1% - 1% DV Oct-18 to 2021 | 3.45 | 50 g | Beta Ointment |
| Lotn 0.1% - 1% DV Dec-18 to 2021 | | 50 ml | Betnovate |
| CLOBETASOL PROPIONATE | | | |
| Crm 0.05% – 1% DV Nov-19 to 2022 | | 30 g | Dermol |
| Oint 0.05% – 1% DV Nov-19 to 2022 | 2.12 | 30 g | Dermol |
| CLOBETASONE BUTYRATE Crm 0.05% | | | |
| DIFLUCORTOLONE VALERATE - Restricted: For continuation only | v | | |
| → Crm 0.1% | <i>y</i> | | |
| ➡ Fatty oint 0.1% | | | |
| HYDROCORTISONE | | | |
| Crm 1%, 100 g – 1% DV Sep-20 to 2022 | | 100 g | Hydrocortisone (PSM) |
| Crm 1%, 30 g | | 30 g | DermAssist |
| Note: DV limit applies to the pack sizes of less than or equal Crm 1%, 500 g | | 500 g | Hydrocortisone (PSM) |
| (DermAssist Crm 1%, 30 g to be delisted 1 September 2020) | | 500 g | |
| HYDROCORTISONE ACETATE | | | |
| Crm 1% | 2.48 | 14.2 g | AFT |
| (AFT Crm 1% to be delisted 1 November 2020) | | 0 | |
| HYDROCORTISONE AND PARAFFIN LIQUID AND LANOLIN | | | |
| Lotn 1% with paraffin liquid 15.9% and lanolin 0.6% - 1% DV Oc | t-20 | | |
| to 2023 | 10.57 | 250 ml | DP Lotn HC |
| HYDROCORTISONE BUTYRATE Crm 0.1% | 6.05 | 100 ~ | Logoid Lingaroom |
| Oint 0.1% – 1% DV Mar-19 to 2021 | | 100 g 100 g | Locoid Lipocream Locoid |
| Milky emul 0.1% – 1% DV Mar-19 to 2021 | | 100 g | Locoid Crelo |
| METHYLPREDNISOLONE ACEPONATE | | | |
| Crm 0.1% | 4.95 | 15 g | Advantan |
| Oint 0.1% | 4.95 | 15 g | Advantan |

| | Price (ex man. excl. GST \$ | Per | Brand or Generic Manufacturer |
|---|-----------------------------------|--------------|-------------------------------------|
| MOMETASONE FUROATE | | | |
| Crm 0.1% – 1% DV Nov-18 to 2021 | | 15 g | Elocon Alcohol Free |
| | 2.50 | 50 g | Elocon Alcohol Free |
| Oint 0.1% - 1% DV Nov-18 to 2021 | 1.51 | 15 g | Elocon |
| | 2.90 | 50 g | Elocon |
| Lotn 0.1% - 1% DV Nov-18 to 2021 | 6.30 | 30 ml | Elocon |
| TRIAMCINOLONE ACETONIDE | | | |
| Crm 0.02% – 1% DV Nov-20 to 2023 | 6.30 | 100 g | Aristocort |
| Oint 0.02% - 1% DV Nov-20 to 2023 | 6.35 | 100 g | Aristocort |
| | | - | |
| Corticosteroids with Anti-Infective Agents | | | |
| BETAMETHASONE VALERATE WITH CLIOQUINOL - Restricted ↓ Crm 0.1% with clioquiniol 3% → Restricted (RS1125) Initiation Either: | see terms below | | |
| For the treatment of intertrigo; or For continuation use. | | | |
| BETAMETHASONE VALERATE WITH SODIUM FUSIDATE [FUSID Crm 0.1% with sodium fusidate (fusidic acid) 2% | IC ACID] | | |
| HYDROCORTISONE WITH MICONAZOLE | | | |
| Crm 1% with miconazole nitrate 2% - 1% DV Sep-18 to 2021 | 2.00 | 15 g | Micreme H |
| HYDROCORTISONE WITH NATAMYCIN AND NEOMYCIN | | Ū | |
| Crm 1% with natamycin 1% and neomycin sulphate 0.5% | 3.35 | 15 g | Pimafucort |
| Oint 1% with natamycin 1% and neomycin sulphate 0.5% | | 15 g | Pimafucort |
| TRIAMCINOLONE ACETONIDE WITH NEOMYCIN SULPHATE, GF | | Ũ | |
| Crm 1 mg with nystatin 100,000 u, neomycin sulphate 2.5 mg ar gramicidin 250 mcg per g | | | |
| Psoriasis and Eczema Preparations | | | |
| ACITRETIN | | | |
| Cap 10 mg – 1% DV Oct-20 to 2023 | 17 86 | 60 | Novatretin |
| Cap 25 mg – 1% DV Oct-20 to 2023 | | 60 60 | Novatretin |
| | | 00 | |
| BETAMETHASONE DIPROPIONATE WITH CALCIPOTRIOL Foam spray 500 mcg with calcipotriol 50 mcg per g | 50.05 | 60 a | Enctilor |
| | | 60 g | Enstilar Daivobet |
| Gel 500 mcg with calcipotriol 50 mcg per g – 1% DV Dec-18 to Oint 500 mcg with calcipotriol 50 mcg per g – 1% DV Dec-18 to | | 60 g 30 g | Daivobet |
| | EVET 13.30 | 50 y | Daivobel |
| CALCIPOTRIOL Oint 50 mag par a | 40.00 | 100 ~ | Doivonov |
| Oint 50 mcg per g | 40.00 | 120 g | Daivonex |
| COAL TAR WITH SALICYLIC ACID AND SULPHUR Oint 12% with salicylic acid 2% and sulphur 4% | | | |
| METHOXSALEN [8-METHOXYPSORALEN] Tab 10 mg Lotn 1.2% | | | |
| PINE TAR WITH TROLAMINE LAURILSULFATE AND FLUORESCI Soln 2.3% with trolamine laurilsulfate and fluorescein sodium – | 1% DV | | _ |
| Nov-20 to 2023 | 4.44 | 500 ml | Pinetarsol |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| Crm 5%, 250 mg sachet |
|---|
| POTASSIUM PERMANGANATE Tab 400 mg Crystals Scalp Preparations BETAMETHASONE VALERATE Scalp app 0.1% - 1% DV Oct-18 to 2021 |
| Tab 400 mg Crystals Scalp Preparations BETAMETHASONE VALERATE Scalp app 0.1% - 1% DV Oct-18 to 2021 |
| Scalp Preparations BETAMETHASONE VALERATE Scalp app 0.1% - 1% DV Oct-18 to 2021 |
| BETAMETHASONE VALERATE Scalp app 0.1% - 1% DV Oct-18 to 2021 |
| Scalp app 0.1% - 1% DV Oct-18 to 2021 |
| CLOBETASOL PROPIONATE Scalp app 0.05% - 1% DV Nov-19 to 2022 |
| Scalp app 0.05% - 1% DV Nov-19 to 2022 |
| HYDROCORTISONE BUTYRATE Scalp lotn 0.1% - 1% DV Mar-19 to 2021 |
| Scalp lotn 0.1% - 1% DV Mar-19 to 2021 7.30 100 ml Locoid Wart Preparations IMIQUIMOD 21.72 24 Perrigo PODOPHYLLOTOXIN 21.72 24 Perrigo Soln 0.5% 33.60 3.5 ml Condyline SILVER NITRATE 33.60 3.5 ml Condyline SILVER NITRATE Sticks with applicator DIPHEMANIL METILSULFATE Powder 2% SUNSCREEN, PROPRIETARY SUNSCREEN, PROPRIETARY SUNSCREEN, PROPRIETARY |
| IMIQUIMOD Crm 5%, 250 mg sachet |
| PODOPHYLLOTOXIN Soln 0.5% |
| PODOPHYLLOTOXIN Soln 0.5% |
| Soln 0.5% |
| SILVER NITRATE Sticks with applicator Other Skin Preparations DIPHEMANIL METILSULFATE Powder 2% SUNSCREEN, PROPRIETARY |
| Sticks with applicator Other Skin Preparations DIPHEMANIL METILSULFATE Powder 2% SUNSCREEN, PROPRIETARY |
| DIPHEMANIL METILSULFATE Powder 2% SUNSCREEN, PROPRIETARY |
| Powder 2% SUNSCREEN, PROPRIETARY |
| SUNSCREEN, PROPRIETARY |
| |
| Loto 19/ DV Mor 20 to 2022 |
| Lotn – 1% DV Mar-20 to 20225.10 200 g Marine Blue Lotion 50+ |
| Antineoplastics |
| FLUOROURACIL SODIUM |
| Crm 5% – 1% DV Sep-18 to 2021 |
| METHYL AMINOLEVULINATE HYDROCHLORIDE – Restricted see terms below Crm 16% |
| ➡ Restricted (RS1127) |
| Dermatologist or plastic surgeon |
| Wound Management Products |
| |

Gel 2.5%

e.g. Orion

| | Price | | Brand or |
|---|--------------------------|----------|-------------------------------|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer |
| | Ψ | 1 61 | Manulacturer |
| Anti-Infective Agents | | | |
| ACETIC ACID | | | |
| Soln 3% | | | |
| Soln 5% | | | |
| ACETIC ACID WITH HYDROXYQUINOLINE, GLYCEROL AND RICIN Jelly 0.94% with hydroxyquinoline sulphate 0.025%, glycerol 5% ar ricinoleic acid 0.75% with applicator | | | |
| CHLORHEXIDINE GLUCONATE | | | |
| Crm 1% | | 50 g | healthE |
| Lotn 1%, 200 ml | 2.98 | 1 | healthE |
| (healthE Crm 1% to be delisted 1 November 2020) (healthE Lotn 1%, 200 ml to be delisted 1 November 2020) | | | |
| CLOTRIMAZOLE | | | |
| Vaginal crm 1% with applicator - 1% DV Jan-20 to 2022 | | 35 g | Clomazol |
| Vaginal crm 2% with applicator – 1% DV Jan-20 to 2022 | 3.00 | 20 g | Clomazol |
| VIICONAZOLE NITRATE Vaginal crm 2% with applicator – 1% DV Nov-20 to 2023 | 6 89 | 40 g | Micreme |
| | | τυg | moreme |
| Vaginal crm 100,000 u per 5 g with applicator(s) – 1% DV Oct-20 f | o 2023 4.00 | 75 g | Nilstat |
| Contracontines | | | |
| Contraceptives | | | |
| Antiandrogen Oral Contraceptives | | | |
| CYPROTERONE ACETATE WITH ETHINYLOESTRADIOL | | | |
| Tab 2 mg with ethinyloestradiol 35 mcg and 7 inert tablets | 4.67 | 168 | Ginet |
| Combined Oral Contraceptives | | | |
| ETHINYLOESTRADIOL WITH DESOGESTREL | | | |
| Tab 20 mcg with desogestrel 150 mcg | | | |
| Tab 30 mcg with desogestrel 150 mcg | | | |
| ETHINYLOESTRADIOL WITH LEVONORGESTREL | | | |
| Tab 20 mcg with levonorgestrel 100 mcg and 7 inert tablets Tab 30 mcg with levonorgestrel 150 mcg and 7 inert tablets | | 84 84 | Microgynon 20 ED Levlen ED |
| Tab 20 mcg with levonorgestrel 100 mcg | | 04 | |
| Tab 30 mcg with levonorgestrel 150 mcg | | | |
| Tab 50 mcg with levonorgestrel 125 mcg | 9.45 | 84 | Microgynon 50 ED |
| THINYLOESTRADIOL WITH NORETHISTERONE Tab 35 mcg with norethisterone 1 mg | | | |
| Tab 35 mcg with norethisterone 1 mg and 7 inert tab – 1% DV Mai | -20 | | |
| to 2022 | | 84 | Brevinor 1/28 |
| Tab 35 mcg with norethisterone 500 mcg | | | |
| NORETHISTERONE WITH MESTRANOL | | | |
| Tab 1 mg with mestranol 50 mcg | | | |

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GENITO-URINARY SYSTEM

| Price (ex man. excl. GST | 7 | Brand or Generic |
|---|-----|-----------------------|
| (ox mail: oxo: oxo: \$ | Per | Manufacturer |
| Contraceptive Devices | | |
| INTRA-UTERINE DEVICE | | |
| IUD 29.1 mm length × 23.2 mm width – 1% DV Nov-19 to 2022 | 1 | Choice TT380 Short |
| IUD 33.6 mm length × 29.9 mm width – 1% DV Nov-19 to 2022 | 1 | Choice TT380 Standard |
| IUD 35.5 mm length × 19.6 mm width - 1% DV Nov-19 to 2022 15.50 | 1 | Choice Load 375 |
| Emergency Contraception | | |
| LEVONORGESTREL | | |
| Tab 1.5 mg | 1 | Postinor-1 |
| • | | |
| Progestogen-Only Contraceptives | | |
| LEVONORGESTREL | | |
| Tab 30 mcg - 1% DV May-20 to 2022 | 84 | Microlut |
| Subdermal implant (2 × 75 mg rods) | 1 | Jadelle |
| Intra-uterine device 52 mg - 1% DV Nov-19 to 31 Oct 2022 | 1 | Mirena |
| Intra-uterine device 13.5 mg - 1% DV Nov-19 to 31 Oct 2022 | 1 | Jaydess |
| MEDROXYPROGESTERONE ACETATE | | |
| Inj 150 mg per ml, 1 ml syringe – 1% DV Dec-19 to 2022 | 1 | Depo-Provera |
| NORETHISTERONE | | |
| Tab 350 mcg - 1% DV Sep-18 to 20216.25 | 84 | Noriday 28 |
| | | - |
| Obstetric Preparations | | |
| Antiprogestogens | | |
| MIFEPRISTONE | | |
| Tab 200 mg | | |
| Oxytocics | | |
| CARBOPROST TROMETAMOL | | |
| Inj 250 mcg per ml, 1 ml ampoule | | |
| DINOPROSTONE | | |
| Pessaries 10 mg | | |
| Vaginal gel 1 mg in 3 g | 1 | Prostin E2 |
| Vaginal gel 2 mg in 3 g69.77 | 1 | Prostin E2 |
| ERGOMETRINE MALEATE | | |
| Inj 500 mcg per ml, 1 ml ampoule | 5 | DBL Ergometrine |
| OXYTOCIN | | 3 |
| Inj 5 iu per ml, 1 ml ampoule – 1% DV Nov-18 to 2021 | 5 | Oxytocin BNM |
| Inj 10 iu per ml, 1 ml ampoule – 1% DV Nov-18 to 2021 | 5 | Oxytocin BNM |
| DXYTOCIN WITH ERGOMETRINE MALEATE | • | , |
| Inj 5 iu with ergometrine maleate 500 mcg per ml, 1 ml ampoule – 1% | | |
| DV Oct-18 to 2021 | 5 | Syntometrine |
| Tocolytics | | - |
| PROCESTERONE Destricted and terms on the part page | | |
| | | |
| PROGESTERONE – Restricted see terms on the next page Cap 100 mg16.50 | 30 | Utrogestan |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price | | | Brand or |
|--------|---------|--------|-----|--------------|
| (ex ma | n. excl | . GST) | | Generic |
| | \$ | | Per | Manufacturer |

➡ Restricted (RS1533)

Initiation

Gynaecologist or obstetrician

Re-assessment required after 12 months

Both:

- 1 For the prevention of pre-term labour*; and
- 2 Either:
 - 2.1 The patient has a short cervix on ultrasound (defined as < 25mm at 16 to 28 weeks); or
 - 2.2 The patient has a history of pre-term birth at less than 28 weeks.

Continuation

Gynaecologist or obstetrician

Re-assessment required after 12 months

All of the following:

- 1 For the prevention of pre-term labour*; and
- 2 Treatment is required for second or subsequent pregnancy; and
- 3 Either:
 - 3.1 The patient has a short cervix on ultrasound (defined as < 25mm at 16 to 28 weeks); or
 - 3.2 The patient has a history of pre-term birth at less than 28 weeks.

Note: Indications marked with * are unapproved indications.

TERBUTALINE - Restricted see terms below

Inj 500 mcg ampoule

⇒ Restricted (RS1130)

Obstetrician

Oestrogens

| Crm 1 mg per g with applicator - 1% DV Oct-20 to 2023 | 6.62 | 15 g | Ovestin | |
|---|------|------|---------|--|
| Pessaries 500 mcg - 1% DV Oct-20 to 2023 | 6.86 | 15 | Ovestin | |

| Urologicals | | |
|---|--------------------|----------------|
| 5-Alpha Reductase Inhibitors | | |
| FINASTERIDE - Restricted see terms below ↓ Tab 5 mg | 100 dicated; or | Ricit |
| Alpha-1A Adrenoceptor Blockers | | |
| TAMSULOSIN HYDROCHLORIDE - Restricted see terms below ↓ Cap 400 mcg - 1% DV Jan-20 to 2022 | 100 | Tamsulosin-Rex |

continued...

GENITO-URINARY SYSTEM

| | I | Price | | | Brand or |
|--|---------|---------|----------|--------|-------------------|
| (ex | man. | | GST) | Der | Generic |
| | | \$ | | Per | Manufacturer |
| ontinued | | | | | |
| Patient has symptomatic benign prostatic hyperplasia; and The patient is intolerant of non-selective alpha blockers or these are | e cont | traindi | icated. | | |
| Urinary Alkalisers | | | | | |
| POTASSIUM CITRATE - Restricted see terms below | | | | | |
| Oral liq 3 mmol per ml – 1% DV Oct-18 to 2021 | | .31.8 | 0 | 200 ml | Biomed |
| Restricted (RS1133) | | | | | |
| nitiation | | | | | |
| Both: | | | | | |
| The patient has recurrent calcium oxalate urolithiasis; and The patient has had more than two renal calculi in the two years pri- | or to t | the ap | oplicati | on. | |
| ODIUM CITRO-TARTRATE | | | | | |
| Grans eff 4 g sachets - 1% DV Oct-20 to 2023 | | 2.2 | 2 | 28 | Ural |
| Urinary Antispasmodics | | | | | |
| DXYBUTYNIN | | | | | |
| Tab 5 mg | | .11.7 | 0 | 500 | Apo-Oxybutynin |
| Oral liq 5 mg per 5 ml | | | | 473 ml | Apo-Oxybutynin |
| OLIFENACIN SUCCINATE – Some items restricted see terms below | | | | | |
| Tab 5 mg - 1% DV Dec-18 to 2021 | | 3.0 | 0 | 30 | Solifenacin Mylan |
| Tab 10 mg - 1% DV Dec-18 to 2021 | | 5.5 | 0 | 30 | Solifenacin Mylan |
| → Restricted (RS1274) | | | | | - |
| nitiation | | | | | |

Patient has overactive bladder and a documented intolerance of, or is non-responsive to, oxybutynin.

| (ex | Price man. excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------------------------|------------|-----------|-------------------------------------|
| Anabolic Agents | | | | |
| DXANDROLONE | | | | |
| Tab 2.5 mg | | | | |
| ◆ Restricted (RS1302) nitiation | | | | |
| For the treatment of burns patients. | | | | |
| Androgen Agonists and Antagonists | | | | |
| CYPROTERONE ACETATE | | | | |
| Tab 50 mg - 1% DV Dec-18 to 2021 | | 7 | 50 | Siterone |
| Tab 100 mg - 1% DV Dec-18 to 2021 | | | 50 | Siterone |
| ESTOSTERONE | | | | |
| Patch 5 mg per day | 90.0 | 0 | 30 | Androderm |
| ESTOSTERONE CIPIONATE | 70 5 | • | | Dana Tastastasaa |
| Inj 100 mg per ml, 10 ml vial | | 0 | 1 | Depo-Testosterone |
| ESTOSTERONE ESTERS Inj testosterone decanoate 100 mg, testosterone isocarproate 60 mg, | | | | |
| testosterone phenylpropionate 60 mg and testosterone propionate | | | | |
| 30 mg per ml, 1 ml ampoule | | | | |
| ESTOSTERONE UNDECANOATE | | | | |
| Cap 40 mg – 1% DV Nov-18 to 2021 Inj 250 mg per ml, 4 ml vial | | | 60 1 | Andriol Testocaps Reandron 1000 |
| | | 0 | I | Realition 1000 |
| Calcium Homeostasis | | | | |
| CALCITONIN | | | | |
| Inj 100 iu per ml, 1 ml ampoule | 121.0 | 0 | 5 | Miacalcic |
| CINACALCET – Restricted see terms below | | | | |
| Tab 30 mg - 1% DV Sep-18 to 2021 | 210.3 | 0 | 28 | Sensipar |
| Restricted (RS1540) nitiation | | | | |
| lephrologist or endocrinologist | | | | |
| Re-assessment required after 6 months | | | | |
| ither: | | | | |
| 1 All of the following: | | | | |
| 1.1 The patient has been diagnosed with a parathyroid carcinoma | | <i>'</i> . | | |
| 1.2 The patient has persistent hypercalcaemia (serum calcium gr first-line treatments including sodium thiosulfate (where appro | | | | |
| 1.3 The patient is symptomatic; or | priatoj di | a bibpi | 1000100 | aco, ana |
| 2 All of the following: | | | | |
| 2.1 The patient has been diagnosed with calciphylaxis (calcific ur | aemic arte | eriolopa | athy); ar | nd |
| 2.2 The patient has symptomatic (e.g. painful skin ulcers) hypere | alcaemia | (serun | n calciur | n greater than or equal to |

3 mmol/L); and 2.3 The patient's condition has not responded to previous first-line treatments including bisphosphonates and sodium thiosulfate.

continued...

| Price (ex man. excl. (\$ | | Per | Brand or Generic Manufacturer |
|--|----------|------------|-------------------------------------|
| continued | | | |
| Continuation | | | |
| Nephrologist or endocrinologist | | | |
| Both: | | | |
| 1 The patient's serum calcium level has fallen to < 3mmol/L; and | | | |
| 2 The patient has experienced clinically significant symptom improvement. | | | |
| Note: This does not include parathyroid adenomas unless these have become malignation | ant. | | |
| ZOLEDRONIC ACID | | | |
| Inj 4 mg per 5 ml, vial – 1% DV May-19 to 2021 | | 1 | Zoledronic acid Mylan |
| → Restricted (RS1602) | | | |
| nitiation – bone metastases | | | |
| Dicologist, haematologist or palliative care specialist | | | |
| Any of the following: | | | |
| 1 Patient has hypercalcaemia of malignancy; or | | | |
| 2 Both: | | | |
| 2.1 Patient has bone metastases or involvement; and | | | |
| 2.2 Patient has severe bone pain resistant to standard first-line treatments; c | JL | | |
| 3 Both: | | | |
| 3.1 Patient has bone metastases or involvement; and3.2 Patient is at risk of skeletal-related events (pathological fracture, spinal c | ord co | mpress | ion, radiation to bone or |
| surgery to bone). | | | |
| nitiation – early breast cancer | | | |
| Dncologist | | | |
| All of the following: | | | |
| 1 Treatment to be used as adjuvant therapy for early breast cancer; and | | اممر مالك. | |
| 2 Patient has been amenorrhoeic for 12 months or greater, either naturally or indu a patheonogeneous state and | ucea, w | htn end | ocrine levels consistent wit |
| a postmenopausal state; and 2. Treatment to be administered at a minimum interval of 6 monthly for a maximum | m of 0 v | | |
| 3 Treatment to be administered at a minimum interval of 6-monthly for a maximum | n oi 2 y | lears. | |
| | | | |
| Corticosteroids | | | |
| | | | |
| BETAMETHASONE | | | |
| Tab 500 mcg | | | |
| Inj 4 mg per ml, 1 ml ampoule | | | |
| BETAMETHASONE SODIUM PHOSPHATE WITH BETAMETHASONE ACETATE | | | |
| Inj 3.9 mg with betamethasone acetate 3 mg per ml, 1 ml ampoule | | | |
| EXAMETHASONE | | | |
| Tab 0.5 mg - 1% DV Oct-18 to 2021 | | 30 | Dexmethsone |
| Tab 4 mg - 1% DV Oct-18 to 2021 | | 30 | Dexmethsone |
| Oral liq 1 mg per ml | | 25 ml | Biomed |
| DEXAMETHASONE PHOSPHATE | | | |
| Inj 4 mg per ml, 1 ml ampoule – 1% DV Jul-20 to 2022 | | 10 | Dexamethasone |
| , Jessing and provide the second s | | | Phosphate |

| Inj 4 mg per ml, 2 ml ampoule – 1% DV Jul-20 to 2022 | 6.37 | 10 | Panpharma Dexamethasone Phosphate Panpharma |
|---|------|-----|--|
| FLUDROCORTISONE ACETATE Tab 100 mcg14 | 4.32 | 100 | Florinef |

Products with Hospital Supply Status (HSS) are in **bold**

Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GST | 7) | Brand or Generic |
|---|-----------------------------|-------|-----------------------|
| | (ex man. exci. GST \$ | Per | Manufacturer |
| HYDROCORTISONE | | | |
| Tab 5 mg - 1% DV Sep-18 to 2021 | 8.10 | 100 | Douglas |
| Tab 20 mg - 1% DV Sep-18 to 2021 | 20.32 | 100 | Douglas |
| Inj 100 mg vial | | 1 | Solu-Cortef |
| METHYLPREDNISOLONE (AS SODIUM SUCCINATE) | | | |
| Tab 4 mg - 1% DV Dec-18 to 2021 | 112.00 | 100 | Medrol |
| Tab 100 mg - 1% DV Dec-18 to 2021 | | 20 | Medrol |
| Inj 40 mg vial – 1% DV Dec-18 to 2021 | | 1 | Solu-Medrol Act-O-Via |
| Inj 125 mg vial - 1% DV Dec-18 to 2021 | | 1 | Solu-Medrol Act-O-Via |
| Inj 500 mg vial - 1% DV Dec-18 to 2021 | | 1 | Solu-Medrol Act-O-Via |
| Inj 1 g vial - 1% DV Dec-18 to 2021 | | 1 | Solu-Medrol |
| METHYLPREDNISOLONE ACETATE | | | |
| Inj 40 mg per ml, 1 ml vial - 1% DV Dec-18 to 2021 | | 5 | Depo-Medrol |
| PREDNISOLONE | | | • |
| Oral liq 5 mg per ml – 1% DV Jun-18 to 2021 | 6.00 | 30 ml | Redipred |
| Enema 200 mcg per ml, 100 ml | | ••• | |
| PREDNISONE | | | |
| Tab 1 mg | | 500 | Apo-Prednisone |
| Tab 2.5 mg | | 500 | Apo-Prednisone |
| Tab 5 mg | | 500 | Apo-Prednisone |
| Tab 20 mg | | 500 | Apo-Prednisone |
| TRIAMCINOLONE ACETONIDE | | | |
| Inj 10 mg per ml, 1 ml ampoule – 5% DV Nov-20 to 2023 | 20.80 | 5 | Kenacort-A 10 |
| Inj 40 mg per ml, 1 ml ampoule – 1% DV Nov-20 to 2023 | | 5 | Kenacort-A 40 |
| TRIAMCINOLONE HEXACETONIDE | | | |

Inj 20 mg per ml, 1 ml vial

Hormone Replacement Therapy

Oestrogens

OESTRADIOL

| Tab 1 mg | | | |
|---------------------------------|-------|----|-----------|
| Patch 25 mcg per day | 6.12 | 8 | Estradot |
| Patch 50 mcg per day | | 8 | Estradot |
| Patch 75 mcg per day | 7.91 | 8 | Estradot |
| Patch 100 mcg per day | 7.91 | 8 | Estradot |
| OESTRADIOL VALERATE | | | |
| Tab 1 mg - 1% DV Sep-18 to 2021 | | 84 | Progynova |
| Tab 2 mg - 1% DV Sep-18 to 2021 | 12.36 | 84 | Progynova |
| OESTROGENS (CONJUGATED EQUINE) | | | |

Tab 300 mcg

Tab 625 mcg

Progestogen and Oestrogen Combined Preparations

OESTRADIOL WITH NORETHISTERONE ACETATE

Tab 1 mg with 0.5 mg norethisterone acetate

Tab 2 mg with 1 mg norethisterone acetate

Tab 2 mg with 1 mg norethisterone acetate (10), and tab 2 mg oestradiol (12) and tab 1 mg oestradiol (6)

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|-----------------|-------------------------------------|
| OESTROGENS WITH MEDROXYPROGESTERONE ACETATE Tab 625 mcg conjugated equine with 2.5 mg medroxyprogesterone acetate Tab 625 mcg conjugated equine with 5 mg medroxyprogesterone acetate | | | |
| Progestogens | | | |
| MEDROXYPROGESTERONE ACETATE Tab 2.5 mg Tab 5 mg Tab 10 mg Other Endocrine Agents | 14.00 | 30 100 30 | Provera Provera Provera |
| CABERGOLINE – Restricted see terms below Tab 0.5 mg – 1% DV Sep-18 to 2021 | | 2 | Dostinex Dostinex |
| → Restricted (RS1319) Initiation Any of the following: Inhibition of lactation; or Patient has pathological hyperprolactinemia; or Patient has acromegaly. | 15.20 | U | POSITIEX |
| CLOMIFENE CITRATE Tab 50 mg | | 10 | Mylan Clomiphen |
| DANAZOL Cap 100 mg Cap 200 mg | | 28 100 | Mylan Azol |
| GESTRINONE Cap 2.5 mg METYRAPONE | | | |
| Cap 250 mg PENTAGASTRIN Inj 250 mcg per ml, 2 ml ampoule | | | |
| Other Oestrogen Preparations | | | |
| ETHINYLOESTRADIOL Tab 10 mcg – 1% DV Sep-18 to 2021 | 17.60 | 100 | NZ Medical and Scientific |
| OESTRADIOL Implant 50 mg OESTRIOL | | | Colonino |
| Tab 2 mg – 1% DV Sep-20 to 2023 | 7.00 | 30 | Ovestin |
| Other Progestogen Preparations | | | |
| MEDROXYPROGESTERONE Tab 100 mg | 101.00 | 100 | Provera HD |
| NORETHISTERONE Tab 5 mg – 1% DV Dec-19 to 2021 | | 100 | Primolut N |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| (e | F x man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|-------------|----------------------|--------|------------------|--|
| Pituitary and Hypothalamic Hormones and Analogues | | | | | |
| CORTICOTRORELIN (OVINE) Inj 100 mcg vial THYROTROPIN ALFA Inj 900 mcg vial | | | | | |
| Adrenocorticotropic Hormones | | | | | |
| TETRACOSACTIDE [TETRACOSACTRIN] Inj 250 mcg per ml, 1 ml ampoule Inj 1 mg per ml, 1 ml ampoule | | | | 1 1 | Synacthen Synacthen Depot |
| GnRH Agonists and Antagonists | | | | | |
| BUSERELIN Inj 1 mg per ml, 5.5 ml vial GONADORELIN Inj 100 mcg vial GOSERELIN Implant 3.6 mg, syringe Implant 10.8 mg, syringe LEUPRORELIN ACETATE Inj 3.75 mg prefilled dual chamber syringe Inj 11.25 mg prefilled dual chamber syringe | 1 2 | 221.6 | о О | 1 1 1 1 | Zoladex Zoladex Lucrin Depot 1-month Lucrin Depot 3-month |
| Gonadotrophins | | | | | |
| CHORIOGONADOTROPIN ALFA Inj 250 mcg in 0.5 ml syringe | | | | | |
| Growth Hormone | | | | | |
| SOMATROPIN - Restricted see terms below Inj 5 mg cartridge - 1% DV Oct-18 to 2021 Inj 10 mg cartridge - 1% DV Oct-18 to 2021 Inj 15 mg cartridge - 1% DV Oct-18 to 2021 → Restricted (RS1549) Initiation - growth hormone deficiency in children Endocrinologist or paediatric endocrinologist <i>Re-assessment required after 12 months</i> Either: | | 69.7 | 5 | 1 1 1 | Omnitrope Omnitrope Omnitrope |
| Growth hormone deficiency causing symptomatic hypoglycaemia, sequelae (e.g. cardiomyopathy, hepatic dysfunction) and diagnose | ed with | GH | < 5 mc | | |

samples in the first 2 weeks of life, or from samples during established hypoglycaemia (whole blood glucose < 2 mmol/l using a laboratory device); or

2 All of the following:

2.1 Height velocity < 25th percentile for age; and adjusted for bone age/pubertal status if appropriate over 6 or 12 months using the standards of Tanner and Davies (1985); and

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | - | Generic |
| \$ | Per | Manufacturer |

- 2.2 A current bone age is < 14 years (female patients) or < 16 years (male patients); and
- 2.3 Peak growth hormone value of < 5.0 mcg per litre in response to two different growth hormone stimulation tests. In children who are 5 years or older, GH testing with sex steroid priming is required; and</p>
- 2.4 If the patient has been treated for a malignancy, they should be disease free for at least one year based upon follow-up laboratory and radiological imaging appropriate for the malignancy, unless there are strong medical reasons why this is either not necessary or appropriate; and
- 2.5 Appropriate imaging of the pituitary gland has been obtained.

Continuation – growth hormone deficiency in children

Endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 A current bone age is 14 years or under (female patients) or 16 years or under (male patients); and
- 2 Height velocity is greater than or equal to 25th percentile for age (adjusted for bone age/pubertal status if appropriate) while on growth hormone treatment, as calculated over six months using the standards of Tanner and Davis (1985); and
- 3 Height velocity is greater than or equal to 2.0 cm per year, as calculated over 6 months; and
- 4 No serious adverse effect that the patients specialist considers is likely to be attributable to growth hormone treatment has occurred; and
- 5 No malignancy has developed since starting growth hormone.

Initiation – Turner syndrome

Endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 The patient has a post-natal genotype confirming Turner Syndrome; and
- 2 Height velocity is < 25th percentile over 6-12 months using the standards of Tanner and Davies (1985); and
- 3 A current bone age is < 14 years.

Continuation – Turner syndrome

Endocrinologist or paediatric endocrinologist *Re-assessment required after 12 months* All of the followino:

- 1 Height velocity greater than or equal to 50th percentile for age (while on growth hormone calculated over 6 to 12 months using the Ranke's Turner Syndrome growth velocity charts); and
- 2 Height velocity is greater than or equal to 2 cm per year, calculated over six months; and
- 3 A current bone age is 14 years or under; and
- 4 No serious adverse effect that the specialist considers is likely to be attributable to growth hormone treatment has occurred; and
- 5 No malignancy has developed since starting growth hormone.

Initiation - short stature without growth hormone deficiency

Endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 The patient's height is more than 3 standard deviations below the mean for age or for bone age if there is marked growth acceleration or delay; and
- 2 Height velocity is < 25th percentile for age (adjusted for bone age/pubertal status if appropriate), as calculated over 6 to 12 months using the standards of Tanner and Davies(1985); and
- 3 A current bone age is < 14 years (female patients) or < 16 years (male patients); and
- 4 The patient does not have severe chronic disease (including malignancy or recognized severe skeletal dysplasia) and is not receiving medications known to impair height velocity.

| Price | | Brand or |
|------------------|------|--------------|
| (ex man. excl.) | GST) | Generic |
| \$ | Per | Manufacturer |

Continuation - short stature without growth hormone deficiency

Endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 Height velocity is greater than or equal to 50th percentile (adjusted for bone age/pubertal status if appropriate) as calculated over 6 to 12 months using the standards of Tanner and Davies (1985); and
- 2 Height velocity is greater than or equal to 2 cm per year as calculated over six months; and
- 3 Current bone age is 14 years or under (female patients) or 16 years or under (male patients); and
- 4 No serious adverse effect that the patient's specialist considers is likely to be attributable to growth hormone treatment has occurred.

Initiation - short stature due to chronic renal insufficiency

Endocrinologist, paediatric endocrinologist or renal physician on the recommendation of a endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 The patient's height is more than 2 standard deviations below the mean; and
- 2 Height velocity is < 25th percentile (adjusted for bone age/pubertal status if appropriate) as calculated over 6 to 12 months using the standards of Tanner and Davies (1985); and
- 3 A current bone age is to 14 years or under (female patients) or to 16 years or under (male patients); and
- 4 The patient is metabolically stable, has no evidence of metabolic bone disease and absence of any other severe chronic disease; and
- 5 The patient is under the supervision of a specialist with expertise in renal medicine; and
- 6 Either:
 - 6.1 The patient has a GFR less than or equal to 30 ml/min/1.73 m² as measured by the Schwartz method (Height(cm)/plasma creatinine (umol/l × 40 = corrected GFR (ml/min/1.73 m²) in a child who may or may not be receiving dialysis; or
 - 6.2 The patient has received a renal transplant and has received < 5mg/ m² /day of prednisone or equivalent for at least 6 months.

Continuation - short stature due to chronic renal insufficiency

Endocrinologist, paediatric endocrinologist or renal physician on the recommendation of a endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 Height velocity is greater than or equal to 50th percentile (adjusted for bone age/pubertal status if appropriate) as calculated over 6 to 12 months using the standards of Tanner and Davies (1985); and
- 2 Height velocity is greater than or equal to 2 cm per year as calculated over six months; and
- 3 A current bone age is 14 years or under (female patients) or 16 years or under (male patients); and
- 4 No serious adverse effect that the patients specialist considers is likely to be attributable to growth hormone has occurred; and
- 5 No malignancy has developed after growth hormone therapy was commenced; and
- 6 The patient has not experienced significant biochemical or metabolic deterioration confirmed by diagnostic results; and
- 7 The patient has not received renal transplantation since starting growth hormone treatment; and
- 8 If the patient requires transplantation, growth hormone prescription should cease before transplantation and a new application should be made after transplantation based on the above criteria.

Initiation - Prader-Willi syndrome

Endocrinologist or paediatric endocrinologist *Re-assessment required after 12 months* All of the following:

| Price | | Brand or | |
|---------------------|-----|--------------|--|
| (ex man. excl. GST) | | Generic | |
| \$ | Per | Manufacturer | |

- 1 The patient has a diagnosis of Prader-Willi syndrome that has been confirmed by genetic testing or clinical scoring criteria; and
- 2 The patient is aged six months or older; and
- 3 A current bone age is < 14 years (female patients) or < 16 years (male patients); and
- 4 Sleep studies or overnight eximetry have been performed and there is no obstructive sleep disorder requiring treatment, or if an obstructive sleep disorder is found, it has been adequately treated under the care of a paediatric respiratory physician and/or ENT surgeon; and
- 5 Either:
 - 5.1 Both:
 - 5.1.1 The patient is aged two years or older; and
 - 5.1.2 There is no evidence of type II diabetes or uncontrolled obesity defined by BMI that has increased by greater than or equal to 0.5 standard deviations in the preceding 12 months; or
 - 5.2 The patient is aged between six months and two years and a thorough upper airway assessment is planned to be undertaken prior to treatment commencement and at six to 12 weeks following treatment initiation.

Continuation – Prader-Willi syndrome

Endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 Height velocity is greater than or equal to 50th percentile (adjusted for bone age/pubertal status if appropriate) as calculated over 6 to 12 months using the standards of Tanner and Davies (1985); and
- 2 Height velocity is greater than or equal to 2 cm per year as calculated over six months; and
- 3 A current bone age is 14 years or under (female patients) or 16 years or under (male patients); and
- 4 No serious adverse effect that the patient's specialist con siders is likely to be attributable to growth hormone treatment has occurred; and
- 5 No malignancy has developed after growth hormone therapy was commenced; and
- 6 The patient has not developed type II diabetes or uncontrolled obesity as defined by BMI that has increased by greater than or equal to 0.5 standard deviations in the preceding 12 months.

Initiation - adults and adolescents

Endocrinologist or paediatric endocrinologist

Re-assessment required after 12 months

All of the following:

- 1 The patient has a medical condition that is known to cause growth hormone deficiency (e.g. surgical removal of the pituitary for treatment of a pituitary tumour); and
- 2 The patient has undergone appropriate treatment of other hormonal deficiencies and psychological illnesses; and
- 3 The patient has severe growth hormone deficiency (see notes); and
- 4 The patient's serum IGF-I is more than 1 standard deviation below the mean for age and sex; and
- 5 The patient has poor quality of life, as defined by a score of 16 or more using the disease-specific quality of life questionnaire for adult growth hormone deficiency (QoL-AGHDA®).

Notes: For the purposes of adults and adolescents, severe growth hormone deficiency is defined as a peak serum growth hormone level of less than or equal to 3 mcg per litre during an adequately performed insulin tolerance test (ITT) or glucagon stimulation test.

Patients with one or more additional anterior pituitary hormone deficiencies and a known structural pituitary lesion only require one test. Patients with isolated growth hormone deficiency require two growth hormone stimulation tests, of which, one should be ITT unless otherwise contraindicated. Where an additional test is required, an arginine provocation test can be used with a peak serum growth hormone level of less than or equal to 0.4 mcg per litre.

The dose of somatropin should be started at 0.2 mg daily and be titrated by 0.1 mg monthly until it is within 1 standard deviation of the mean normal value for age and sex; and

The dose of somatropin not to exceed 0.7 mg per day for male patients, or 1 mg per day for female patients.

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

70

At the commencement of treatment for hypopituitarism, patients must be monitored for any required adjustment in replacement doses of corticosteroid and levothyroxine.

Continuation - adults and adolescents

Endocrinologist or paediatric endocrinologist *Re-assessment required after 12 months* Either:

1 All of the following:

- 1.1 The patient has been treated with somatropin for < 12 months; and
- 1.2 There has been an improvement in the Quality of Life Assessment defined as a reduction of at least 8 points on the Quality of Life Assessment of Growth Hormone Deficiency in Adults (QoL-AGHDA®) score from baseline; and
- 1.3 Serum IGF-I levels have increased to within ±1SD of the mean of the normal range for age and sex; and
- 1.4 The dose of somatropin does not exceed 0.7 mg per day for male patients, or 1 mg per day for female patients; or
- 2 All of the following:
 - 2.1 The patient has been treated with somatropin for more than 12 months; and
 - 2.2 The patient has not had a deterioration in Quality of Life defined as a 6 point or greater increase from their lowest QoL-AGHDA® score on treatment (other than due to obvious external factors such as external stressors); and
 - 2.3 Serum IGF-I levels have continued to be maintained within ±1SD of the mean of the normal range for age and sex (other than for obvious external factors); and
 - 2.4 The dose of somatropin has not exceeded 0.7 mg per day for male patients or 1 mg per day for female patients.

Thyroid and Antithyroid Preparations

CARBIMAZOI F Tab 5 mg IODINE Soln BP 50 mg per ml LEVOTHYROXINE Tab 25 mcg Tab 50 mcg Tab 100 mcg LIOTHYRONINE SODIUM Tab 20 mcg → Restricted (RS1301) Initiation For a maximum of 14 days' treatment in patients with thyroid cancer who are due to receive radioiodine therapy. Ini 20 mcg vial POTASSIUM IODATE Tab 170 mg POTASSIUM PERCHLORATE Cap 200 mg PROPYLTHIOURACIL - Restricted see terms below 100 PTU → Restricted (RS1276) Initiation Both: 1 The patient has hyperthyroidism: and 2 The patient is intolerant of carbimazole or carbimazole is contraindicated.

Note: Propylthiouracil is not recommended for patients under the age of 18 years unless the patient is pregnant and other treatments are contraindicated.

e.g. Brand indicates brand example only. It is not a contracted product.

| Price | | Brand or |
|--|-------------|------------------------------|
| (ex man. excl. G \$ | GST) Per | Generic Manufacturer |
| PROTIRELIN | | |
| Inj 100 mcg per ml, 2 ml ampoule | | |
| Vasopressin Agents | | |
| ARGIPRESSIN [VASOPRESSIN] Inj 20 u per ml, 1 ml ampoule | | |
| DESMOPRESSIN ACETATE – Some items restricted see terms below | | |
| Tab 100 mcg | 30 | Minirin |
| Tab 200 mcg | 30 6 ml | Minirin Dosmonrossin-DH&T |
| Inj 4 mcg per ml, 1 ml ampoule | 0 111 | Desmopressin-PH&T |
| Inj 15 mcg per ml, 1 ml ampoule | | |
| Nasal drops 100 mcg per ml | | |
| → Restricted (RS1339) | | |
| Initiation – Nocturnal enuresis | | |
| Either: | | |
| 1 The nasal forms of desmopressin are contraindicated; or | | |
| 2 An enuresis alarm is contraindicated. | | |
| Note: Cranial diabetes insipidus and the nasal forms of desmopressin are contraindical | ted. | |
| TERLIPRESSIN | | |
| lnj 0.1 mg per ml, 8.5 ml ampoule | 5 | Glypressin |
| Inj 1 mg per 8.5 ml ampoule | 5 | Glypressin |



| | Price (ex man. excl. GS \$ | T) Per | Brand or Generic Manufacturer |
|---|----------------------------------|-----------|-------------------------------------|
| Antibacterials | | | |
| Aminoglycosides | | | |
| AMIKACIN - Restricted see terms below Inj 5 mg per ml, 10 ml syringe | | | |
| Inj 5 mg per ml, 5 ml syringe | | 1 | Biomed |
| Inj 15 mg per ml, 5 ml syringe Inj 250 mg per ml, 2 ml vial – 1% DV Aug-18 to 2021 | | 5 | DBL Amikacin |
| ➡ Restricted (RS1041) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory special | ist | | |
| GENTAMICIN SULPHATE Inj 10 mg per ml, 1 ml ampoule | 25.00 | 5 | DBL Gentamicin |
| Inj 40 mg per ml, 2 ml ampoule | | 5 10 | Pfizer |
| PAROMOMYCIN – Restricted see terms below | | 10 | |
| Cap 250 mg | 126.00 | 16 | Humatin |
| ➡ Restricted (RS1603) | | | |
| Clinical microbiologist, infectious disease specialist or gastroenterologist | | | |
| STREPTOMYCIN SULPHATE – Restricted see terms below | | | |
| Inj 400 mg per ml, 2.5 ml ampoule | | | |
| → Restricted (RS1043) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory special | ist | | |
| TOBRAMYCIN Powder | | | |
| Powder → Restricted (RS1475) | | | |
| Initiation | | | |
| For addition to orthopaedic bone cement. | | | |
| Inj 40 mg per ml, 2 ml vial − 1% DV Sep-18 to 2021 → Restricted (R\$1044) | 15.00 | 5 | Tobramycin Mylan |
| Clinical microbiologist, infectious disease specialist or respiratory special | ist | | |
| Inj 100 mg per ml, 5 ml vial → Restricted (R\$1044) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory special | ist | | |
| Solution for inhalation 60 mg per ml, 5 ml | | 56 dose | ТОВІ |
| → Restricted (RS1435) | | 00 0000 | 1001 |
| Initiation | | | |
| Patient has cystic fibrosis. | | | |
| Carbapenems | | | |
| ERTAPENEM – Restricted see terms below ↓ Inj 1 g vial – 1% DV Aug-19 to 2022 → Restricted (RS1045) | 70.00 | 1 | Invanz |
| Clinical microbiologist or infectious disease specialist | | | |
| IMIPENEM WITH CILASTATIN – Restricted see terms below | ~~ ~~ | | Iminonom Olloctotic |
| Inj 500 mg with 500 mg cilastatin vial − 1% DV Jul-19 to 2022 | | 1 | Imipenem+Cilastatin RBX |
| → Restricted (RS1046) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| | | | |

t Item restricted (see \Rightarrow above); **t** Item restricted (see \Rightarrow below) *e.g. Brand* indicates brand example only. It is not a contracted product.

| | Price | - | Brand or |
|---|--------------------------|----------|-----------------------------|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer |
| MEROPENEM – Restricted see terms below | | | |
| Inj 500 mg vial | 4 00 | 1 | Meropenem Ranbaxy |
| Inj 1 q vial | | 1 | Meropenem Ranbaxy |
| → Restricted (RS1047) | | | moroponom nanoaxy |
| Clinical microbiologist or infectious disease specialist | | | |
| Cephalosporins and Cephamycins - 1st Generation | | | |
| | | | |
| CEFALEXIN | | | |
| Cap 250 mg – 1% DV Nov-19 to 2022 | | 20 | Cephalexin ABM |
| Cap 500 mg | | 20 | Cephalexin ABM |
| Grans for oral liq 25 mg per ml - 1% DV Oct-18 to 2021 | | 100 ml | Cefalexin Sandoz |
| Grans for oral liq 50 mg per ml – 1% DV Oct-18 to 2021 | 11./5 | 100 ml | Cefalexin Sandoz |
| CEFAZOLIN | | | |
| Inj 500 mg vial – 1% DV Nov-20 to 2023 | | 5 | AFT |
| Inj 1 g vial – 1% DV Nov-20 to 2023 | 3.49 | 5 | AFT |
| Cephalosporins and Cephamycins - 2nd Generation | | | |
| CEFACLOR | | | |
| Cap 250 mg – 1% DV Oct-19 to 2022 | | 100 | Ranbaxy-Cefaclor |
| Grans for oral liq 25 mg per ml – 1% DV Oct-19 to 2022 | | 100 ml | Ranbaxy-Cefaclor |
| CEFOXITIN | | | • |
| Inj 1 g vial | 58.00 | 10 | Cefoxitin Actavis |
| | | 10 | Ocioxian Addavio |
| CEFUROXIME Tab 250 mg - 1% DV Feb-20 to 2022 | 45.00 | 50 | Zinnat |
| Inj 750 mg vial | | 50 10 | Zinna Cefuroxime Actavis |
| Inj 750 mg viai | | 10 | Cefuroxime Actavis |
| | | | |
| Cephalosporins and Cephamycins - 3rd Generation | | | |
| CEFOTAXIME | | | |
| Inj 500 mg vial | | 1 | Cefotaxime Sandoz |
| Inj 1 g vial – 1% DV Nov-20 to 2023 | 45.00 | 10 | DBL Cefotaxime |
| CEFTAZIDIME – Restricted see terms below | | | |
| Inj 1 g vial – 1% DV Dec-20 to 2023 | | 5 | Ceftazidime Mylan |
| | 2.69 | 1 | Ceftazidime-AFT |
| (Ceftazidime Mylan Inj 1 g vial to be delisted 1 December 2020) | | | |
| ➡ Restricted (RS1048) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory special | list | | |
| CEFTRIAXONE | | | |
| Inj 500 mg vial – 1% DV Jan-20 to 2022 | | 1 | Ceftriaxone-AFT |
| Inj 1 g vial - 1% DV Jan-20 to 2022 | | 5 | Ceftriaxone-AFT |
| Inj 2 g vial – 1% DV Jan-20 to 2022 | 1.98 | 1 | Ceftriaxone-AFT |
| Cephalosporins and Cephamycins - 4th Generation | | | |
| CEFEPIME – Restricted see terms below | | | |
| Inj 1 g vial − 1% DV Sep-18 to 2021 | | 1 | Cefepime-AFT |
| Inj 2 g vial − 1% DV Sep-18 to 2021 | | 1 | Cefepime-AFT |
| → Restricted (RS1049) | | · | |
| Clinical microbiologist or infectious disease specialist | | | |
| • | | | |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated. INFECTIONS

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|-------------|--------------------------------------|
| Cephalosporins and Cephamycins - 5th Generat | ion | | |
| EFTAROLINE FOSAMIL – Restricted see terms below Inj 600 mg vial * Restricted (RS1446) hitiation – multi-resistant organisn salvage therapy linical microbiologist or infectious disease specialist iither: 1 for patients where alternative therapies have failed; or 2 for patients who have a contraindication or hypersensitivity | | 10 pies. | Zinforo |
| Macrolides | | | |
| ZITHROMYCIN – Restricted see terms below Tab 250 mg – 1% DV Sep-18 to 2021 Tab 500 mg – 1% DV Sep-18 to 2021 Grans for oral liq 200 mg per 5 ml (40 mg per ml) – 1% DV DV | 0.93 ec-18 | 30 2 | Apo-Azithromycin Apo-Azithromycin |
| to 2021 → Restricted (RS1598) itiation – bronchiolitis obliterans syndrome, cystic fibrosis a | | 15 ml | Zithromax |
| ny of the following: | | | |
| Patient has received a lung transplant, stem cell transplant bronchiolitis obliterans syndrome*; or Patient has received a lung transplant and requires prophy Patient has cystic fibrosis and has chronic infection with Ps negative organisms*; or Patient has an atypical Mycobacterium infection. | laxis for bronchiolitis oblite | erans sync | drome*; or |
| ote: Indications marked with * are unapproved indications itiation – non-cystic fibrosis bronchiectasis * espiratory specialist or paediatrician <i>le-assessment required after 12 months</i> Il of the following: | | | |
| For prophylaxis of exacerbations of non-cystic fibrosis bron Patient is aged 18 and under; and Either: | nchiectasis*; and | | |
| 3.1 Patient has had 3 or more exacerbations of their browner.3.2 Patient has had 3 acute admissions to hospital for the 12 month period. | | | |
| lote: Indications marked with * are unapproved indications. A m brosis will be subsidised in the community. continuation – non-cystic fibrosis bronchiectasis * lespiratory specialist or paediatrician <i>be accessment required after 12 months</i> . | aximum of 24 months of a | zithromyc | in treatment for non-cysti |

Re-assessment required after 12 months

All of the following:

- 1 The patient has completed 12 months of azithromycin treatment for non-cystic fibrosis bronchiectasis; and
- 2 Following initial 12 months of treatment, the patient has not received any further azithromycin treatment for non-cystic
- fibrosis bronchiectasis for a further 12 months, unless considered clinically inappropriate to stop treatment; and
- 3 The patient will not receive more than a total of 24 months' azithromycin cumulative treatment (see note).

| | | | | INFECTIONS |
|--|-----------------|--------------------------|---------------|--|
| | | Price excl. GST \$ |) Per | Brand or Generic Manufacturer |
| continued Note: Indications marked with * are unapproved indications. A m ibrosis will be subsidised in the community. | aximum of 24 | months of | azithromyo | cin treatment for non-cyst |
| nitiation – other indications | | | | |
| Re-assessment required after 5 days | | | | |
| or any other condition. | | | | |
| Continuation – other indications | | | | |
| Re-assessment required after 5 days | | | | |
| or any other condition. | | | | |
| CLARITHROMYCIN - Restricted see terms below Tab 250 mg | | 2 00 | 14 | Ana Clarithromusin |
| Tab 200 mg | | | 14 | Apo-Clarithromycin Apo-Clarithromycin |
| Grans for oral liq 50 mg per ml | | | 50 ml | Klacid |
| Inj 500 mg vial – 1% DV Dec-17 to 31 Aug 2020 | | | 1 | Martindale |
| Restricted (RS1709) | | | | |
| nitiation – Tab 250 mg and oral liquid | | | | |
| ny of the following: | | | | |
| 1 Atypical mycobacterial infection; or | | | | |
| 2 Mycobacterium tuberculosis infection where there is drug r | esistance or ir | ntolerance | to standard | d pharmaceutical agents; |
| 3 Helicobacter pylori eradication; or | | | | |
| 4 Prophylaxis of infective endocarditis associated with surgic | al or dental pr | ocedures i | f amoxicilli | n is contra-indicated. |
| nitiation – Tab 500 mg | | | | |
| lelicobacter pylori eradication. | | | | |
| nitiation – Infusion | | | | |
| In the following: | | | | |
| Atypical mycobacterial infection; or Mycobacterium tuberculosis infection where there is drug r | ocietanoo or ir | toloranco | to standar | d pharmacoutical agapte: |
| 3 Community-acquired pneumonia. | | liuleranice | io stanuari | a phannaceutical agents, |
| , , , | | | | |
| RYTHROMYCIN (AS ETHYLSUCCINATE) | | 10.05 | 100 | |
| Tab 400 mg Grans for oral liq 200 mg per 5 ml | | | 100 100 ml | E-Mycin E-Mycin |
| Grans for oral lig 400 mg per 5 ml. | | | 100 ml | E-Mycin |
| | | 0.77 | 100 111 | |
| RYTHROMYCIN (AS LACTOBIONATE) | | 10.00 | 1 | Eruthropin IV |
| Inj 1 g vial – 1% DV Dec-19 to 2022 | | . 10.00 | I | Erythrocin IV |
| RYTHROMYCIN (AS STEARATE) – Restricted: For continuat | ion only | | | |
| Tab 250 mg | | | | |
| Tab 500 mg | | | | |
| ROXITHROMYCIN – Some items restricted see terms below | | | | |
| Tab dispersible 50 mg | | | 10 | Rulide D |
| Tab 150 mg - 1% DV Sep-19 to 2022 | | | 50 50 | Arrow-Roxithromyci |
| Tab 300 mg − 1% DV Sep-19 to 2022 | | . 10.33 | 50 | Arrow-Roxithromyci |
| nitiation | | | | |
| Initiation | | | | |

Only for use in patients under 12 years of age.

| | Price (ex man. excl. GS \$ | T) Per | Brand or Generic Manufacturer |
|---|----------------------------------|-----------|-------------------------------------|
| Penicillins | | | |
| AMOXICILLIN | | | |
| Cap 250 mg - 1% DV Apr-20 to 2022 | | 500 | Alphamox |
| Cap 500 mg - 1% DV Apr-20 to 2022 | | 500 | Alphamox |
| Grans for oral liq 125 mg per 5 ml - 1% DV Nov-20 to 2023 | 1.40 | 100 ml | Alphamox 125 |
| Grans for oral liq 250 mg per 5 ml - 1% DV Nov-20 to 2023 | 1.73 | 100 ml | Alphamox 250 |
| Inj 250 mg vial | 10.67 | 10 | Ibiamox |
| Inj 500 mg vial | 12.41 | 10 | Ibiamox |
| Inj 1 g vial | 17.29 | 10 | Ibiamox |
| AMOXICILLIN WITH CLAVULANIC ACID | | | |
| Tab 500 mg with clavulanic acid 125 mg | | 20 | Augmentin |
| Grans for oral lig 25 mg with clavulanic acid 6.25 mg per ml | | 100 ml | Augmentin |
| Grans for oral lig 50 mg with clavulanic acid 12.5 mg per ml | | 100 ml | Curam |
| Inj 500 mg with clavulanic acid 100 mg vial | | 10 | m-Amoxiclav |
| Inj 1,000 mg with clavulanic acid 200 mg vial | | 10 | m-Amoxiclav |
| BENZATHINE BENZYLPENICILLIN | | | |
| Inj 900 mg (1.2 million units) in 2.3 ml syringe – 1% DV Dec-18 | to 2021 344.93 | 10 | Bicillin LA |
| | | 10 | |
| BENZYLPENICILLIN SODIUM [PENICILLIN G] | 05.00 | 05 | Den Denisillin O Cadium |
| Inj 600 mg (1 million units) vial – 1% DV Nov-20 to 2023 | | 25 | Pan-Penicillin G Sodium |
| (Pan-Penicillin G Sodium Inj 600 mg (1 million units) vial to be deliste | 11.09 1 November 2020 | 10 | Sandoz |
| | u i novembei 2020 |) | |
| FLUCLOXACILLIN | | | - · · · |
| Cap 250 mg - 1% DV Sep-18 to 2021 | | 250 | Staphlex |
| Cap 500 mg - 1% DV Sep-18 to 2021 | | 500 | Staphlex |
| Grans for oral liq 25 mg per ml – 1% DV Oct-18 to 2021 | | 100 ml | AFT |
| Grans for oral liq 50 mg per ml – 1% DV Oct-18 to 2021 | | 100 ml | AFT |
| Inj 250 mg vial | | 10 | Flucloxin |
| Inj 500 mg vial | | 10 | Flucloxin |
| Inj 1 g vial – 1% DV Nov-20 to 2023 | 5.70 | 5 | Flucil |
| PHENOXYMETHYLPENICILLIN [PENICILLIN V] | | | |
| Cap 250 mg - 1% DV Sep-18 to 2021 | | 50 | Cilicaine VK |
| Cap 500 mg - 1% DV Sep-18 to 2021 | 4.26 | 50 | Cilicaine VK |
| Grans for oral liq 125 mg per 5 ml - 1% DV Jan-20 to 2022 | 2.99 | 100 ml | AFT |
| Grans for oral liq 250 mg per 5 ml - 1% DV Jan-20 to 2022 | 3.99 | 100 ml | AFT |
| PIPERACILLIN WITH TAZOBACTAM – Restricted see terms below | 1 | | |
| Inj 4 g with tazobactam 0.5 g vial | | 10 | PipTaz Sandoz |
| , , | | | PiperTaz Sandoz |
| → Restricted (RS1053) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory spe | cialist | | |
| PROCAINE PENICILLIN | | | |
| Inj 1.5 g in 3.4 ml syringe | | 5 | Cilicaine |
| TICARCILLIN WITH CLAVULANIC ACID – Restricted see terms be | | č | |
| | | | |
| Inj 3 g with clavulanic acid 0.1 mg vial → Restricted (RS1054) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory spe | cialist | | |

Clinical microbiologist, infectious disease specialist or respiratory specialist

e.g. Brand indicates brand example only. It is not a contracted product.

INFECTIONS

| | Price (ex man. excl. GST \$ | Per | Brand or Generic Manufacturer |
|--|--|--|--|
| Quinolones | | | |
| CIPROFLOXACIN – Restricted see terms below | | | |
| Tab 250 mg – 1% DV Nov-20 to 2023 | 2.42 | 28 | Cipflox |
| Tab 500 mg – 1% DV Nov-20 to 2023 | 3.40 | 28 | Cipflox |
| Tab 750 mg - 1% DV Nov-20 to 2023 | 5.95 | 28 | Cipflox |
| Oral liq 50 mg per ml | | | |
| Oral liq 100 mg per ml | 00.00 | 10 | Oinflau |
| Inj 2 mg per ml, 100 ml bag – 1% DV Oct-18 to 2021 → Restricted (RS1055) | | 10 | Cipflox |
| Clinical microbiologist or infectious disease specialist | | | |
| | | | |
| MOXIFLOXACIN – Restricted see terms below | 50.00 | F | Avalov |
| Tab 400 mg Inj 1.6 mg per ml, 250 ml bottle – 1% DV Apr-20 to 2022 | | 5 1 | Avelox Moxifloxacin Kabi |
| The stricted (RS1644) | | I | |
| Initiation – Mycobacterium infection | | | |
| Infectious disease specialist, clinical microbiologist or respiratory s | specialist | | |
| Any of the following: | | | |
| 1 Both: | | | |
| 1.1 Active tuberculosis: and | | | |
| 1.2 Any of the following: | | | |
| | | | |
| | line medications: or | | |
| 1.2.1 Documented resistance to one or more first- 1.2.2 Suspected resistance to one or more first-lin | ' | sis assun | ned to be contracted in an |
| 1.2.1 Documented resistance to one or more first- | ne medications (tuberculo | | |
| 1.2.1 Documented resistance to one or more first- 1.2.2 Suspected resistance to one or more first-lin area with known resistance), as part of regir 1.2.3 Impaired visual acuity (considered to preclude) | ne medications (tuberculo nen containing other seco de ethambutol use); or | ond-line a | gents; or |
| 1.2.1 Documented resistance to one or more first- 1.2.2 Suspected resistance to one or more first-lin area with known resistance), as part of regir 1.2.3 Impaired visual acuity (considered to preclud 1.2.4 Significant pre-existing liver disease or hepa | ne medications (tuberculo nen containing other seco de ethambutol use); or ttotoxicity from tuberculos | ond-line a | gents; or ations; or |
| 1.2.1 Documented resistance to one or more first- 1.2.2 Suspected resistance to one or more first-lin area with known resistance), as part of regir 1.2.3 Impaired visual acuity (considered to preclud 1.2.4 Significant pre-existing liver disease or hepa 1.2.5 Significant documented intolerance and/or s | ne medications (tuberculo nen containing other seco de ethambutol use); or ttotoxicity from tuberculos | ond-line a | gents; or ations; or |
| 1.2.1 Documented resistance to one or more first- 1.2.2 Suspected resistance to one or more first-lin area with known resistance), as part of regin 1.2.3 Impaired visual acuity (considered to preclud 1.2.4 Significant pre-existing liver disease or hepa 1.2.5 Significant documented intolerance and/or s or | ne medications (tuberculo men containing other seco de ethambutol use); or totoxicity from tuberculos ide effects following a rea | ond-line a is medica asonable | gents; or ations; or trial of first-line medications; |
| 1.2.1 Documented resistance to one or more first-lin area with known resistance), as part of regir 1.2.3 Impaired visual acuity (considered to preclud 1.2.4 Significant pre-existing liver disease or hepa 1.2.5 Significant documented intolerance and/or s or 2 Mycobacterium avium-intracellulare complex not respondir | ne medications (tuberculo nen containing other seco de ethambutol use); or totoxicity from tuberculos ide effects following a rea ng to other therapy or who | ond-line a is medica asonable ere such t | gents; or ations; or trial of first-line medications; herapy is contraindicated; or |
| 1.2.1 Documented resistance to one or more first-lin area with known resistance), as part of regir 1.2.3 Impaired visual acuity (considered to preclud 1.2.4 Significant pre-existing liver disease or hepa 1.2.5 Significant documented intolerance and/or s or 2 Mycobacterium avium-intracellulare complex not respondir 3 Patient is under five years of age and has had close contact | ne medications (tuberculo nen containing other seco de ethambutol use); or totoxicity from tuberculos ide effects following a rea ng to other therapy or who | ond-line a is medica asonable ere such t | gents; or ations; or trial of first-line medications; herapy is contraindicated; or |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance, as part of regin | ne medications (tuberculo nen containing other seco de ethambutol use); or totoxicity from tuberculos ide effects following a rea ng to other therapy or who | ond-line a is medica asonable ere such t | gents; or ations; or trial of first-line medications; herapy is contraindicated; or |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resistance, as part of reginarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resisting liver disease or here a so or a so or | ne medications (tuberculo nen containing other seco de ethambutol use); or totoxicity from tuberculos ide effects following a rea ng to other therapy or who | ond-line a is medica asonable ere such t | gents; or ations; or trial of first-line medications; herapy is contraindicated; or |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resistance, as part of reginarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resistancea with known resi | the medications (tuberculo nen containing other seco de ethambutol use); or totoxicity from tuberculos ide effects following a rea ng to other therapy or who ct with a confirmed multi- | ond-line a sis medica asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resistance, as part of reginarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resistancea with known resi | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or who ct with a confirmed multi- | ond-line a asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or tant tuberculosis case. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance and/or so reginarea with known resistance and/or so reginarea with known resistance and/or so reginarea with known resistance and has had close contained interfectious disease specialist or clinical microbiologist with the previous that is unresistened with previous and that is unresistened with previous known resistened with the previous kn | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or who ct with a confirmed multi- | ond-line a asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or tant tuberculosis case. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of regir 1.2.3 Impaired visual acuity (considered to preclud 1.2.4 Significant pre-existing liver disease or hepa 1.2.5 Significant documented intolerance and/or s or 2 Mycobacterium avium-intracellulare complex not respondir 3 Patient is under five years of age and has had close contactinitation – Pneumonia Infectious disease specialist or clinical microbiologist Either: 1 Immunocompromised patient with pneumonia that is unres 2 Pneumococcal pneumonia or other invasive pneumococca | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or who ct with a confirmed multi- | ond-line a asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or tant tuberculosis case. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resisting is part of reginarea with known resisting is part of reginarea with known resisting is part of reginarea with known resistence and/or so reginarea with known resistence and/or so reginarea with known resistence and has had close contarea initiation – Penetrating eye injury | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a read ing to other therapy or who ct with a confirmed multi- sponsive to first-line treatr al disease highly resistant | ond-line a asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or tant tuberculosis case. |
| 1.2.1 Documented resistance to one or more first- inarea with known resistance), as part of reginarea with known resistance with preumonia that is unrese. Pneumococcal pneumonia or other invasive pneumococca initiation – Penetrating eye injury Ophthalmologist Five days treatment for patients requiring prophylaxis following a properties of the set of the set of the set of the resistance with preuminarea with known resistance with the set of the resistance with the set of the set of the set of the resistance with the resistance with the set of the resistance with the resistance with the resistance with the set of the resistance with the resistancea with the resistance with the resistance with the resistance | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a read ing to other therapy or who ct with a confirmed multi- sponsive to first-line treatr al disease highly resistant | ond-line a asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or tant tuberculosis case. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance with preumonia that is unresistener. 1 Immunocompromised patient with pneumonia that is unresistener. 2 Pneumococcal pneumonia or other invasive pneumococcal initiation – Penetrating eye injury Dophthalmologist Five days treatment for patients requiring prophylaxis following a partialition – Mycoplasma genitalium | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a read ing to other therapy or who ct with a confirmed multi- sponsive to first-line treatr al disease highly resistant | ond-line a asonable ere such t drug resis | gents; or ations; or trial of first-line medications; herapy is contraindicated; or tant tuberculosis case. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance with preumonia that is unresistered with preumococcal preumococcal preumonia or other invasive preumococcal initiation – Penetrating eye injury Ophthalmologist Five days treatment for patients requiring prophylaxis following a partialium All of the following: | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea- ng to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr I disease highly resistant benetrating eye injury. | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance with preumonia that is unresistener. 1 Immunocompromised patient with pneumonia that is unresistener. 2 Pneumococcal pneumonia or other invasive pneumococcal initiation – Penetrating eye injury Dophthalmologist Five days treatment for patients requiring prophylaxis following a partialition – Mycoplasma genitalium | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea- ng to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr I disease highly resistant benetrating eye injury. | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance with responding the following a part of reginarea with known resistence with the following: 1 Has nucleic acid amplification test (NAAT) confirmed Myconarea 2 Either: | The medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr al disease highly resistant penetrating eye injury. | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance and responding the following: 1 Has nucleic acid amplification test (NAAT) confirmed Mycci 2 Either: 2.1 Has tried and failed to clear infection using azithron | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr and isease highly resistant benetrating eye injury. | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance with responding the following a part of reginarea with known resistence with the following: 1 Has nucleic acid amplification test (NAAT) confirmed Myconarea 2 Either: | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr and isease highly resistant benetrating eye injury. | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance and/or so or 2 Mycobacterium avium-intracellulare complex not respondin 3 Patient is under five years of age and has had close contains integes as specialist or clinical microbiologist Either: 1 Immunocompromised patient with pneumonia that is unress 2 Pneumococcal pneumonia or other invasive pneumococcal initiation – Penetrating eye injury Dophthalmologist Five days treatment for patients requiring prophylaxis following a pinitiation – Mycoplasma genitalium All of the following: 1 Has nucleic acid amplification test (NAAT) confirmed Mycore 2 Either: 2.1 Has tried and failed to clear infection using azithron 2.2 Has laboratory confirmed azithromycin resistance; a 3 Treatment is only for 7 days. | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea ing to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr and isease highly resistant benetrating eye injury. | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |
| 1.2.1 Documented resistance to one or more first-linarea with known resistance), as part of reginarea with known resistance, as part of reginarea with known resistance, and the state of the s | the medications (tuberculo men containing other second de ethambutol use); or totoxicity from tuberculos ide effects following a rea- ng to other therapy or whe ct with a confirmed multi- sponsive to first-line treatr I disease highly resistant penetrating eye injury. Splasma genitalium and is nycin; or and | ond-line a sis medica asonable ere such t drug resis nent; or to other a | gents; or ations; or trial of first-line medications; herapy is contraindicated; or stant tuberculosis case. antibiotics. |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|----------|-------------------------------------|
| Tetracyclines | | | |
| DEMECLOCYCLINE HYDROCHLORIDE Tab 150 mg Cap 150 mg Cap 300 mg DOXYCYCLINE → Tab 50 mg - Restricted: For continuation only Tab 100 mg - Inj 5 mg per ml, 20 ml vial MINOCYCLINE Tab 50 mg → Cap 100 mg - Restricted: For continuation only | 64.43 | 500 | Doxine |
| TETRACYCLINE Tab 250 mg Cap 500 mg (<i>Tetracyclin Wolff Cap 500 mg to be delisted 1 December 2020</i>) TIGECYCLINE – Restricted see terms below ↓ Inj 50 mg vial → Restricted (R\$1059) Clinical microbiologist or infectious disease specialist | | 28 30 | Accord Tetracyclin Wolff |
| Other Antibacterials | | | |
| AZTREONAM – Restricted see terms below ↓ Inj 1 g vial → Restricted (RS1277) Clinical microbiologist or infectious disease specialist CHLORAMPHENICOL – Restricted see terms below ↓ Inj 1 g vial → Restricted (RS1277) Clinical microbiologist or infectious disease specialist CLINDAMYCIN – Restricted see terms below | | 10 | Azactam |
| | 4.61 | 24 | Dalacin C |
| ↓ Oral liq 15 mg per ml ↓ Inj 150 mg per ml, 4 ml ampoule - 1% DV Oct-19 to 2022 → Restricted (RS1061) Clinical microbiologist or infectious disease specialist | | 10 | Dalacin C |
| COLISTIN SULPHOMETHATE [COLESTIMETHATE] – Restricted se Inj 150 mg per ml, 1 ml vial Restricted (RS1062) Clinical microbiologist, infectious disease specialist or respiratory speci DAPTOMYCIN – Restricted see terms below | 65.00 | 1 | Colistin-Link |
| Inj 500 mg vial | 243.52 | 1 | Cubicin |
| FOSFOMYCIN – Restricted see terms on the next page Powder for oral solution, 3 g sachet | | | e.g. UroFos |

| | Price | | Brand or |
|--|---------------------|---------|---------------------------|
| | (ex man. excl. GST) | Per | Generic |
| | \$ | rei | Manufacturer |
| → Restricted (RS1315) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| LINCOMYCIN – Restricted see terms below | | | |
| Inj 300 mg per ml, 2 ml vial | | | |
| → Restricted (RS1065) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| LINEZOLID – Restricted see terms below | | | _ |
| Tab 600 mg – 1% DV Oct-18 to 2021 | | 10 | Zyvox |
| Oral liq 20 mg per ml – 1% DV Dec-18 to 2021 | | 150 ml | Zyvox |
| ↓ Inj 2 mg per ml, 300 ml bottle – 1% DV Feb-19 to 2021 | | 1 | Linezolid Kabi |
| → Restricted (RS1066) Clinical microbiologist or infectious disease specialist | | | |
| | | | |
| METHENAMINE (HEXAMINE) HIPPURATE | 40.01 | 100 | Linnov |
| Tab 1 g | 40.01 | 100 | Hiprex |
| NITROFURANTOIN | | | |
| Tab 50 mg - 1% DV Apr-19 to 2021 | | 100 | Nifuran |
| Tab 100 mg - 1% DV Apr-19 to 2021 | | 100 | Nifuran |
| PIVMECILLINAM – Restricted see terms below | | | |
| Tab 200 mg | | | |
| → Restricted (RS1322) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| SODIUM FUSIDATE [FUSIDIC ACID] – Restricted see terms below | | | |
| Tab 250 mg | | 12 | Fucidin |
| → Restricted (RS1064) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| SULPHADIAZINE – Restricted see terms below | | | |
| Tab 500 mg | | | |
| → Restricted (RS1067) | adicina anacialist | | |
| Clinical microbiologist, infectious disease specialist or maternal-foetal m | ledicine specialist | | |
| TEICOPLANIN – Restricted see terms below | 50.50 | | - · · · · · |
| ↓ Inj 400 mg vial – 1% DV Jul-20 to 2021 | | 1 | Teicoplanin Mylan |
| → Restricted (RS1068) Clinical microbiologist or infectious disease specialist | | | |
| . | | | |
| TRIMETHOPRIM | | | |
| Tab 100 mg Tab 300 mg – 1% DV Oct-18 to 2021 | 16 50 | 50 | ТМР |
| 5 | | 50 | |
| TRIMETHOPRIM WITH SULPHAMETHOXAZOLE [CO-TRIMOXAZOLI | =] | | |
| Tab 80 mg with sulphamethoxazole 400 mg Oral lig 8 mg with sulphamethoxazole 40 mg per ml | 2.07 | 100 ml | Donrim |
| Inj 16 mg with sulphamethoxazole 80 mg per ml, 5 ml ampoule | 2.97 | 100 111 | Deprim |
| | | | |
| VANCOMYCIN – Restricted see terms below | 0.05 | 4 | Mulan |
| Inj 500 mg vial – 1% DV Oct-20 to 2023 → Restricted (RS1069) | 2.35 | 1 | Mylan |
| Clinical microbiologist or infectious disease specialist | | | |
| טווווינמו ווווניטטוטוטעוזי טו וווובנווטעז עוזבמשב ארבומווזי | | | |

INFECTIONS



| (ex mai | Price n. excl. | GST) | Per | Brand or Generic Manufacturor |
|--|--|--|--|--|
| | \$ | | Per | Manufacturer |
| Antifungals | | | | |
| Imidazoles | | | | |
| KETOCONAZOLE ↓ Tab 200 mg → Restricted (RS1410) Dncologist | | | | |
| Polyene Antimycotics | | | | |
| AMPHOTERICIN B Inj (liposomal) 50 mg vial3 | ,450.00 |) | 10 | AmBisome |
| → Restricted (RS1071) | | | | |
| nitiation Clinical microbiologist, haematologist, infectious disease specialist, oncologist, Either: | respirat | tory sp | ecialist c | r transplant specialist |
| Proven or probable invasive fungal infection, to be prescribed under an Both: | establis | shed pr | otocol; c | r |
| 2.1 Possible invasive fungal infection; and2.2 A multidisciplinary team (including an infectious disease physicia treatment to be appropriate. | an or a c | clinical | microbic | logist) considers the |
| Inj 50 mg vial | | | | |
| | respira | tory sp | ecialist c | r transplant specialist |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, | respira | tory sp | ecialist c | r transplant specialist |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, | 17.09 |) | ecialist c 50 50 | r transplant specialist Nilstat Nilstat |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u | 17.09 |) | 50 | Nilstat |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below | 17.09 15.47 | , , | 50 50 | Nilstat Nilstat |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 | 17.09 15.47 2.75 |) , ; | 50 50 28 | Nilstat Nilstat Mylan |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles ELUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 | 17.09 15.47 2.75 0.65 |) , ; | 50 50 28 1 | Nilstat Nilstat Mylan Mylan |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, IYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 | 17.09 15.47 2.75 0.65 12.89 | ; | 50 50 28 1 28 | Nilstat Nilstat Mylan Mylan Mylan |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, IYSTATIN Tab 500,000 u Cap 500,000 u Triazoles CLUCONAZOLE - Restricted see terms below Cap 50 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Cap 200 mg - 1% DV Nov-20 to 200 mg - 1% | 17.09 15.47 2.75 0.65 12.89 98.50 | ; | 50 50 28 1 | Nilstat Nilstat Mylan Mylan |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles ELUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 | | 50 50 28 1 28 35 ml | Nilstat Nilstat Mylan Mylan Mylan Diflucan |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, VYSTATIN Tab 500,000 u Cap 500,000 u Triazoles *LUCONAZOLE - Restricted see terms below Cap 50 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Cap 200 mg - 1% DV Nov-20 to 2023 Cap 200 mg - 1% DV Nov-20 to 2023 Ing 2 mg per ml, 50 ml vial - 1% DV Oct-19 to 2022 Ing 2 mg per ml, 100 ml vial - 1% DV Oct-19 to 2022 Restricted (RS1072) | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 | | 50 50 28 1 28 35 ml 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, YYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE - Restricted see terms below Cap 50 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Inj 2 mg per ml, 50 ml vial - 1% DV Oct-19 to 2022 Inj 2 mg per ml, 100 ml vial - 1% DV Oct-19 to 2022 Restricted (RS1072) Consultant | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 | | 50 50 28 1 28 35 ml 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Cap 10 mg per 5 ml Inj 2 mg per ml, 50 ml vial – 1% DV Oct-19 to 2022 Inj 2 mg per ml, 100 ml vial – 1% DV Oct-19 to 2022 Restricted (RS1072) Consultant TRACONAZOLE – Restricted see terms below | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 3.45 | | 50 50 28 1 28 35 ml 1 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris Fluconazole-Claris |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE - Restricted see terms below Cap 50 mg - 1% DV Nov-20 to 2023 Cap 150 mg - 1% DV Nov-20 to 2023 Cap 200 mg - 1% DV Nov-20 to 2023 Cap 200 mg - 1% DV Nov-20 to 2023 Inj 2 mg per ml, 50 ml vial - 1% DV Oct-19 to 2022 Inj 2 mg per ml, 50 ml vial - 1% DV Oct-19 to 2022 Festricted (RS1072) Consultant TRACONAZOLE - Restricted see terms below Cap 100 mg - 1% DV Nov-19 to 2022 | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 3.45 | | 50 50 28 1 28 35 ml 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, YYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Cap 150 mg per sml Inj 2 mg per ml, 50 ml vial – 1% DV Oct-19 to 2022 Inj 2 mg per ml, 100 ml vial – 1% DV Oct-19 to 2022 Fastricted (RS1072) Consultant TRACONAZOLE – Restricted see terms below Cap 100 mg – 1% DV Nov-19 to 2022 | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 3.45 | | 50 50 28 1 28 35 ml 1 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris Fluconazole-Claris |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles ELUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Cap 150 mg per 5 ml Inj 2 mg per ml, 50 ml vial – 1% DV Oct-19 to 2022 Inj 2 mg per ml, 100 ml vial – 1% DV Oct-19 to 2022 Restricted (RS1072) Consultant TRACONAZOLE – Restricted see terms below Cap 100 mg – 1% DV Nov-19 to 2022 Oral liquid 10 mg per ml Restricted (RS1073) | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 3.45 3.45 | ;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;; | 50 50 28 1 28 35 ml 1 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris Fluconazole-Claris |
| Clinical microbiologist, haematologist, infectious disease specialist, oncologist, NYSTATIN Tab 500,000 u Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Inj 2 mg per ml, 100 ml vial – 1% DV Oct-19 to 2022 Inj 2 mg per ml, 100 ml vial – 1% DV Oct-19 to 2022 Consultant TRACONAZOLE – Restricted see terms below Cap 100 mg – 1% DV Nov-19 to 2022 Oral liquid 10 mg per ml Restricted (RS1073) Clinical immunologist, clinical microbiologist, dermatologist or infectious diseas | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 3.45 3.45 | ;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;; | 50 50 28 1 28 35 ml 1 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris Fluconazole-Claris |
| Cap 500,000 u Triazoles FLUCONAZOLE – Restricted see terms below Cap 50 mg – 1% DV Nov-20 to 2023 Cap 150 mg – 1% DV Nov-20 to 2023 Cap 200 mg – 1% DV Nov-20 to 2023 Oral liquid 50 mg per 5 ml Inj 2 mg per ml, 50 ml vial – 1% DV Oct-19 to 2022 Inj 2 mg per ml, 100 ml vial – 1% DV Oct-19 to 2022 Restricted (RS1072) Consultant TRACONAZOLE – Restricted see terms below Cap 100 mg – 1% DV Nov-19 to 2022 | 17.09 15.47 2.75 0.65 12.89 98.50 2.80 3.45 3.45 |) ; ; ; ; ; | 50 50 28 1 28 35 ml 1 1 | Nilstat Nilstat Mylan Mylan Diflucan Fluconazole-Claris Fluconazole-Claris |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| \$ Per Manufacturer | Price Brand or (ex man. excl. GST) Generic |
|---------------------|---|
|---------------------|---|

➡ Restricted (RS1074)

Initiation

Haematologist or infectious disease specialist *Re-assessment required after 6 weeks* Both:

Both:

- 1 Either:
 - 1.1 Patient has acute myeloid leukaemia; or
 - 1.2 Patient is planned to receive a stem cell transplant and is at high risk for aspergillus infection; and
- 2 Patient is to be treated with high dose remission induction therapy or re-induction therapy.

Continuation

Haematologist or infectious disease specialist

Re-assessment required after 6 weeks

Both:

1 Patient has previously received posaconazole prophylaxis during remission induction therapy; and

- 2 Any of the following:
 - 2.1 Patient is to be treated with high dose remission re-induction therapy; or
 - 2.2 Patient is to be treated with high dose consolidation therapy; or
 - 2.3 Patient is receiving a high risk stem cell transplant.

VORICONAZOLE - Restricted see terms below

| t | Tab 50 mg - 1% DV Sep-18 to 2021 | 56 | Vttack |
|---|--|-------|------------|
| t | Tab 200 mg - 1% DV Sep-18 to 2021 | 56 | Vttack |
| t | Powder for oral suspension 40 mg per ml - 1% DV Dec-18 to 20211,437.00 | 70 ml | Vfend |
| | Inj 200 mg vial - 1% DV Oct-19 to 2022 | 1 | Neo Health |
| | | | |

→ Restricted (RS1075)

Initiation - Proven or probable aspergillus infection

Clinical microbiologist, haematologist or infectious disease specialist Both:

- 1 Patient is immunocompromised; and
- 2 Patient has proven or probable invasive aspergillus infection.

Initiation – Possible aspergillus infection

Clinical microbiologist, haematologist or infectious disease specialist All of the following:

- 1 Patient is immunocompromised; and
- 2 Patient has possible invasive aspergillus infection; and
- 3 A multidisciplinary team (including an infectious disease physician) considers the treatment to be appropriate.

Initiation - Resistant candidiasis infections and other moulds

Clinical microbiologist, haematologist or infectious disease specialist

All of the following:

- 1 Patient is immunocompromised; and
- 2 Either:
 - 2.1 Patient has fluconazole resistant candidiasis; or
 - 2.2 Patient has mould strain such as Fusarium spp. and Scedosporium spp; and
- 3 A multidisciplinary team (including an infectious disease physician or clinical microbiologist) considers the treatment to be appropriate.

Other Antifungals

| CA | SPOFUNGIN - Restricted see terms on the next page | | | |
|----|---|--------|---|------------|
| | Inj 50 mg vial - 1% DV Dec-19 to 2022 | 220.28 | 1 | Max Health |
| t | Inj 70 mg vial - 1% DV Dec-19 to 2022 | 284.63 | 1 | Max Health |

Products with Hospital Supply Status (HSS) are in **bold**

Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | (ex man | Price . excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|------------|------------------------|----------|-----------|-------------------------------------|
| → Restricted (RS1076) | | | | | |
| Initiation Clinical microbiologist, haematologist, infectious disease specialist, onco Either: | ologist, r | espira | atory sp | oecialist | or transplant specialist |
| 1 Proven or probable invasive fungal infection, to be prescribed un 2 Both: | der an e | establi | shed p | rotocol; | or |
| 2.1 Possible invasive fungal infection; and2.2 A multidisciplinary team (including an infectious disease p treatment to be appropriate. | hysicia | n or a | clinical | microbi | ologist) considers the |
| FLUCYTOSINE – Restricted see terms below | | | | | |
| Cap 500 mg | | | | | |
| Restricted (RS1279) Clinical microbiologist or infectious disease specialist | | | | | |
| TERBINAFINE | | | | | |
| Tab 250 mg | | 1.3 | 3 | 14 | Deolate |
| Antimycobacterials | | | | | |
| Antileprotics | | | | | |
| CLOFAZIMINE – Restricted see terms below | | | | | |
| Cap 50 mg | | | | | |
| → Restricted (RS1077) | | | | | |
| Clinical microbiologist, dermatologist or infectious disease specialist | | | | | |
| DAPSONE – Restricted see terms below Tab 25 mg | | 268 5 | 0 | 100 | Dapsone |
| ↓ Tab 20 mg | | | | 100 | Dapsone |
| → Restricted (RS1078) | | | | | · |
| Clinical microbiologist, dermatologist or infectious disease specialist | | | | | |
| Antituberculotics | | | | | |
| CYCLOSERINE – Restricted see terms below | | | | | |
| Cap 250 mg → Restricted (RS1079) | | | | | |
| Clinical microbiologist, infectious disease specialist or respiratory specia | list | | | | |
| ETHAMBUTOL HYDROCHLORIDE – Restricted see terms below | | | | | |
| ↓ Tab 100 mg | | | | | |
| Tab 400 mg | | 49.3 | 4 | 56 | Myambutol |
| → Restricted (RS1080) | 1:-+ | | | | |
| Clinical microbiologist, infectious disease specialist or respiratory specia | liist | | | | |
| ISONIAZID – Restricted see terms below Tab 100 mg – 1% DV Oct-18 to 2021 | | 220 | 0 | 100 | PSM |
| → Restricted (RS1281) | | 0 | 0 | 100 | |
| Clinical microbiologist, dermatologist, paediatrician, public health physic | ian or in | ternal | medic | ine phys | ician |
| ISONIAZID WITH RIFAMPICIN - Restricted see terms below | | | | | |
| Tab 100 mg with rifampicin 150 mg - 1% DV Sep-18 to 2021 | | .85.5 | 4 | 100 | Rifinah |
| ■ Tab 150 mg with rifampicin 300 mg – 1% DV Sep-18 to 2021 | | | | 100 | Rifinah |
| Restricted (RS1282) | | | | | |

Clinical microbiologist, dermatologist, paediatrician, public health physician or internal medicine physician

INFECTIONS

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|---------------|-------------------------------------|
| PARA-AMINOSALICYLIC ACID – Restricted see terms below | | | |
| Grans for oral lig 4 g | | 30 | Paser |
| → Restricted (RS1083) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory spe | ecialist | | |
| PROTIONAMIDE – Restricted see terms below | | | |
| Tab 250 mg | 305.00 | 100 | Peteha |
| → Restricted (RS1084) | | 100 | 1 otonu |
| Clinical microbiologist, infectious disease specialist or respiratory spe | ecialist | | |
| PYRAZINAMIDE – Restricted see terms below | | | |
| Tab 500 mg | | | |
| → Restricted (RS1085) | | | |
| Clinical microbiologist, infectious disease specialist or respiratory spe | ocialist | | |
| | coldiist | | |
| RIFABUTIN – Restricted see terms below | 000 75 | 20 | Mucobutin |
| Cap 150 mg | | 30 | Mycobutin |
| → Restricted (RS1086) | int or reapiratory apopia | iot | |
| Clinical microbiologist, gastroenterologist, infectious disease speciali | ist or respiratory special | ISL | |
| RIFAMPICIN – Restricted see terms below | | | |
| Cap 150 mg - 1% DV Nov-20 to 2023 | | 100 | Rifadin |
| Cap 300 mg - 1% DV Nov-20 to 2023 | | 100 | Rifadin |
| Oral liq 100 mg per 5 ml – 1% DV Nov-20 to 2023 | | 60 ml | Rifadin |
| Inj 600 mg vial – 1% DV Nov-20 to 2023 | 134.98 | 1 | Rifadin |
| → Restricted (RS1087) | a diatulaiana ay ay dalla ka | مريما مرافا م | -1 |
| Clinical microbiologist, dermatologist, internal medicine physician, pa | | aitir priys | GIAIT |
| Antiparasitics | | | |
| Anthelmintics | | | |
| ALBENDAZOLE – Restricted see terms below | | | |
| Tab 200 mg | | | |
| Tab 400 mg | | | |
| - Postrieted (PS1099) | | | |
| | | | |
| | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below | | 4 | Stromectol |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below I Tab 3 mg | | 4 | Stromectol |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below Tab 3 mg | | 4 | Stromectol |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below ↓ Tab 3 mg → Restricted (RS1283) Clinical microbiologist, dermatologist or infectious disease specialist | | 4 | Stromectol |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below ↓ Tab 3 mg → Restricted (RS1283) Clinical microbiologist, dermatologist or infectious disease specialist MEBENDAZOLE | | · | |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below ↓ Tab 3 mg → Restricted (RS1283) Clinical microbiologist, dermatologist or infectious disease specialist MEBENDAZOLE Tab 100 mg | | 4 24 | Stromectol De-Worm |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below ↓ Tab 3 mg | | · | |
| Oral liq 100 mg per 5 ml PRAZIQUANTEL | | · | |
| Clinical microbiologist or infectious disease specialist VERMECTIN – Restricted see terms below ↓ Tab 3 mg | | · | |
| Clinical microbiologist or infectious disease specialist VERMECTIN - Restricted see terms below ↓ Tab 3 mg | | · | |
| Clinical microbiologist or infectious disease specialist VERMECTIN - Restricted see terms below ↓ Tab 3 mg | 24.19 | · | |

→ Restricted (RS1090) Clinical microbiologist or infectious disease specialist

| | Price | | Brand or |
|---|---------------------------|--------|-------------------------|
| | (ex man. excl. GST) \$ | Per | Generic Manufacturer |
| ARTESUNATE – Restricted see terms below | | | |
| Inj 60 mg vial | | | |
| → Restricted (RS1091) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| ATOVAQUONE WITH PROGUANIL HYDROCHLORIDE - Restricted | ed see terms below | | |
| Tab 62.5 mg with proguanil hydrochloride 25 mg | | 12 | Malarone Junior |
| Tab 250 mg with proguanil hydrochloride 100 mg | 64.00 | 12 | Malarone |
| → Restricted (RS1092) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| CHLOROQUINE PHOSPHATE – Restricted see terms below | | | |
| Tab 250 mg | | | |
| → Restricted (RS1093) Clinical microbiologist, dermatologist, infectious disease specialist or | rhaumatalagiat | | |
| | meumatologist | | |
| MEFLOQUINE – Restricted see terms below I Tab 250 mg | | | |
| Tab 250 mg → Restricted (RS1094) | | | |
| Clinical microbiologist, dermatologist, infectious disease specialist or | rheumatologist | | |
| METRONIDAZOLE | riouniatologiot | | |
| Tab 200 mg | 10.45 | 100 | Trichozole |
| Tab 400 mg | | 100 | Trichozole |
| Oral lig benzoate 200 mg per 5 ml | | 100 ml | Flagyl-S |
| Injection 5 mg per ml, 100 ml bottle | 1.39 | 100 ml | AFT |
| Inj 5 mg per ml, 100 ml bottle | 34.80 | 20 | Colpocin-T |
| Inj 5 mg per ml, 100 ml bag | | 10 | Baxter |
| Suppos 500 mg | 24.48 | 10 | Flagyl |
| (Trichozole Tab 200 mg to be delisted 1 September 2020) | | | |
| (Trichozole Tab 400 mg to be delisted 1 September 2020) | | | |
| NITAZOXANIDE – Restricted see terms below | | | |
| Tab 500 mg | 1,680.00 | 30 | Alinia |
| ↓ Oral liq 100 mg per 5 ml | | | |
| → Restricted (RS1095) Clinical microbiologist or infectious disease specialist | | | |
| | | | |
| ORNIDAZOLE Tab 500 mg | 22.05 | 10 | Arrow-Ornidazole |
| 5 | | 10 | Anow-Oniuazoie |
| PENTAMIDINE ISETHIONATE – Restricted see terms below | 016.00 | 5 | Pentacarinat |
| Inj 300 mg vial - 1% DV Nov-19 to 2022 → Restricted (RS1096) | 216.00 | Э | Pentacarinat |
| Clinical microbiologist or infectious disease specialist | | | |
| PRIMAQUINE – Restricted see terms below | | | |
| ↓ Tab 15 mg | | | |
| Tab 7.5 mg | | | |
| → Restricted (RS1097) | | | |
| Clinical microbiologist or infectious disease specialist | | | |
| PYRIMETHAMINE – Restricted see terms below | | | |
| ↓ Tab 25 mg | | | |
| → Restricted (RS1098) | | | |
| Clinical microbiologist, infectious disease specialist or maternal-foetal | medicine specialist | | |

INFECTIONS

| | (ex man | Price . excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------|------------------------|------|-----|-------------------------------------|
| QUININE DIHYDROCHLORIDE – Restricted see terms below Inj 60 mg per ml, 10 ml ampoule Inj 300 mg per ml, 2 ml vial Restricted (RS1099) Clinical microbiologist or infectious disease specialist QUININE SULPHATE Tab 300 mg SODIUM STIBOGLUCONATE – Restricted see terms below Inj 100 mg per ml, 1 ml vial Restricted (RS1100) Clinical microbiologist or infectious disease specialist SPIRAMYCIN – Restricted see terms below I Tab 500 mg Restricted (RS1101) Maternal-foetal medicine specialist | | 61.9 | 1 | 500 | Q 300 |
| Antiretrovirals Non-Nucleoside Reverse Transcriptase Inhibitors Restricted (RS1571) Initiation – Confirmed HIV Patient has confirmed HIV Infection. | | | | | |
| Initiation – Prevention of maternal transmission Either: 1 Prevention of maternal foetal transmission; or | | | | | |

2 Treatment of the newborn for up to eight weeks.

Initiation – Post-exposure prophylaxis following non-occupational exposure to HIV

Both:

- 1 Treatment course to be initiated within 72 hours post exposure; and
- 2 Any of the following:
 - 2.1 Patient has had unprotected receptive anal intercourse with a known HIV positive person; or
 - 2.2 Patient has shared intravenous injecting equipment with a known HIV positive person; or
 - 2.3 Patient has had non-consensual intercourse and the clinician considers that the risk assessment indicates prophylaxis is required.

Initiation – Percutaneous exposure

Patient has percutaneous exposure to blood known to be HIV positive.

EFAVIRENZ – **Restricted** see terms above

| t | Tab 200 mg | .190.15 | 90 | Stocrin |
|----|---------------------------------------|---------|--------|-----------------------|
| t | Tab 600 mg | 63.38 | 30 | Stocrin |
| t | Oral liq 30 mg per ml | | | |
| | RAVIRINE – Restricted see terms above | | | |
| t | Tab 200 mg | .770.00 | 60 | Intelence |
| NE | VIRAPINE – Restricted see terms above | | | |
| | Tab 200 mg - 1% DV Sep-18 to 2021 | | 60 | Nevirapine Alphapharm |
| t | Oral suspension 10 mg per ml | .203.55 | 240 ml | Viramune Suspension |

| | | Price excl. GS \$ | ST) Per | Brand or Generic Manufacturer |
|--|---------|-------------------------|--------------|-------------------------------------|
| Nucleoside Reverse Transcriptase Inhibitors | | | | |
| → Restricted (RS1572) | | | | |
| nitiation – Confirmed HIV Patient has confirmed HIV infection. | | | | |
| nitiation – Prevention of maternal transmission | | | | |
| Either: | | | | |
| 1 Prevention of maternal foetal transmission; or | | | | |
| 2 Treatment of the newborn for up to eight weeks. | | | | |
| nitiation – Post-exposure prophylaxis following non-occupational Both: | exposu | re to HIV | | |
| Treatment course to be initiated within 72 hours post exposure; a Any of the following: | Ind | | | |
| 2.1 Patient has had unprotected receptive anal intercourse w2.2 Patient has shared intravenous injecting equipment with a2.3 Patient has had non-consensual intercourse and the clini prophylaxis is required. | a known | HIV posi | tive person; | or |
| nitiation – Percutaneous exposure | | | | |
| Patient has percutaneous exposure to blood known to be HIV positive. | | | | |
| ABACAVIR SULPHATE – Restricted see terms above | | | | |
| Tab 300 mg - 1% DV Jul-19 to 2022 Oral lig 20 mg per ml | | | 60 240 ml | Ziagen Ziagen |
| ABACAVIR SULPHATE WITH LAMIVUDINE – Restricted see terms a | | -00.01 | 240 111 | Ziagon |
| Tab 600 mg with lamivudine 300 mg – 1% DV Jul-19 to 2022 | | .63.00 | 30 | Kivexa |
| EFAVIRENZ WITH EMTRICITABINE AND TENOFOVIR DISOPROXIL | - Restr | icted see | e terms abo | ve |
| t Tab 600 mg with emtricitabine 200 mg and tenofovir disoproxil 245 | | | | |
| (300 mg as a maleate) – 1% DV Jun-19 to 2022 | 1 | 06.88 | 30 | Mylan |
| EMTRICITABINE – Restricted see terms above | | | | |
| t Cap 200 mg - 1% DV Jul-19 to 2022 | 3 | 307.20 | 30 | Emtriva |
| LAMIVUDINE – Restricted see terms above Tab 150 mg – 1% DV Nov-20 to 2023 | | 01 50 | 60 | Lamivudine |
| | | .04.30 | 00 | Alphapharm |
| | | | | |
| STAVUDINE – Restricted see terms above | | | | |
| t Cap 40 mg | | | | |
| Powder for oral soln 1 mg per ml | | | | |
| ZIDOVI IDINE [AZT] – Bestricted see terms above | | | | |

| ZIDOVUDINE [AZT] – Restricted see terms above | | | |
|--|--------|--------|---------------|
| t Cap 100 mg | 152.25 | 100 | Retrovir |
| t Oral liq 10 mg per ml | | 200 ml | Retrovir |
| 1 Inj 10 mg per ml, 20 ml vial | | 5 | Retrovir IV |
| ZIDOVUDINE [AZT] WITH LAMIVUDINE - Restricted see terms above t Tab 300 mg with lamivudine 150 mg | | 60 | Alphapharm |
| | | 00 | ripriapriarii |

| Price Brand or (ex man. excl. GST) Generic \$ Per Manufacturer |
|--|
|--|

Protease Inhibitors

| → Restricted (RS1573) Initiation - Confirmed HIV Patient has confirmed HIV infection. Initiation - Prevention of maternal transmission Either: Prevention of maternal foetal transmission; or Treatment of the newborn for up to eight weeks. Initiation - Post-exposure prophylaxis following non-occupational exposure to HIV Both: | | |
|--|-------------|----------|
| 1 Treatment course to be initiated within 72 hours post exposure; and | | |
| Any of the following: 2.1 Patient has had unprotected receptive anal intercourse with a known HIV pos 2.2 Patient has shared intravenous injecting equipment with a known HIV positive 2.3 Patient has had non-consensual intercourse and the clinician considers that the prophylaxis is required. | e person; o | r |
| Initiation – Percutaneous exposure | | |
| Patient has percutaneous exposure to blood known to be HIV positive. | | |
| ATAZANAVIR SULPHATE - Restricted see terms above | | |
| t Cap 150 mg - 1% DV Jun-19 to 2022 | 60 60 | Teva |
| t Cap 200 mg - 1% DV Jun-19 to 2022 | 60 | Teva |
| DARUNAVIR – Restricted see terms above | 60 | Prezista |
| Tab 400 mg 335.00 Tab 600 mg 476.00 | 60 60 | Prezista |
| INDINAVIR – Restricted see terms above t Cap 200 mg t Cap 400 mg | 00 | 11021314 |
| LOPINAVIR WITH RITONAVIR – Restricted see terms above | | |
| t Tab 100 mg with ritonavir 25 mg 183.75 | 60 | Kaletra |
| t Tab 200 mg with ritonavir 50 mg | 120 | Kaletra |
| Cral liq 80 mg with ritonavir 20 mg per ml735.00 | 300 ml | Kaletra |
| RITONAVIR - Restricted see terms above t Tab 100 mg - 1% DV Jul-19 to 2022 | 30 | Norvir |
| Strand Transfer Inhibitors | | |

➡ Restricted (RS1574)

Initiation – Confirmed HIV

Patient has confirmed HIV infection.

Initiation – Prevention of maternal transmission Either:

- 1 Prevention of maternal foetal transmission; or
- 2 Treatment of the newborn for up to eight weeks.

Initiation – Post-exposure prophylaxis following non-occupational exposure to HIV Both:

| | P (ex man. | rice excl. G \$ | | Per | Brand or Generic Manufacturer |
|---|-----------------|-----------------------|--------|-------------|-------------------------------------|
| ontinued | | | | | |
| 1 Treatment course to be initiated within 72 hours post exposu | ire; and | | | | |
| 2 Any of the following:2.1 Patient has had unprotected receptive anal intercours | o with a know | ир ЦIV | nociti | | n: or |
| 2.1 Patient has hared intravenous injecting equipment w | | | | | |
| 2.3 Patient has had non-consensual intercourse and the prophylaxis is required. | | | | | |
| nitiation – Percutaneous exposure | | | | | |
| atient has percutaneous exposure to blood known to be HIV positi | ve. | | | | |
| OLUTEGRAVIR - Restricted see terms on the previous page | | | | | |
| Tab 50 mg | | 90.00 | | 30 | Tivicay |
| ALTEGRAVIR POTASSIUM - Restricted see terms on the previo | | | | | |
| Tab 400 mg Tab 600 mg | , | | | 60 | Isentress |
| Tab 600 mg | 1,0 | 90.00 | | 60 | Isentress HD |
| Antivirals | | | | | |
| Hepatitis B | | | | | |
| DEFOVIR DIPIVOXIL – Restricted see terms below | | | | | |
| Tab 10 mg | 6 | 70.00 | | 30 | Hepsera |
| Restricted (RS1104) | | | | | |
| itiation astroenterologist or infectious disease specialist | | | | | |
| Il of the following: | | | | | |
| 1 Patient has confirmed Hepatitis B infection (HBsAg+); and | | | | | |
| Documented resistance to lamivudine defined as: | | | | | |
| 2 Patient has raised serum ALT (> 1 × ULN); and | | | | | - 10 fold |
| 3 Patient has HBV DNA greater than 100,000 copies per mL, of 4 Detection of M204I or M204V mutation; and | or viral load g | reater | nan o | r equal 1 | o 10-told over hadir; and |
| 5 Either: | | | | | |
| 5.1 Both: | | | | | |
| 5.1.1 Patient is cirrhotic; and | | | | | |
| 5.1.2 Adefovir dipivoxil to be used in combination with | ith lamivudine | e; or | | | |
| 5.2 Both: | | | | | |
| 5.2.1 Patient is not cirrhotic; and | | | | | |
| 5.2.2 Adefovir dipivoxil to be used as monotherapy. | | | | | |
| NTECAVIR | | | | | |
| Tab 0.5 mg - 1% DV Nov-18 to 2021 | | 52.00 | | 30 | Entecavir Sandoz |
| | | 0.05 | | 00 | 7-41 |
| Tab 100 mg – 1% DV Nov-20 to 2023 Oral liq 5 mg per ml | | | n | 28 40 ml | Zetlam Zeffix |
| ENOFOVIR DISOPROXIL | 2 | 10.00 | 2 | 40 IIII | CCIIIX |
| | | | | | |

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|---------------------|--|
| Hepatitis C | | | |
| GLECAPREVIR WITH PIBRENTASVIR Note: the supply of treatment is via PHARMAC's approved dirr PHARMAC's website https://www.pharmac.govt.nz/hepatitis-c- | | Further de | etails can be found on |
| Tab 100 mg with pibrentasvir 40 mg LEDIPASVIR WITH SOFOSBUVIR - Restricted see terms below | 24,750.00 | 84 | Maviret |
| | 24,363.46 | 28 | Harvoni |
| Note: Only for use in patients with approval by the Hepatitis C Treat HepCTP at its regular meetings and approved subject to eligibility a Pharmaceutical Schedule). | | | |
| Herpesviridae | | | |
| ACICLOVIR Tab dispersible 200 mg − 1% DV Oct-19 to 2022 Tab dispersible 400 mg − 1% DV Oct-19 to 2022 Tab dispersible 800 mg − 1% DV Oct-19 to 2022 Inj 250 mg vial − 1% DV Sep-18 to 2021 CIDOFOVIR − Restricted see terms below I Inj 75 mg per ml, 5 ml vial → Restricted (RS1108) Clinical microbiologist, infectious disease specialist, otolaryngologis FOSCARNET SODIUM − Restricted see terms below I Inj 24 mg per ml, 250 ml bottle → Restricted (RS1109) Clinical microbiologist or infectious disease specialist GANCICLOVIR − Restricted see terms below I Inj 500 mg vial → Restricted (RS1110) Clinical microbiologist or infectious disease specialist | 5.38 5.98 9.60 | 25 56 35 5 | Lovir Lovir Aciclovir-Claris Cymevene |
| VALACICLOVIR Tab 500 mg - 1% DV Sep-18 to 2021 Tab 1,000 mg - 1% DV Sep-18 to 2021 | | 30 30 | Vaclovir Vaclovir |
| VALGANCICLOVIR – Restricted see terms below ↓ Tab 450 mg – 1% DV May-19 to 2021 → Restricted (RS1112) Initiation – Transplant cytomegalovirus prophylaxis Limited to 3 months treatment Patient has undergone a solid organ transplant and requires valgan Initiation – Lung transplant cytomegalovirus prophylaxis Limited to 6 months treatment Both: | | 60 laxis. | Valganciclovir Mylan |
| 1 Patient has undergone a lung transplant; and | | | |

Price

1 Patient has undergone a lung transplant; and

2 Either:

INFECTIONS

Brand or

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|-------------|-------------------------------------|
| | φ | Fel | Manulaciulei |
| continued 2.1 The donor was cytomegalovirus positive and the pa 2.2 The recipient is cytomegalovirus positive. | tient is cytomegalovirus ne | egative; o | r |
| Initiation – Cytomegalovirus in immunocompromised patients Both: | 6 | | |
| Patient is immunocompromised; and Any of the following: | | | |
| 2.1 Patient has cytomegalovirus syndrome or tissue inv2.2 Patient has rapidly rising plasma CMV DNA in abse2.3 Patient has cytomegalovirus retinitis. | | | |
| HIV Prophylaxis and Treatment | | | |
| EMTRICITABINE WITH TENOFOVIR DISOPROXIL - Restricted | see terms below | | |
| Tab 200 mg with tenofovir disoproxil 245 mg (300.6 mg as a s | | | _ |
| - 1% DV Jun-19 to 2022 | 61.15 | 30 | Teva |
| Initiation – Confirmed HIV | | | |
| Patient has confirmed HIV infection. | | | |
| Initiation – Prevention of maternal transmission Either: | | | |
| Prevention of maternal foetal transmission; or Treatment of the newborn for up to eight weeks. | | | |
| Initiation – Post-exposure prophylaxis following non-occupat Both: | - | | |
| Treatment course to be initiated within 72 hours post expose Any of the following: | | | |
| 2.1 Patient has had unprotected receptive anal intercou2.2 Patient has shared intravenous injecting equipment2.3 Patient has had non-consensual intercourse and the prophylaxis is required. | with a known HIV positive | person; | or |
| Initiation – Percutaneous exposure | | | |
| Patient has percutaneous exposure to blood known to be HIV pos | itive. | | |
| Initiation – Pre-exposure prophylaxis | | | |
| Re-assessment required after 3 months | | | |
| All of the following: | | | |
| 1 Applicant has an up to date knowledge of the safety issues | | | exposure prophylaxis (refer |
| to local health pathways or https://ashm.org.au/HIV/PrEP/ 2 Patient has undergone testing for HIV, syphilis and Hep B | • /· | | acks: and |
| Patient has had renal function testing (creatinine, phosphal is not contraindicated for treatment; and | | | |
| Patient has received advice regarding the reduction of risk those risks; and | of HIV and sexually transr | nitted infe | ections and how to reduce |
| 5 Patient has tested HIV negative and is not at risk of HIV se 6 Either: | roconversion; and | | |
| | | | |

6.1 All of the following:

90

- 6.1.1 Patient is male or transgender; and
- 6.1.2 Patient has sex with men; and

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST |) | Generic |
| \$ | Per | Manufacturer |

continued...

- 6.1.3 Patient is likely to have multiple episodes of condomless anal intercourse in the next 3 months; and
- 6.1.4 Any of the following:
 - 6.1.4.1 Patient has had at least one episode of condomless receptive anal intercourse with one or more casual male partners in the last 3 months; or
 - 6.1.4.2 A diagnosis of rectal chlamydia, rectal gonorrhoea, or infectious syphilis within the last 3 months; or
 - 6.1.4.3 Patient has used methamphetamine in the last three months; or
- 6.2 All of the following:
 - 6.2.1 Patient has a regular partner who has HIV infection; and
 - 6.2.2 Partner is either not on treatment or has a detectable viral load; and
 - 6.2.3 Condoms have not been consistently used.

Continuation - Pre-exposure prophylaxis

Re-assessment required after 3 months

All of the following:

- 1 Applicant has an up to date knowledge of the safety issues and is competent to prescribe pre-exposure prophylaxis (refer to local health pathways or https://ashm.org.au/HIV/PrEP/ for training materials); and
- 2 Patient has undergone testing for HIV, syphilis and Hep B if not immune in the previous two weeks; and
- 3 Patient has had renal function testing (creatinine, phosphate and urine protein/creatinine ratio) within the last 12 months and is not contraindicated for treatment; and
- 4 Patient has received advice regarding the reduction of risk of HIV and sexually transmitted infections and how to reduce those risks; and
- 5 Patient has tested HIV negative and is not at risk of HIV seroconversion; and
- 6 Either:
 - 6.1 All of the following:
 - 6.1.1 Patient is male or transgender; and
 - 6.1.2 Patient has sex with men; and
 - 6.1.3 Patient is likely to have multiple episodes of condomless anal intercourse in the next 3 months; and
 - 6.1.4 Any of the following:
 - 6.1.4.1 Patient has had at least one episode of condomless receptive anal intercourse with one or more casual male partners in the last 3 months; or
 - 6.1.4.2 A diagnosis of rectal chlamydia, rectal gonorrhoea, or infectious syphilis within the last 3 months; or
 - 6.1.4.3 Patient has used methamphetamine in the last three months; or
 - 6.2 All of the following:
 - 6.2.1 Patient has a regular partner who has HIV infection; and
 - 6.2.2 Partner is either not on treatment or has a detectable viral load; and
 - 6.2.3 Condoms have not been consistently used.

Influenza

OSELTAMIVIR - Restricted see terms below

Note: The restriction on the use of oseltamivir to hospitalised patients means that supply into the community for a new course is not permitted. Supply of a part original pack on discharge where initiated as a hospital inpatient is permitted.

- Tab 75 mg
- Powder for oral suspension 6 mg per ml

→ Restricted (RS1307)

Initiation

Either:

- 1 Only for hospitalised patient with known or suspected influenza; or
- 2 For prophylaxis of influenza in hospitalised patients as part of a DHB hospital approved infections control plan.

| Price | | | Brand or |
|---------------|--|-----|--------------|
| (ex man. excl | | | Generic |
| \$ | | Per | Manufacturer |

ZANAMIVIR

Note: The restriction on the use of zanamivir to hospitalised patients means that supply into the community for a new course is not permitted. Supply of a part original pack on discharge where initiated as a hospital inpatient is permitted.

→ Restricted (RS1369)

Initiation

Either:

- 1 Only for hospitalised patient with known or suspected influenza; or
- 2 For prophylaxis of influenza in hospitalised patients as part of a DHB hospital approved infections control plan.

Immune Modulators

INTERFERON ALFA-2A

- Inj 3 m iu prefilled syringe
- Inj 6 m iu prefilled syringe
- Inj 9 m iu prefilled syringe

(Any Inj 3 m iu prefilled syringe to be delisted 1 December 2020) (Any Inj 6 m iu prefilled syringe to be delisted 1 December 2020) (Any Inj 9 m iu prefilled syringe to be delisted 1 December 2020)

INTERFERON ALFA-2B

- Inj 18 m iu, 1.2 ml multidose pen
- Inj 30 m iu, 1.2 ml multidose pen
- Inj 60 m iu, 1.2 ml multidose pen

INTERFERON GAMMA - Restricted see terms below

- Inj 100 mcg in 0.5 ml vial
- ➡ Restricted (RS1113)

Initiation

Patient has chronic granulomatous disease and requires interferon gamma.

PEGYLATED INTERFERON ALFA-2A - Restricted see terms below

| t | Inj 180 mcg prefilled syringe500.00 | 4 | Pegasys |
|---|-------------------------------------|---|---------|
|---|-------------------------------------|---|---------|

➡ Restricted (RS1762)

Initiation – Chronic hepatitis C - genotype 1, 4, 5 or 6 infection or co-infection with HIV or genotype 2 or 3 post liver transplant

Limited to 48 weeks treatment

Any of the following:

- 1 Patient has chronic hepatitis C, genotype 1, 4, 5 or 6 infection; or
- 2 Patient has chronic hepatitis C and is co-infected with HIV; or
- 3 Patient has chronic hepatitis C genotype 2 or 3 and has received a liver transplant.

Notes: Consider stopping treatment if there is absence of a virological response (defined as at least a 2-log reduction in viral load) following 12 weeks of treatment since this is predictive of treatment failure.

Consider reducing treatment to 24 weeks if serum HCV RNA level at Week 4 is undetectable by sensitive PCR assay (less than 50IU/ml) AND Baseline serum HCV RNA is less than 400,000IU/ml.

Continuation - Chronic hepatitis C - genotype 1 infection

Gastroenterologist, infectious disease specialist or general physician

Re-assessment required after 48 weeks

All of the following:

92

1 Patient has chronic hepatitis C, genotype 1; and

| | (ex man. excl. \$ | Per | Generic Manufacturer |
|---|----------------------|----------------|-------------------------------|
| continued | | | |
| 2 Patient has had previous treatment with pegylated interferon3 Either: | and ribavirin; and | | |
| 3.1 Patient has responder relapsed; or3.2 Patient was a partial responder; and | | | |
| 4 Patient is to be treated in combination with boceprevir. | | | |
| Initiation – Chronic Hepatitis C - genotype 1 infection treatment Gastroenterologist, infectious disease specialist or general physicial <i>Limited to 48 weeks</i> treatment All of the following: | | rs prior | |
| Patient has chronic hepatitis C, genotype 1; and Patient has had previous treatment with pegylated interferon Any of the following: | and ribavirin; and | | |
| 3.1 Patient has responder relapsed; or3.2 Patient was a partial responder; or3.3 Patient received interferon treatment prior to 2004; ar | nd | | |
| 4 Patient is to be treated in combination with boceprevir. Initiation – Chronic hepatitis C - genotype 2 or 3 infection without Limited to 6 months treatment | out co-infection w | rith HIV | |
| Patient has chronic hepatitis C, genotype 2 or 3 infection. Initiation – Hepatitis B | _ | | |
| Gastroenterologist, infectious disease specialist or general physician <i>Limited to 48 weeks</i> treatment All of the following: | n | | |
| Patient has confirmed Hepatitis B infection (HBsAg positive f Patient is Hepatitis B treatment-naive; and ALT > 2 times Upper Limit of Normal; and HBV DNA < 10 log10 IU/ml; and Either: | for more than 6 mo | onths); and | |
| 5.1 HBeAg positive; or5.2 Serum HBV DNA greater than or equal to 2,000 units Stage F2 or moderate fibrosis); and | /ml and significant | fibrosis (grea | ater than or equal to Metavir |
| 6 Compensated liver disease; and 7 No continuing alcohol abuse or intravenous drug use; and 8 Not co-infected with HCV, HIV or HDV; and 9 Neither ALT nor AST > 10 times upper limit of normal; and 10 No history of hypersensitivity or contraindications to pegylate | ed interferon. | | |
| Notes: Approved dose is 180 mcg once weekly. The recommended dose of Pegylated Interferon alfa-2a is 180 mcg In patients with renal insufficiency (calculated creatinine clearance is be reduced to 135 mcg once weekly. | ess than 50ml/min | | |
| In patients with neutropaenia and thrombocytopaenia, dose should Pegylated Interferon alfa-2a is not approved for use in children. Initiation – myeloproliferative disorder or cutaneous T cell lymp | | ordance with t | he datasheet guidelines. |
| Re-assessment required after 12 months Any of the following: 1 Patient has a cutaneous T cell lymphoma*; or | | | |
| 2 All of the following: | | | |
| | | | continued. |

INFECTIONS

Brand or

Generic

Price

(ex man. excl. GST)

| Price | | | Brand or |
|--------------|---------------------|-----|--------------|
| (ex man. exc | (ex man. excl. GST) | | Generic |
| \$ | | Per | Manufacturer |

continued...

- 2.1 Patient has a myeloproliferative disorder*; and
- 2.2 Patient is intolerant of hydroxyurea; and
- 2.3 Treatment with anagrelide and busulfan is not clinically appropriate; or
- 3 Both:
 - 3.1 Patient has a myeloproliferative disorder; and
 - 3.2 Patient is pregnant, planning pregnancy or lactating.

Continuation – myeloproliferative disorder or cutaneous T cell lymphoma

Re-assessment required after 12 months

All of the following:

- 1 No evidence of disease progression; and
- 2 The treatment remains appropriate and patient is benefitting from treatment; and
- 3 Either:

94

- 3.1 Patient has a cutaneous T cell lymphoma*; or
- 3.2 Both:
 - 3.2.1 Patient has a myeloproliferative disorder*; and
 - 3.2.2 Either:
 - 3.2.2.1 Remains intolerant of hydroxyurea and treatment with anagrelide and busulfan remains clinically inappropriate; or
 - 3.2.2.2 Patient is pregnant, planning pregnancy or lactating.
- Note: Indications marked with * are unapproved indications

| | Price | | Brand or |
|--|-----------------------------|------------------------|--|
| | (ex man. excl. GST \$ | Per | Generic Manufacturer |
| | Ŷ | 1 01 | Manufacturor |
| Anticholinesterases | | | |
| DROPHONIUM CHLORIDE – Restricted see terms below | | | |
| Inj 10 mg per ml, 15 ml vial | | | |
| Inj 10 mg per ml, 1 ml ampoule → Restricted (RS1015) | | | |
| nitiation | | | |
| or the diagnosis of myasthenia gravis. | | | |
| EOSTIGMINE METILSULFATE | | | |
| Inj 2.5 mg per ml, 1 ml ampoule | | 50 | AstraZeneca |
| EOSTIGMINE METILSULFATE WITH GLYCOPYRRONIUM B | ROMIDE | | |
| Inj 2.5 mg with glycopyrronium bromide 0.5 mg per ml, 1 ml | ampoule20.90 | 10 | Max Health |
| YRIDOSTIGMINE BROMIDE | | | |
| Tab 60 mg - 1% DV Nov-19 to 2022 | 45.79 | 100 | Mestinon |
| Antirheumatoid Agents | | | |
| | | | |
| IYDROXYCHLOROQUINE – Restricted see terms below | 7.00 | 100 | Diaguanil |
| ↓ Tab 200 mg – 1% DV Sep-18 to 2021 | | 100 | Plaquenil |
| | | | |
| nitiation | | | |
| nitiation .ny of the following: | | | |
| ny of the following: 1 Rheumatoid arthritis; or | | | |
| any of the following: 1 Rheumatoid arthritis; or 2 Systemic or discoid lupus erythematosus; or | | | |
| 1 Rheumatoid arthritis; or 2 Systemic or discoid lupus erythematosus; or 3 Malaria treatment or suppression; or | unus and lister planus, and | tonoouo | |
| ny of the following: 1 Rheumatoid arthritis; or 2 Systemic or discoid lupus erythematosus; or 3 Malaria treatment or suppression; or 4 Relevant dermatological conditions (cutaneous forms of light and set of the set of t | upus and lichen planus, cu | taneous v | vasculitides and mucosal |
| Rheumatoid arthritis; or Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luulceration). | upus and lichen planus, cu | taneous v | vasculitides and mucosal |
| Rheumatoid arthritis; or Rystemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE | | taneous v 30 | |
| Rheumatoid arthritis; or Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luulceration). | | | vasculitides and mucosal Apo-Leflunomide Arava |
| Rheumatoid arthritis; or Rystemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE | 2.90 6.00 | | Apo-Leflunomide |
| Invo of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg – 1% DV Dec-20 to 2023 Tab 20 mg – 1% DV Dec-20 to 2023 | 2.90 6.00 | 30 | Apo-Leflunomide Arava |
| Any of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg - 1% DV Dec-20 to 2023 Tab 20 mg - 1% DV Dec-20 to 2023 Apo-Leflunomide Tab 10 mg to be delisted 1 December 2020) | | 30 | Apo-Leflunomide Arava Apo-Leflunomide |
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| Any of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg – 1% DV Dec-20 to 2023 Tab 20 mg – 1% DV Dec-20 to 2023 Apo-Leflunomide Tab 10 mg to be delisted 1 December 2020) PENICILLAMINE Tab 125 mg | | 30 30 100 | Apo-Leflunomide Arava Apo-Leflunomide Arava D-Penamine |
| Inv of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg – 1% DV Dec-20 to 2023 Tab 20 mg – 1% DV Dec-20 to 2023 Apo-Leflunomide Tab 10 mg to be delisted 1 December 2020) Apo-Leflunomide Tab 20 mg to be delisted 1 December 2020) PENICILLAMINE Tab 125 mg Tab 250 mg | | 30 30 | Apo-Leflunomide Arava Apo-Leflunomide Arava |
| Inv of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg – 1% DV Dec-20 to 2023 Tab 20 mg – 1% DV Dec-20 to 2023 Apo-Leflunomide Tab 10 mg to be delisted 1 December 2020) Apo-Leflunomide Tab 20 mg to be delisted 1 December 2020) PENICILLAMINE Tab 125 mg Tab 250 mg GODIUM AUROTHIOMALATE | | 30 30 100 | Apo-Leflunomide Arava Apo-Leflunomide Arava D-Penamine |
| Inv of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg – 1% DV Dec-20 to 2023 Tab 20 mg – 1% DV Dec-20 to 2023 Apo-Leflunomide Tab 10 mg to be delisted 1 December 2020) Apo-Leflunomide Tab 20 mg to be delisted 1 December 2020) PENICILLAMINE Tab 125 mg Tab 250 mg | | 30 30 100 | Apo-Leflunomide Arava Apo-Leflunomide Arava D-Penamine |
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| Inv of the following: Rheumatoid arthritis; or Systemic or discoid lupus erythematosus; or Malaria treatment or suppression; or Relevant dermatological conditions (cutaneous forms of luceration). EFLUNOMIDE Tab 10 mg - 1% DV Dec-20 to 2023 Tab 20 mg - 1% DV Dec-20 to 2023 Apo-Leflunomide Tab 10 mg to be delisted 1 December 2020) Apo-Leflunomide Tab 20 mg to be delisted 1 December 2020) PENICILLAMINE Tab 125 mg Tab 250 mg CODIUM AUROTHIOMALATE Inj 10 mg in 0.5 ml ampoule Inj 20 mg in 0.5 ml ampoule Inj 50 mg in 0.5 ml ampoule Drugs Affecting Bone Metabolism | | 30 30 100 100 | Apo-Leflunomide Arava Apo-Leflunomide Arava D-Penamine D-Penamine |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price man. excl. GS | <u>۲۱</u> | Brand or Generic |
|---|------------------------|-------------|-------------------------|
| | \$ | Per | Manufacturer |
| PAMIDRONATE DISODIUM | | | |
| Inj 3 mg per ml, 10 ml vial | 5.98 | 1 | Pamisol |
| Inj 6 mg per ml, 10 ml vial | | 1 | Pamisol |
| Inj 9 mg per ml, 10 ml vial | | 1 | Pamisol |
| RISEDRONATE SODIUM | | | |
| Tab 35 mg - 1% DV Oct-19 to 2022 | | 4 | Risedronate Sandoz |
| ZOLEDRONIC ACID | | | |
| Inj 5 mg per 100 ml, vial – 1% DV Oct-19 to 2022 | 60.00 | 100 ml | Aclasta |
| ➡ Restricted (RS1663) | | | |
| Initiation – Inherited bone fragility disorders | | | |
| Any specialist | | | |
| Patient has been diagnosed with an inherited bone fragility disorder (e.g. or | steogenesis ir | mperfecta). | |
| Initiation – Osteoporosis | - | | |
| Any specialist | | | |
| Therapy limited to 3 doses | | | |
| Both: | | | |
| 1 Any of the following: | | | |
| 1.1 History of one significant osteonorotic fracture demonstrated | vilicologically | and docume | ented hone mineral dens |

- 1.1 History of one significant osteoporotic fracture demonstrated radiologically and documented bone mineral density (BMD) greater than or equal to 2.5 standard deviations below the mean normal value in young adults (i.e. T-Score less than or equal to -2.5) (see Note); or
- 1.2 History of one significant osteoporotic fracture demonstrated radiologically, and either the patient is elderly, or densitometry scanning cannot be performed because of major logistical, technical or pathophysiological reasons. It is unlikely that this provision would apply to many patients under 75 years of age; or
- 1.3 History of two significant osteoporotic fractures demonstrated radiologically; or
- 1.4 Documented T-Score greater than or equal to -3.0 (see Note); or
- 1.5 A 10-year risk of hip fracture greater than or equal to 3%, calculated using a published risk assessment algorithm (e.g. FRAX or Garvan) which incorporates BMD measurements (see Note); or
- 1.6 Patient has had a Special Authority approval for alendronate (Underlying cause Osteoporosis) prior to 1 February 2019 or has had a Special Authority approval for raloxifene; and
- 2 The patient will not be prescribed more than 5 mg of zoledronic acid in a 12-month period.

Initiation - glucocorticosteroid therapy

Any specialist

Re-assessment required after 12 months

All of the following:

- 1 The patient is receiving systemic glucocorticosteroid therapy (greater than or equal to 5 mg per day prednisone equivalents) and has already received or is expected to receive therapy for at least three months; and
- 2 Any of the following:
 - 2.1 The patient has documented BMD greater than or equal to 1.5 standard deviations below the mean normal value in young adults (i.e. T-Score less than or equal to -1.5) (see Note); or
 - 2.2 The patient has a history of one significant osteoporotic fracture demonstrated radiologically; or
 - 2.3 The patient has had a Special Authority approval for alendronate (Underlying cause glucocorticosteroid therapy) prior to 1 February 2019 or has had a Special Authority approval for raloxifene; and
- 3 The patient will not be prescribed more than 5 mg of zoledronic acid in the 12-month approval period.

Continuation - glucocorticosteroid therapy

Any specialist

Re-assessment required after 12 months Both:

1 The patient is continuing systemic glucocorticosteriod therapy (greater than or equal to 5 mg per day prednisone

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

equivalents); and

2 The patient will not be prescribed more than 5 mg of zoledronic acid in the 12-month approval period.

Initiation – Paget's disease

Any specialist

Re-assessment required after 12 months

All of the following:

- 1 Paget's disease; and
- 2 Any of the following:
 - 2.1 Bone or articular pain; or
 - 2.2 Bone deformity; or
 - 2.3 Bone, articular or neurological complications; or
 - 2.4 Asymptomatic disease, but risk of complications; or
 - 2.5 Preparation for orthopaedic surgery; and
- 3 The patient will not be prescribed more than 5 mg of zoledronic acid in the 12-month approval period.

Continuation - Paget's disease

Any specialist

Re-assessment required after 12 months Both:

- 1 Any of the following:
 - 1.1 The patient has relapsed (based on increases in serum alkaline phosphatase); or
 - 1.2 The patient's serum alkaline phosphatase has not normalised following previous treatment with zoledronic acid; or
 - 1.3 Symptomatic disease (prescriber determined); and
- 2 The patient will not be prescribed more than 5 mg of zoledronic acid in the 12-month approval period.

Notes:

- 1 BMD (including BMD used to derive T-Score) must be measured using dual-energy x-ray absorptiometry (DXA). Quantitative ultrasound and quantitative computed tomography (QCT) are not acceptable.
- 2 Evidence suggests that patients aged 75 years and over who have a history of significant osteoporotic fracture demonstrated radiologically are very likely to have a T-Score less than or equal to -2.5 and, therefore, do not require BMD measurement for treatment with bisphosphonates.
- 3 Osteoporotic fractures are the incident events for severe (established) osteoporosis and can be defined using the WHO definitions of osteoporosis and fragility fracture. The WHO defines severe (established) osteoporosis as a T-score below -2.5 with one or more associated fragility fractures. Fragility fractures are fractures that occur as a result of mechanical forces that would not ordinarily cause fracture (minimal trauma). The WHO has quantified this as forces equivalent to a fall from a standing height or less.
- 4 A vertebral fracture is defined as a 20% or greater reduction in height of the anterior or mid portion of a vertebral body relative to the posterior height of that body, or a 20% or greater reduction in any of these heights compared to the vertebral body above or below the affected vertebral body.

Other Drugs Affecting Bone Metabolism

| DENOSUMAB – Restricted see terms below | | | |
|--|--------|---|--------|
| Inj 60 mg prefilled syringe | 326.00 | 1 | Prolia |
| → Restricted (RS1665) | | | |
| Initiation | | | |
| All of the following: | | | |
| • | | | |

1 The patient has severe, established osteoporosis; and

| Price | | Brand or | |
|-------------------|-----|--------------|--|
| (ex man. excl. GS | | Generic | |
| \$ | Per | Manufacturer | |

continued...

- 2 Either:
 - 2.1 The patient is female and postmenopausal; or
 - 2.2 The patient is male or non-binary; and
- 3 Any of the following:
 - 3.1 History of one significant osteoporotic fracture demonstrated radiologically and documented bone mineral density (BMD) greater than or equal to 2.5 standard deviations below the mean normal value in young adults (i.e. T-Score less than or equal to -2.5) (see Note); or
 - 3.2 History of one significant osteoporotic fracture demonstrated radiologically, and either the patient is elderly, or densitometry scanning cannot be performed because of major logistical, technical or pathophysiological reasons; or
 - 3.3 History of two significant osteoporotic fractures demonstrated radiologically; or
 - 3.4 Documented T-Score less than or equal to -3.0 (see Note); or
 - 3.5 A 10-year risk of hip fracture greater than or equal to 3%, calculated using a published risk assessment algorithm (e.g. FRAX or Garvan) which incorporates BMD measurements (see Note); or
 - 3.6 Patient has had a Special Authority approval for alendronate (Underlying cause Osteoporosis) prior to 1 February 2019 or has had a Special Authority approval for raloxifene; and
- 4 Zoledronic acid is contraindicated because the patient's creatinine clearance is less than 35 mL/min; and
- 5 The patient has experienced at least one symptomatic new fracture after at least 12 months' continuous therapy with a funded antiresorptive agent at adequate doses (see Notes); and
- 6 The patient must not receive concomitant treatment with any other funded antiresorptive agent for this condition or teriparatide.

Notes:

- 1 BMD (including BMD used to derive T-Score) must be measured using dual-energy x-ray absorptiometry (DXA). Quantitative ultrasound and quantitative computed tomography (QCT) are not acceptable.
- 2 Evidence suggests that patients aged 75 years and over who have a history of significant osteoporotic fracture demonstrated radiologically are very likely to have a T-Score less than or equal to -2.5 and, therefore, do not require BMD measurement for treatment with denosumab.
- 3 Osteoporotic fractures are the incident events for severe (established) osteoporosis and can be defined using the WHO definitions of osteoporosis and fragility fracture. The WHO defines severe (established) osteoporosis as a T-score below -2.5 with one or more associated fragility fractures. Fragility fractures are fractures that occur as a result of mechanical forces that would not ordinarily cause fracture (minimal trauma). The WHO has quantified this as forces equivalent to a fall from a standing height or less.
- 4 A vertebral fracture is defined as a 20% or greater reduction in height of the anterior or mid portion of a vertebral body relative to the posterior height of that body, or a 20% or greater reduction in any of these heights compared to the vertebral body above or below the affected vertebral body.
- 5 Antiresorptive agents and their adequate doses for the purposes of this Special Authority are defined as: risedronate sodium tab 35 mg once weekly; alendronate sodium tab 70 mg or tab 70 mg with cholecalciferol 5,600 iu once weekly; raloxifene hydrochloride tab 60 mg once daily. If an intolerance of a severity necessitating permanent treatment withdrawal develops during the use of one antiresorptive agent, an alternate antiresorptive agent must be trialled so that the patient achieves the minimum requirement of 12 months' continuous therapy.

| RALOXIFENE – Restricted see terms below | | |
|---|--------|--------|
| I Tab 60 mg | 28 | Evista |
| ➡ Restricted (RS1666) | | |
| Initiation | | |

Any of the following:

1 History of one significant osteoporotic fracture demonstrated radiologically and documented bone mineral density (BMD) greater than or equal to 2.5 standard deviations below the mean normal value in young adults (i.e. T-Score less than or equal to -2.5) (see Notes); or

continued...

| Price | | Brand or | |
|---------------------|-----|--------------|--|
| (ex man. excl. GST) | _ | Generic | |
| \$ | Per | Manufacturer | |

continued...

- 2 History of one significant osteoporotic fracture demonstrated radiologically, and either the patient is elderly, or densitometry scanning cannot be performed because of major logistical, technical or pathophysiological reasons. It is unlikely that this provision would apply to many patients under 75 years of age; or
- 3 History of two significant osteoporotic fractures demonstrated radiologically; or
- 4 Documented T-Score greater than or equal to -3.0 (see Notes); or
- 5 A 10-year risk of hip fracture greater than or equal to 3%, calculated using a published risk assessment algorithm (e.g. FRAX or Garvan) which incorporates BMD measurements (see Notes); or
- 6 Patient has had a Special Authority approval for zoledronic acid (Underlying cause Osteoporosis) or has had a Special Authority approval for alendronate (Underlying cause Osteoporosis) prior to 1 February 2019.

Notes:

- 1 BMD (including BMD used to derive T-Score) must be measured using dual-energy x-ray absorptiometry (DXA). Quantitative ultrasound and quantitative computed tomography (QCT) are not acceptable.
- 2 Evidence suggests that patients aged 75 years and over who have a history of significant osteoporotic fracture demonstrated radiologically are very likely to have a T-Score less than or equal to -2.5 and, therefore, do not require BMD measurement for raloxifene funding.
- 3 Osteoporotic fractures are the incident events for severe (established) osteoporosis, and can be defined using the WHO definitions of osteoporosis and fragility fracture. The WHO defines severe (established) osteoporosis as a T-score below -2.5 with one or more associated fragility fractures. Fragility fractures are fractures that occur as a result of mechanical forces that would not ordinarily cause fracture (minimal trauma). The WHO has quantified this as forces equivalent to a fall from a standing height or less.
- 4 A vertebral fracture is defined as a 20% or greater reduction in height of the anterior or mid portion of a vertebral body relative to the posterior height of that body, or a 20% or greater reduction in any of these heights compared to the vertebral body above or below the affected vertebral body.

TERIPARATIDE - Restricted see terms below

→ Restricted (RS1143)

Initiation

Limited to 18 months treatment

All of the following:

- 1 The patient has severe, established osteoporosis; and
- 2 The patient has a documented T-score less than or equal to -3.0 (see Notes); and
- 3 The patient has had two or more fractures due to minimal trauma; and
- 4 The patient has experienced at least one symptomatic new fracture after at least 12 months' continuous therapy with a funded antiresorptive agent at adequate doses (see Notes).

Notes:

- 1 The bone mineral density (BMD) measurement used to derive the T-score must be made using dual-energy x-ray absorptiometry (DXA). Quantitative ultrasound and quantitative computed tomography (QCT) are not acceptable
- 2 Antiresorptive agents and their adequate doses for the purposes of this restriction are defined as: alendronate sodium tab 70 mg or tab 70 mg with colecalciferol 5,600 iu once weekly; raloxifene hydrochloride tab 60 mg once daily; zoledronic acid 5 mg per year. If an intolerance of a severity necessitating permanent treatment withdrawal develops during the use of one antiresorptive agent, an alternate antiresorptive agent must be trialled so that the patient achieves the minimum requirement of 12 months' continuous therapy.
- 3 A vertebral fracture is defined as a 20% or greater reduction in height of the anterior or mid portion of a vertebral body relative to the posterior height of that body, or a 20% or greater reduction in any of these heights compared to the vertebral body above or below the affected vertebral body.

Enzymes

HYALURONIDASE

Inj 1,500 iu ampoule

| | | Price excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|--------------------------------|---------------------------|--|
| | | Ψ | 1 61 | Manufacturer |
| Hyperuricaemia and Antigout | | | | |
| ALLOPURINOL | | | | |
| Tab 100 mg - 1% DV Nov-20 to 2023 | | | 500 | DP-Allopurinol |
| Tab 300 mg - 1% DV Nov-20 to 2023 | | .28.57 | 500 | DP-Allopurinol |
| BENZBROMARONE - Restricted see terms below | | | | |
| Tab 50 mg Tab 100 mg | | 45.00 | 100 | Benzbromaron AL 100 |
| → Restricted (RS1489) | | .43.00 | 100 | Denzbromaron AL 100 |
| nitiation | | | | |
| Any specialist | | | | |
| All of the following: | | | | |
| 1 Patient has been diagnosed with gout; and | | | | |
| 2 Any of the following: | | | | |
| 2.1 The patient has a serum urate level greater than 0.3 | | | | |
| 600 mg/day and addition of probenecid at doses of 2.2 The patient has experienced intolerable side effects | | | | |
| and serum urate remains greater than 0.36 mmol/l c | | | | |
| maximum tolerated dose; or | lespile use of | properieciu | 11 00565 | or up to 2 y per day or |
| 2.3 Both: | | | | |
| 2.3.1 The patient has renal impairment such that p | robenecid is o | contraindicat | ed or like | lv to be ineffective and |
| serum urate remains greater than 0.36 mmol | | | | |
| 2.3.2 The patient has a rate of creatinine clearance | e greater than | or equal to 2 | 20 ml/min | ; or |
| 2.4 All of the following: | | | | |
| 2.4.1 The patient is taking azathioprine and require | es urate-lower | ing therapy; | and | |
| 2.4.2 Allopurinol is contraindicated; and | | | | |
| 2.4.3 Appropriate doses of probenecid are ineffect function; and | ive or probene | ecid cannot t | be used d | ue to reduced renal |
| 3 The patient is receiving monthly liver function tests. | | | | |
| Notes: Benzbromarone has been associated with potentially fatal he glomerular filtration rate is 30 ml/minute or less, probenecid ma batients with renal impairment is defined as treatment to the creati emains greater than 0.36 mmol/l, a gradual increase of the dose of the dose of the dose | ay not be effect nine clearance | ctive. Optima e-adjusted de | al treatme ose of allo | ent with allopurinol in opurinol then, if serum urat |
| The New Zealand Rheumatology Association has developed informat www.rheumatology.org.nz/home/resources-2/ | | | | |
| COLCHICINE | | | | |
| Tab 500 mcg – 1% DV Jan-19 to 2021 | | 9.58 | 100 | Colgout |
| EBUXOSTAT – Restricted see terms below | | | | |
| Tab 80 mg | | | 28 | Adenuric |
| Tab 120 mg | | .39.50 | 28 | Adenuric |
| → Restricted (RS1760) | | | | |

Initiation

Any specialist

Both:

1 Patient has been diagnosed with gout; and

2 Any of the following:

2.1 The patient has a serum urate level greater than 0.36 mmol/l despite treatment with allopurinol at doses of at least

continued...

| P | rice | | Brand or |
|----------|-----------|-----|--------------|
| (ex man. | excl. GST | | Generic |
| | \$ | Per | Manufacturer |

continued...

600 mg/day and addition of probenecid at doses of up to 2 g per day or maximum tolerated dose; or

- 2.2 The patient has experienced intolerable side effects from allopurinol such that treatment discontinuation is required and serum urate remains greater than 0.36 mmol/l despite use of probenecid at doses of up to 2 g per day or maximum tolerated dose; or
- 2.3 The patient has renal impairment such that probenecid is contraindicated or likely to be ineffective and serum urate remains greater than 0.36 mmol/l despite optimal treatment with allopurinol (see Note); or
- 2.4 The patient has previously had an initial Special Authority approval for benzbromarone for treatment of gout...

Note: In chronic renal insufficiency, particularly when the glomerular filtration rate is 30 ml/minute or less, probenecid may not be effective. The efficacy and safety of febuxostat have not been fully evaluated in patients with severe renal impairment (creatinine clearance less than 30 ml/minute). No dosage adjustment of febuxostat is necessary in patients with mild or moderate renal impairment. Optimal treatment with allopurinol in patients with renal impairment is defined as treatment to the creatinine clearance-adjusted dose of allopurinol then, if serum urate remains greater than 0.36 mmol/l, a gradual increase of the dose of allopurinol to 600 mg or the maximum tolerated dose.

PROBENECID

Tab 500 mg

RASBURICASE - Restricted see terms below

- Inj 1.5 mg vial
- → Restricted (RS1016)

Haematologist

Muscle Relaxants and Related Agents

ATRACURIUM BESYLATE

| P | | | | |
|---|--|--------------|----|----------------------|
| | Inj 10 mg per ml, 2.5 ml ampoule - 1% DV Jun-18 to 2021 10.0 | 00 | 5 | Tracrium |
| | Inj 10 mg per ml, 5 ml ampoule - 1% DV Jun-18 to 2021 12. | 50 | 5 | Tracrium |
| F | ACLOFEN | | | |
| - | Tab 10 mg - 1% DV Oct-18 to 20214. | 0 1 | 00 | Pacifen |
| | Oral lig 1 mg per ml | <u>1</u> 0 1 | 00 | raciicii |
| | | | 1 | Lioresal Intrathecal |
| | Inj 0.05 mg per ml, 1 ml ampoule | | • | |
| | Inj 2 mg per ml, 5 ml ampoule - 1% DV Apr-19 to 2021 | 98 | 5 | Medsurge |
| C | CLOSTRIDIUM BOTULINUM TYPE A TOXIN | | | |
| | Inj 100 u vial | 50 | 1 | Botox |
| | Inj 300 u vial | 50 | 1 | Dysport |
| | lnį 500 u vial | | 2 | Dysport |
| г | DANTROLENE | | | 7 - 1 |
| L | · · · · · · · · · · · · · · · · · · · | -0 1 | 00 | Dantrium |
| | Cap 25 mg | | 00 | Bantinann |
| | Cap 50 mg | | 00 | Dantrium |
| | Inj 20 mg vial | 00 | 6 | Dantrium IV |
| Ν | /IVACURIUM CHLORIDE | | | |
| | Inj 2 mg per ml, 5 ml ampoule | 92 | 5 | Mivacron |
| | Inj 2 mg per ml, 10 ml ampoule67. | 17 | 5 | Mivacron |
| ~ | DRPHENADRINE CITRATE | | | |
| C | | | 00 | Norflex |
| | Tab 100 mg - 1% DV Jun-18 to 202118. | 04 I | 00 | Nornex |
| F | ANCURONIUM BROMIDE | | | |
| | Inj 2 mg per ml, 2 ml ampoule | | | |
| F | OCURONIUM BROMIDE | | | |
| | Inj 10 mg per ml, 5 ml ampoule – 1% DV Aug-20 to 2022 | 14 1 | 10 | Hameln |
| | | 17 | | |
| 5 | UXAMETHONIUM CHLORIDE | | | |
| | Inj 50 mg per ml, 2 ml ampoule78. | 00 5 | 50 | AstraZeneca |
| | | | | |

Products with Hospital Supply Status (HSS) are in **bold**

Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|---|---|-----------------------|-------------------------------------|
| ECURONIUM BROMIDE Inj 10 mg vial | | | |
| Reversers of Neuromuscular Blockade | | | |
| UGAMMADEX – Restricted see terms below Inj 100 mg per ml, 2 ml vial Inj 100 mg per ml, 5 ml vial | 1,200.00 3,000.00 | 10 10 | Bridion Bridion |
| ny of the following: Patient requires reversal of profound neuromuscular bloc undertaken using rocuronium (i.e. suxamethonium is con 2 Severe neuromuscular degenerative disease where the u Patient has an unexpectedly difficult airway that cannot be neuromuscular blockade: or | ntraindicated or undesirabluse of neuromuscular block | e); or kade is ree | quired; or |
| 4 The duration of the patient's surgery is unexpectedly sho 5 Neostigmine or a neostigmine/anticholinergic combinatio disease, morbid obesity or COPD); or | | ample the | patient has ischaemic hea |

6 Patient has a partial residual block after conventional reversal.

Non-Steroidal Anti-Inflammatory Drugs

CELECOXIB

| CELECONID | | | |
|---|-----|-----|-------------------|
| Cap 100 mg3 | .63 | 60 | Celecoxib Pfizer |
| Cap 200 mg2 | .30 | 30 | Celecoxib Pfizer |
| DICLOFENAC SODIUM | | | |
| Tab EC 25 mg - 1% DV Oct-18 to 20211 | .23 | 50 | Diclofenac Sandoz |
| Tab 50 mg dispersible1 | .50 | 20 | Voltaren D |
| Tab EC 50 mg - 1% DV Oct-18 to 20211 | .23 | 50 | Diclofenac Sandoz |
| Tab long-acting 75 mg - 1% DV Oct-18 to 2021 | .80 | 500 | Apo-Diclo SR |
| Tab long-acting 100 mg - 1% DV Oct-18 to 2021 | .15 | 500 | Apo-Diclo SR |
| Inj 25 mg per ml, 3 ml ampoule13 | .20 | 5 | Voltaren |
| Suppos 12.5 mg2 | .04 | 10 | Voltaren |
| Suppos 25 mg2 | .44 | 10 | Voltaren |
| Suppos 50 mg4 | .22 | 10 | Voltaren |
| Suppos 100 mg7 | .00 | 10 | Voltaren |
| | | | |

ETORICOXIB - Restricted see terms below

- I Tab 60 mg
- ↓ Tab 90 mg
- ↓ Tab 120 mg
- → Restricted (RS1290)

Initiation

102

For in-vivo investigation of allergy only.

| BUPROFEN | \$.11.71 | Per | Manufacturer |
|---|------------------|--------------|----------------------------|
| | .11.71 | | |
| | .11.71 | | |
| Tab 200 mg | | 1,000 | Relieve |
| → Tab 400 mg - Restricted: For continuation only | | | |
| → Tab 600 mg - Restricted: For continuation only | F 00 | 20 | Ibunratan CD DNM |
| Tab long-acting 800 mg – 1% DV Apr-20 to 2021 Oral lig 20 mg per ml – 1% DV May-19 to 2021 | | 30 200 ml | Ibuprofen SR BNM Ethics |
| Ini 5 mg per ml, 2 ml ampoule | 1.00 | 200 111 | Luncs |
| Inj 10 mg per ml, 2 ml vial | | | |
| | | | |
| | | | |
| Cap 25 mg Cap 50 mg | | | |
| Cap long-acting 75 mg | | | |
| Inj 1 mg vial | | | |
| Suppos 100 mg | | | |
| KETOPROFEN | | | |
| Cap long-acting 200 mg | 12 07 | 28 | Oruvail SR |
| | . 12.07 | 20 | |
| MEFENAMIC ACID - Restricted: For continuation only → Cap 250 mg | | | |
| | | | |
| VAPROXEN | 00.00 | 500 | N - flam 050 |
| Tab 250 mg - 1% DV Dec-18 to 2021 | | 500 | Noflam 250 Noflam 500 |
| Tab 500 mg – 1% DV Dec-18 to 2021 Tab long-acting 750 mg – 1% DV Oct-18 to 2021 | | 250 28 | Naprosyn SR 750 |
| Tab long-acting 1 g – 1% DV Oct-18 to 2021 | | 28 | Naprosyn SR 1000 |
| | 0.2 1 | 20 | Naprosyn on 1000 |
| PARECOXIB Inj 40 mg vial | 00.00 | 10 | Durpootot |
| | 100.00 | 10 | Dynastat |
| SULINDAC | | | |
| Tab 100 mg | | | |
| Tab 200 mg | | | |
| TENOXICAM | | | |
| Tab 20 mg – 1% DV Oct-19 to 2022 | | 100 | Tilcotil |
| Inj 20 mg vial | 9.95 | 1 | AFT |
| Topical Products for Joint and Muscular Pain | | | |
| Topical Products for Joint and Muscular Pall | | | |
| CAPSAICIN – Restricted see terms below | | | |
| Crm 0.025% | 9.95 | 45 g | Zostrix |
| → Restricted (RS1309) | | | |

Initiation

Patient has osteoarthritis that is not responsive to paracetamol and oral non-steroidal anti-inflammatories are contraindicated.

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|--|-----------------------------------|--------------|---------------------------------------|
| Agents for Parkinsonism and Related Disorders | 1 | | |
| Agents for Essential Tremor, Chorea and Relate | d Disorders | | |
| RILUZOLE - Restricted see terms below I Tab 50 mg - 1% DV Aug-18 to 2021 | duration of 5 years or lea | | Rilutek |
| 5.3 The patient is able to swallow. Continuation Re-assessment required after 18 months All of the following: The patient has not undergone a tracheostomy; and The patient has not experienced respiratory failure; and Any of the following: The patient is ambulatory; or The patient is able to use upper limbs; or The patient is able to swallow. | | | |
| TETRABENAZINE Tab 25 mg – 1% DV Oct-19 to 2022 | 91.10 | 112 | Motetis |
| Anticholinergics | | | |
| BENZATROPINE MESYLATE Tab 2 mg Inj 1 mg per ml, 2 ml ampoule – 1% DV Dec-20 to 2023 (Cogentin Inj 1 mg per ml, 2 ml ampoule to be delisted 1 Decemb PROCYCLIDINE HYDROCHLORIDE Tab 5 mg | 95.00 | 60 5 | Benztrop Cogentin Phebra |
| Dopamine Agonists and Related Agents | | | |
| AMANTADINE HYDROCHLORIDE Cap 100 mg APOMORPHINE HYDROCHLORIDE Inj 10 mg per ml, 2 ml ampoule – 1% DV Jan-20 to 2023 Inj 10 mg per ml, 5 ml ampoule – 1% DV Feb-20 to 2023 BROMOCRIPTINE Tab 2.5 mg | | 60 5 5 | Symmetrel Movapo Movapo |

t Item restricted (see → above); t Item restricted (see → below)

104

e.g. Brand indicates brand example only. It is not a contracted product.

NERVOUS SYSTEM

| | Price | | Brand or |
|---|--------------------------|-------------|---|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer |
| ENTACAPONE | * | | |
| Tab 200 mg – 1% DV Sep-18 to 2021 | | 100 | Entapone |
| EVODOPA WITH BENSERAZIDE | | | |
| Tab dispersible 50 mg with benserazide 12.5 mg | 13 25 | 100 | Madopar Rapid |
| Cap 50 mg with benserazide 12.5 mg | | 100 | Madopar 62.5 |
| Cap 100 mg with benserazide 25 mg | | 100 | Madopar 125 |
| Cap long-acting 100 mg with benserazide 25 mg | | 100 | Madopar HBS |
| Cap 200 mg with benserazide 50 mg | | 100 | Madopar 250 |
| EVODOPA WITH CARBIDOPA | | | |
| Tab 100 mg with carbidopa 25 mg | | 100 | Sinemet |
| Tab long-acting 100 mg with carbipoda 25 mg | | | |
| Tab long-acting 200 mg with carbidopa 50 mg | | 100 | Sinemet CR |
| Tab 250 mg with carbidopa 25 mg | | 100 | Sinemet |
| PRAMIPEXOLE HYDROCHLORIDE | | | |
| Tab 0.25 mg - 1% DV Oct-19 to 2022 | 6.12 | 100 | Ramipex |
| Tab 1 mg - 1% DV Oct-19 to 2022 | 20.73 | 100 | Ramipex |
| ROPINIROLE HYDROCHLORIDE | | | |
| Tab 0.25 mg - 1% DV Mar-20 to 2022 | 2.85 | 84 | Ropin |
| Tab 1 mg - 1% DV Mar-20 to 2022 | 3.95 | 84 | Ropin |
| Tab 2 mg - 1% DV Mar-20 to 2022 | 5.48 | 84 | Ropin |
| Tab 5 mg – 1% DV Mar-20 to 2022 | | 84 | Ropin |
| ELEGILINE HYDROCHLORIDE | | | |
| Tab 5 mg | | | |
| OLCAPONE | | | |
| Tab 100 mg | | 100 | Tasmar |
| | | | |
| Anaesthetics | | | |
| General Anaesthetics | | | |
| DESFLURANE | | | |
| Soln for inhalation 100%, 240 ml bottle | | 6 | Suprane |
| DEXMEDETOMIDINE | , | | |
| Inj 100 mcg per ml, 2 ml vial | | 5 | Precedex |
| TOMIDATE | | U U | |
| Inj 2 mg per ml, 10 ml ampoule | | | |
| | | | |
| SOFLURANE Soln for inhalation 100%, 250 ml bottle | 1 020 00 | 6 | Aerrane |
| | 1,020.00 | 0 | Aenane |
| | | | |
| ETAMINE | 405.00 | - | Diama d |
| ETAMINE Inj 1 mg per ml, 100 ml bag <i>–</i> 1% DV Feb-20 to 2022 | | 5 | Biomed |
| ETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 | | 5 | Biomed |
| ETAMINE Inj 1 mg per ml, 100 ml bag <i></i> 1% DV Feb-20 to 2022 | 70.00 31.50 | | Biomed Ketalar |
| XETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 Inj 100 mg per ml, 2 ml vial – 1% DV Jan-19 to 2021 | | 5 | Biomed |
| KETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 Inj 100 mg per ml, 2 ml vial – 1% DV Jan-19 to 2021 | 70.00 31.50 | 5 | Biomed Ketalar |
| KETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 Inj 100 mg per ml, 2 ml vial – 1% DV Jan-19 to 2021 METHOHEXITAL SODIUM Inj 10 mg per ml, 50 ml vial | 70.00 31.50 | 5 | Biomed Ketalar |
| KETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 Inj 100 mg per ml, 2 ml vial – 1% DV Jan-19 to 2021 METHOHEXITAL SODIUM Inj 10 mg per ml, 50 ml vial PROPOFOL | 70.00 31.50 155.60 | 5 5 | Biomed Ketalar Ketamine-Claris |
| KETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 Inj 100 mg per ml, 2 ml vial – 1% DV Jan-19 to 2021 METHOHEXITAL SODIUM Inj 10 mg per ml, 50 ml vial PROPOFOL Inj 10 mg per ml, 20 ml ampoule – 10% DV Dec-19 to 2022 | | 5 5 5 | Biomed Ketalar Ketamine-Claris Fresofol 1% MCT/LCT |
| XETAMINE Inj 1 mg per ml, 100 ml bag – 1% DV Feb-20 to 2022 Inj 10 mg per ml, 10 ml syringe – 1% DV Feb-20 to 2022 Inj 100 mg per ml, 2 ml vial – 1% DV Jan-19 to 2021 METHOHEXITAL SODIUM Inj 10 mg per ml, 50 ml vial PROPOFOL | | 5 5 | Biomed Ketalar Ketamine-Claris |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|--------|-------------------------------------|
| SEVOFLURANE Soln for inhalation 100%, 250 ml bottle THIOPENTAL [THIOPENTONE] SODIUM Inj 500 mg ampoule | | 6 | Baxter |
| Local Anaesthetics | | | |
| ARTICAINE HYDROCHLORIDE Inj 1% | | | |
| ARTICAINE HYDROCHLORIDE WITH ADRENALINE Inj 4% with adrenaline 1:100,000, 1.7 ml dental cartridge Inj 4% with adrenaline 1:100,000, 2.2 ml dental cartridge Inj 4% with adrenaline 1:200,000, 1.7 ml dental cartridge Inj 4% with adrenaline 1:200,000, 2.2 ml dental cartridge | | | |
| BENZOCAINE Gel 20% | | | |
| BENZOCAINE WITH TETRACAINE HYDROCHLORIDE Gel 18% with tetracaine hydrochloride 2% | | | e.g. ZAP Topical Anaesthetic Gel |
| BUPIVACAINE HYDROCHLORIDE Inj 5 mg per ml, 4 ml ampoule – 1% DV Oct-20 to 2023 Inj 2.5 mg per ml, 20 ml ampoule | | 5 | Marcain Isobaric |
| Inj 2.5 mg per ml, 20 ml ampoule sterile pack – 1% DV Aug-20 Inj 5 mg per ml, 10 ml ampoule sterile pack – 1% DV Aug-20 Inj 5 mg per ml, 20 ml ampoule | | 5 5 | Marcain Marcain |
| Inj 5 mg per ml, 20 ml ampoule sterile pack – 1% DV Aug-20 t Inj 1.25 mg per ml, 100 ml bag Inj 1.25 mg per ml, 200 ml bag | to 2023 16.56 | 5 | Marcain |
| Inj 2.5 mg per ml, 100 ml bag – 1% DV Oct-20 to 2023 Inj 2.5 mg per ml, 200 ml bag Inj 1.25 mg per ml, 500 ml bag | 150.00 | 5 | Marcain |
| BUPIVACAINE HYDROCHLORIDE WITH ADRENALINE Inj 2.5 mg per ml with adrenaline 1:400,000, 20 ml vial – 1% D | V Aug 10 | | |
| to 2022 | | 5 | Marcain with Adrenaline |
| Inj 5 mg per ml with adrenaline 1:200,000, 20 ml vial – 1% DV to 2022 | | 5 | Marcain with Adrenaline |
| BUPIVACAINE HYDROCHLORIDE WITH FENTANYL Inj 0.625 mg with fentanyl 2 mcg per ml, 100 ml bag Inj 0.625 mg with fentanyl 2 mcg per ml, 200 ml bag – 1% DV | Δnr-20 | | |
| to 2022 Inj 1.25 mg with fentanyl 2 mcg per ml, 100 ml syringe | | 5 | Biomed |
| Inj 1.25 mg with fentanyl 2 mcg per ml, 100 ml bag – 1% DV N to 2022 Inj 1.25 mg with fentanyl 2 mcg per ml, 200 ml bag – 1% DV N | | 5 | Bupafen |
| to 2022 Inj 1.25 mg with fentanyl 2 mcg per ml, 50 ml syringe | | 5 | Bupafen |
| Inj 1.25 mg with fentanyl 2 mcg per ml, 15 ml syringe Inj 1.25 mg with fentanyl 2 mcg per ml, 20 ml syringe | | 5 5 | Biomed Biomed |

e.g. Brand indicates brand example only. It is not a contracted product.

NERVOUS SYSTEM

| | Price | T \ | Brand or | |
|--|-------------------------|------------|-------------------------|--|
| | (ex man. excl. GS \$ | I) Per | Generic Manufacturer | |
| BUPIVACAINE HYDROCHLORIDE WITH GLUCOSE | | | | |
| Inj 0.5% with glucose 8%, 4 ml ampoule | | 5 | Marcain Heavy | |
| OCAINE HYDROCHLORIDE | | | | |
| Paste 5% | | | | |
| Soln 15%, 2 ml syringe | | | | |
| Soln 4%, 2 ml syringe | 25.46 | 1 | Biomed | |
| COCAINE HYDROCHLORIDE WITH ADRENALINE | | | | |
| Paste 15% with adrenaline 0.06% | | | | |
| Paste 25% with adrenaline 0.06% | | | | |
| ETHYL CHLORIDE | | | | |
| Spray 100% | | | | |
| IDOCAINE [LIGNOCAINE] | | | | |
| Crm 4% | 5.40 | 5 g | LMX4 | |
| | 27.00 | 30 g | LMX4 | |
| IDOCAINE [LIGNOCAINE] HYDROCHLORIDE | | | | |
| Gel 2% - 1% DV Nov-18 to 2021 | 4.87 | 20 g | Orion | |
| Soln 4% | | | | |
| Spray 10% – 1% DV Jul-19 to 2022 | | 50 ml | Xylocaine | |
| Oral (gel) soln 2% | | 200 ml | Mucosoothe | |
| Inj 1%, 20 ml ampoule, sterile pack Inj 2%, 20 ml ampoule, sterile pack | | | | |
| Inj 1%, 5 ml ampoule | 8 75 | 25 | Lidocaine-Claris | |
| Inj 1%, 20 ml vial – 1% DV Jul-19 to 2022 | | 5 | Lidocaine-Claris | |
| Inj 2%, 5 ml ampoule - 1% DV Nov-19 to 2022 | | 25 | Lidocaine-Claris | |
| Inj 2%, 20 ml vial - 1% DV Jul-19 to 2022 | | 5 | Lidocaine-Claris | |
| Gel 2%, 11 ml urethral syringe - 1% DV Apr-20 to 2022 | | 10 | Instillagel Lido | |
| IDOCAINE [LIGNOCAINE] HYDROCHLORIDE WITH ADRENALINE | | | | |
| Inj 1% with adrenaline 1:100,000, 5 ml ampoule - 1% DV Nov-19 | | | | |
| to 2022 | | 10 | Xylocaine | |
| Inj 1% with adrenaline 1:200,000, 20 ml vial | 50.00 | 5 | Xylocaine | |
| Inj 2% with adrenaline 1:80,000, 1.7 ml dental cartridge | | | | |
| Inj 2% with adrenaline 1:80,000, 1.8 ml dental cartridge | | | | |
| Inj 2% with adrenaline 1:80,000, 2.2 ml dental cartridge | 60.00 | F | Vulaasina | |
| Inj 2% with adrenaline 1:200,000, 20 ml vial | | 5 | Xylocaine | |
| IDOCAINE [LIGNOCAINE] HYDROCHLORIDE WITH ADRENALINE | | = HYDROC | HLORIDE | |
| Soln 4% with adrenaline 0.1% and tetracaine hydrochloride 0.5%, s | | | Teniesies | |
| syringe | | 1 | Topicaine | |
| IDOCAINE [LIGNOCAINE] HYDROCHLORIDE WITH CHLORHEXIDI | | 10 | Dfiner | |
| Gel 2% with chlorhexidine 0.05%, 10 ml urethral syringe | | 10 | Pfizer | |
| IDOCAINE [LIGNOCAINE] HYDROCHLORIDE WITH PHENYLEPHR | INE HYDROCHLC | RIDE | | |
| Nasal spray 5% with phenylephrine hydrochloride 0.5% | | | | |
| IDOCAINE [LIGNOCAINE] WITH PRILOCAINE | | • | | |
| Crm 2.5% with prilocaine 2.5% | | 30 g | EMLA | |
| Patch 25 mcg with prilocaine 25 mcg | | 20 | EMLA | |
| Crm 2.5% with prilocaine 2.5%, 5 g | 45.00 | 5 | EMLA | |
| | | 50 | 0 | |
| Inj 3%, 1.8 ml dental cartridge | | 50 | Scandonest 3% | |
| Inj 3%, 2.2 ml dental cartridge | 43.00 | 50 | Scandonest 3% | |

| | Price (ex man. excl. GST) | | Brand or Generic |
|---|------------------------------|-----|---------------------|
| | (ex man. excl. GST) \$ | Per | Manufacturer |
| PRILOCAINE HYDROCHLORIDE Inj 0.5%, 50 ml vial Inj 2%, 5 ml ampoule | | 5 | Citanest |
| PRILOCAINE HYDROCHLORIDE WITH FELYPRESSIN Inj 3% with felypressin 0.03 iu per ml, 1.8 ml dental cartridge Inj 3% with felypressin 0.03 iu per ml, 2.2 ml dental cartridge | | | |
| ROPIVACAINE HYDROCHLORIDE | | | |
| Inj 2 mg per ml, 10 ml ampoule – 1% DV Nov-20 to 2023 | | 5 | Ropivacaine Kabi |
| Inj 2 mg per ml, 20 ml ampoule – 1% DV Nov-20 to 2023 | | 5 | Ropivacaine Kabi |
| Inj 2 mg per ml, 100 ml bag – 1% DV Nov-20 to 2023 | | 5 | Ropivacaine Kabi |
| Inj 2 mg per ml, 200 ml bag – 1% DV Nov-20 to 2023 | 40.95 | 5 | Ropivacaine Kabi |
| Inj 7.5 mg per ml, 10 ml ampoule - 1% DV Nov-20 to 2023 | 10.40 | 5 | Ropivacaine Kabi |
| Inj 7.5 mg per ml, 20 ml ampoule - 1% DV Nov-20 to 2023 | | 5 | Ropivacaine Kabi |
| Inj 10 mg per ml, 10 ml ampoule - 1% DV Nov-20 to 2023 | | 5 | Ropivacaine Kabi |
| Inj 10 mg per ml, 20 ml ampoule - 1% DV Nov-20 to 2023 | | 5 | Ropivacaine Kabi |
| ROPIVACAINE HYDROCHLORIDE WITH FENTANYL | | | |
| Inj 2 mg with fentanyl 2 mcg per ml, 100 ml bag | | 5 | Naropin |
| Inj 2 mg with fentanyl 2 mcg per ml, 200 ml bag | | 5 | Naropin |
| FETRACAINE FAMETHOCAINE HYDROCHLORIDE | | | |

TETRACAINE [AMETHOCAINE] HYDROCHLORIDE

Gel 4%

Analgesics

Non-Opioid Analgesics

| Tab dispersible 300 mg - 1% DV Oct-19 to 2022 | 4.50 | 100 | Ethics Aspirin |
|---|-------|------|----------------|
| CAPSAICIN – Restricted see terms below | 40.50 | 45 | Zerticulup |
| ↓ Crm 0.075% → Restricted (RS1145) | 12.50 | 45 g | Zostrix HP |

Initiation

For post-herpetic neuralgia or diabetic peripheral neuropathy.

METHOXYFLURANE - Restricted see terms below

Soln for inhalation 99.9%, 3 ml bottle

➡ Restricted (RS1292)

Initiation

Both:

1 Patient is undergoing a painful procedure with an expected duration of less than one hour; and

2 Only to be used under supervision by a medical practitioner or nurse who is trained in the use of methoxyflurane.

NEFOPAM HYDROCHLORIDE

Tab 30 mg

| | Price | | Brand or |
|---|-------------------------|---------------|-----------------------------|
| | (ex man. excl. GS \$ | Per | Generic Manufacturer |
| PARACETAMOL – Some items restricted see terms below | | | |
| Tab soluble 500 mg | | | |
| Tab 500 mg | | | |
| Oral liq 120 mg per 5 ml - 20% DV Nov-20 to 2023 | 5.45 | 1,000 ml | Paracare |
| Oral liq 250 mg per 5 ml – 20% DV Nov-20 to 2023 | 6.25 | 1,000 ml | Paracare Double Strength |
| Inj 10 mg per ml, 100 ml vial – 1% DV Nov-20 to 2023 | 8.90 | 10 | Paracetamol Kabi |
| Suppos 25 mg - 1% DV Nov-19 to 2022 | | 20 | Biomed |
| Suppos 50 mg - 1% DV Nov-19 to 2022 | | 20 | Biomed |
| Suppos 125 mg - 1% DV Nov-18 to 2021 | | 10 | Gacet |
| Suppos 250 mg - 1% DV Nov-18 to 2021 | | 10 | Gacet |
| Suppos 500 mg - 1% DV Feb-19 to 2021 | | 50 | Gacet |
| ➡ Restricted (RS1146) | | | |
| Initiation | | | |
| Intravenous paracetamol is only to be used where other routes ar | e unavailable or imprac | tical, or whe | e there is reduced |
| absorption. The need for IV paracetamol must be re-assessed ev | very 24 hours. | | |
| SUCROSE | | | |
| Oral liq 25% - 1% DV Feb-20 to 2022 | | 25 ml | Biomed |
| I Oral lig 66.7% (preservative free) | | | |
| → Restricted (RS1763) | | | |
| initiation | | | |
| For use in neonatal patients only. | | | |
| | | | |
| Onicid Analyseice | | | |
| Opioid Analgesics | | | |
| ALFENTANIL | | | |
| Inj 0.5 mg per ml, 2 ml ampoule – 1% DV Nov-20 to 2023 CODEINE PHOSPHATE | | 10 | Hameln |

| CODEINE PHOSPHATE Tab 15 mg - 1% DV Nov-20 to 2023 | 100 | PSM |
|---|-----|------------------|
| Tab 30 mg – 1% DV Nov-20 to 2023 | 100 | PSM |
| Tab 60 mg - 1% DV Nov-20 to 202314.25 | 100 | PSM |
| DIHYDROCODEINE TARTRATE | | |
| Tab long-acting 60 mg – 1% DV Oct-19 to 2022 | 60 | DHC Continus |
| FENTANYL | | |
| Inj 10 mcg per ml, 10 ml syringe | | |
| Inj 50 mcg per ml, 2 ml ampoule - 1% DV Nov-18 to 2021 | 10 | Boucher and Muir |
| Inj 10 mcg per ml, 50 ml bag210.00 | 10 | Biomed |
| Inj 10 mcg per ml, 50 ml syringe 165.00 | 10 | Biomed |
| Inj 50 mcg per ml, 10 ml ampoule - 1% DV Nov-18 to 2021 | 10 | Boucher and Muir |
| Inj 10 mcg per ml, 100 ml bag - 1% DV Nov-19 to 2022 | 5 | Biomed |
| Inj 20 mcg per ml, 50 ml syringe - 1% DV Oct-18 to 2021 | 1 | Biomed |
| Inj 20 mcg per ml, 100 ml bag | | |
| Patch 12.5 mcg per hour2.95 | 5 | Fentanyl Sandoz |
| Patch 25 mcg per hour | 5 | Fentanyl Sandoz |
| Patch 50 mcg per hour6.65 | 5 | Fentanyl Sandoz |
| Patch 75 mcg per hour9.25 | 5 | Fentanyl Sandoz |
| Patch 100 mcg per hour 11.40 | 5 | Fentanyl Sandoz |

| | Price | | Brand or |
|--|-------------------|---------|-----------------------|
| | (ex man. excl. GS | | Generic |
| | \$ | Per | Manufacturer |
| METHADONE HYDROCHLORIDE | | | |
| Tab 5 mg - 1% DV Sep-19 to 2022 | | 10 | Methatabs |
| Oral liq 2 mg per ml – 1% DV Oct-18 to 2021 | | 200 ml | Biodone |
| Oral liq 5 mg per ml - 1% DV Oct-18 to 2021 | | 200 ml | Biodone Forte |
| Oral liq 10 mg per ml – 1% DV Oct-18 to 2021 | | 200 ml | Biodone Extra Forte |
| Inj 10 mg per ml, 1 ml vial | 61.00 | 10 | AFT |
| MORPHINE HYDROCHLORIDE | | | |
| Oral liq 1 mg per ml – 1% DV Dec-18 to 2021 | 9.28 | 200 ml | RA-Morph |
| Oral liq 2 mg per ml - 1% DV Dec-18 to 2021 | 16.24 | 200 ml | RA-Morph |
| Oral liq 5 mg per ml – 1% DV Dec-18 to 2021 | 19.44 | 200 ml | RA-Morph |
| Oral liq 10 mg per ml – 1% DV Dec-18 to 2021 | 27.74 | 200 ml | RA-Morph |
| MORPHINE SULPHATE | | | |
| Tab long-acting 10 mg | 1.93 | 10 | Arrow-Morphine LA |
| Tab immediate-release 10 mg - 1% DV Nov-20 to 2023 | | 10 | Sevredol |
| Tab immediate-release 20 mg - 1% DV Nov-20 to 2023 | | 10 | Sevredol |
| Tab long-acting 30 mg | | 10 | Arrow-Morphine LA |
| Tab long-acting 60 mg | | 10 | Arrow-Morphine LA |
| Cap long-acting 10 mg - 1% DV Jan-20 to 2022 | | 10 | m-Eslon |
| Cap long-acting 30 mg - 1% DV Jan-20 to 2022 | | 10 | m-Eslon |
| Cap long-acting 60 mg - 1% DV Jan-20 to 2022 | 6.12 | 10 | m-Eslon |
| Cap long-acting 100 mg - 1% DV Jan-20 to 2022 | | 10 | m-Eslon |
| Inj 1 mg per ml, 100 ml bag - 1% DV Nov-20 to 2023 | | 5 | Biomed |
| Inj 1 mg per ml, 10 ml syringe - 1% DV Nov-20 to 2023 | 24.50 | 5 | Biomed |
| Inj 1 mg per ml, 50 ml syringe - 1% DV Nov-20 to 2023 | | 5 | Biomed |
| Inj 1 mg per ml, 2 ml syringe | | | |
| Inj 2 mg per ml, 30 ml syringe | | 10 | Biomed |
| Inj 5 mg per ml, 1 ml ampoule | 6.27 | 5 | DBL Morphine Sulphate |
| Inj 10 mg per ml, 1 ml ampoule | 4.47 | 5 | DBL Morphine Sulphate |
| Inj 10 mg per ml, 100 mg cassette | | | |
| Inj 10 mg per ml, 100 ml bag | | | |
| Inj 15 mg per ml, 1 ml ampoule | 4.76 | 5 | DBL Morphine Sulphate |
| Inj 30 mg per ml, 1 ml ampoule | 6.19 | 5 | DBL Morphine Sulphate |
| Inj 200 mcg in 0.4 ml syringe | | | |
| Inj 300 mcg in 0.3 ml syringe | | | |
| (Arrow-Morphine LA Tab long-acting 10 mg to be delisted 1 October 20 | 20) | | |
| MORPHINE TARTRATE | | | |
| Inj 80 mg per ml, 1.5 ml ampoule | | 5 | DBL Morphine Tartrate |
| (DBL Morphine Tartrate Inj 80 mg per ml, 1.5 ml ampoule to be delisted | | 20) | |
| OXYCODONE HYDROCHLORIDE | , | , | |
| Tab controlled-release 5 mg – 1% DV May-19 to 2021 | 2 15 | 20 | Oxycodone Sandoz |
| Tab controlled-release 10 mg – 1% DV May-19 to 2021 | | 20 | Oxycodone Sandoz |
| Tab controlled-release 20 mg - 1% DV May-19 to 2021 | | 20 | Oxycodone Sandoz |
| Tab controlled-release 40 mg - 1% DV May-19 to 2021 | | 20 | Oxycodone Sandoz |
| Tab controlled-release 80 mg - 1% DV May-19 to 2021 | | 20 | Oxycodone Sandoz |
| Cap immediate-release 5 mg – 1% DV Sep-18 to 2021 | | 20 | OxyNorm |
| Cap immediate-release 10 mg - 1% DV Sep-18 to 2021 | | 20 | OxyNorm |
| Cap immediate-release 20 mg – 1% DV Sep-18 to 2021 | | 20 | OxyNorm |
| Oral lig 5 mg per 5 ml | | 250 ml | OxyNorm |
| Inj 1 mg per ml, 100 ml bag | | 200 111 | expression |
| Inj 10 mg per ml, 1 ml ampoule – 1% DV Sep-18 to 2021 | 7 28 | 5 | OxyNorm |
| Inj 10 mg per ml, 2 ml ampoule – 1% DV Sep-18 to 2021 | | 5 | OxyNorm |
| Inj 50 mg per ml, 1 ml ampoule – 1% DV Sep-18 to 2021 | | 5 | OxyNorm |
| | | | |

t Item restricted (see \rightarrow above); **f** Item restricted (see \rightarrow below)

e.g. Brand indicates brand example only. It is not a contracted product.

110

| | Price . excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|-----------------------------|-------|-------------------------------------|
| PARACETAMOL WITH CODEINE | | | |
| Tab paracetamol 500 mg with codeine phosphate 8 mg | 18.21 | 1,000 | Paracetamol + Codeine (Relieve) |
| PETHIDINE HYDROCHLORIDE | | | |
| Tab 50 mg - 1% DV Sep-18 to 2021 | 4.46 | 10 | PSM |
| Inj 5 mg per ml, 10 ml syringe | | | |
| Inj 5 mg per ml, 100 ml bag | | | |
| Inj 10 mg per ml, 100 ml bag | | | |
| Inj 10 mg per ml, 50 ml syringe | | | |
| Inj 50 mg per ml, 1 ml ampoule | 4.98 | 5 | DBL Pethidine Hydrochloride |
| Inj 50 mg per ml, 2 ml ampoule | 5.12 | 5 | DBL Pethidine Hydrochloride |
| REMIFENTANIL | | | |
| Inj 1 mg vial – 1% DV Oct-20 to 2023 | 13.95 | 5 | Remifentanil-AFT |
| Inj 2 mg vial - 1% DV Oct-20 to 2023 | 19.95 | 5 | Remifentanil-AFT |
| TRAMADOL HYDROCHLORIDE | | | |
| Tab sustained-release 100 mg - 1% DV Nov-20 to 2023 | 1.52 | 20 | Tramal SR 100 |
| Tab sustained-release 150 mg - 1% DV Nov-20 to 2023 | | 20 | Tramal SR 150 |
| Tab sustained-release 200 mg - 1% DV Nov-20 to 2023 | 2.75 | 20 | Tramal SR 200 |
| Cap 50 mg | 2.25 | 100 | Arrow-Tramadol |
| Oral soln 10 mg per ml | | | |
| Inj 10 mg per ml, 100 ml bag | | | |
| Inj 50 mg per ml, 1 ml ampoule - 1% DV Oct-20 to 2023 | | 5 | Tramal 50 |
| Inj 50 mg per ml, 2 ml ampoule – 1% DV Oct-20 to 2023 | 3.83 | 5 | Tramal 100 |
| Antidepressants | | | |
| Cyclic and Related Agents | | | |
| AMITRIPTYLINE | | | |

| AMITRIPTYLINE | | |
|---|-----|---------------------|
| Tab 10 mg 1.96 | 100 | Arrow-Amitriptyline |
| Tab 25 mg 1.52 | 100 | Arrow-Amitriptyline |
| Tab 50 mg2.51 | 100 | Arrow-Amitriptyline |
| CLOMIPRAMINE HYDROCHLORIDE | | |
| Tab 10 mg - 1% DV Oct-18 to 2021 | 100 | Apo-Clomipramine |
| Tab 25 mg – 1% DV Oct-18 to 2021 | 100 | Apo-Clomipramine |
| DOSULEPIN [DOTHIEPIN] HYDROCHLORIDE - Restricted: For continuation only | | |
| → Cap 25 mg | 50 | Dosulepin Mylan |
| DOXEPIN HYDROCHLORIDE – Restricted: For continuation only | | |
| → Cap 10 mg | | |
| → Cap 25 mg | | |
| ➡ Cap 50 mg | | |
| IMIPRAMINE HYDROCHLORIDE | | |
| Tab 10 mg | 50 | Tofranil |
| 6.58 | 60 | Tofranil |
| Tab 25 mg | 50 | Tofranil |
| MAPROTILINE HYDROCHLORIDE | | |
| Tab 25 mg | | |
| Tab 75 mg | | |
| | | |

| | | Price excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|-------|---------------------------|------------|--|
| MIANSERIN HYDROCHLORIDE - Restricted: For continuation on | nly | | | |
| → Tab 30 mg | | | | |
| VORTRIPTYLINE HYDROCHLORIDE Tab 10 mg – 1% DV Oct-19 to 2022 | | 0.44 | 100 | Normross |
| Tab 25 mg – 1% DV Oct-19 to 2022 Tab 25 mg – 1% DV Oct-19 to 2022 | | 2.44 5.98 | 100 180 | Norpress Norpress |
| | | | 100 | Norpress |
| Monoamine-Oxidase Inhibitors - Non-Selective | | | | |
| PHENELZINE SULPHATE | | | | |
| Tab 15 mg | | | | |
| TRANYLCYPROMINE SULPHATE | | | | |
| Tab 10 mg | | | | |
| Monoamine-Oxidase Type A Inhibitors | | | | |
| MOCLOBEMIDE | | 0.40 | | • · |
| Tab 150 mg – 1% DV Apr-19 to 2021 | | | 60 | Aurorix |
| Tab 300 mg – 1% DV Apr-19 to 2021 | | 9.80 | 60 | Aurorix |
| Other Antidepressants | | | | |
| MIRTAZAPINE | | | | |
| Tab 30 mg - 1% DV Oct-18 to 2021 | | | 30 | Apo-Mirtazapine |
| Tab 45 mg - 1% DV Oct-18 to 2021 | | 3.48 | 30 | Apo-Mirtazapine |
| VENLAFAXINE | | | | |
| Cap 37.5 mg | | | 84 | Enlafax XR Enlafax XR |
| Cap 75 mg Cap 150 mg | | | 84 84 | Enlafax XR |
| | | | 04 | |
| Selective Serotonin Reuptake Inhibitors | | | | |
| | | 1 50 | 0.4 | DOM Ottologram |
| Tab 20 mg - 1% DV Sep-18 to 2021 | | 1.32 | 84 | PSM Citalopram |
| ESCITALOPRAM Tab 10 mg | | 4 4 4 | 28 | Essitaloprom Apotox |
| Tab 20 mg | | | 20 28 | Escitalopram-Apotex Escitalopram-Apotex |
| FLUOXETINE HYDROCHLORIDE | | | 20 | Loonaloprant/ipotox |
| Tab dispersible 20 mg, scored | | 9.93 | 30 | Arrow-Fluoxetine |
| | | 1.98 | | Fluox |
| Cap 20 mg | | 7.49 | 90 | Arrow-Fluoxetine |
| | | 2.91 | 84 | Fluox |
| PAROXETINE | | 0.01 | 00 | Louomine |
| Tab 20 mg – 1% DV Mar-20 to 2022 | | 3.01 | 90 | Loxamine |
| SERTRALINE | | 0.00 | 00 | Catuana |
| Tab 50 mg – 1% DV Mar-20 to 2022 Tab 100 mg – 1% DV Mar-20 to 2022 | | | 30 30 | Setrona Setrona |
| 5 | ····· | 1.01 | 30 | |
| Antiepilepsy Drugs | | | | |
| Agents for the Control of Status Epilepticus | | | | |
| • • • • | | | | |
| CLONAZEPAM Inj 1 mg per ml, 1 ml ampoule | | | | |

| | Price (ex man. excl. GS1 |) | Brand or Generic |
|---|-----------------------------|--------|---------------------|
| | \$ | Per | Manufacturer |
| DIAZEPAM | | | |
| Inj 5 mg per ml, 2 ml ampoule | | 5 | Hospira |
| Rectal tubes 5 mg | | 5 | Stesolid |
| Rectal tubes 10 mg | | 5 | Stesolid |
| LORAZEPAM | | | |
| Inj 2 mg vial | | | |
| Inj 4 mg per ml, 1 ml vial | | | |
| PARALDEHYDE | | | |
| Inj 5 ml ampoule | | | |
| PHENYTOIN SODIUM | | | |
| Inj 50 mg per ml, 2 ml ampoule | 88.63 | 5 | Hospira |
| Inj 50 mg per ml, 5 ml ampoule | | 5 | Hospira |
| | | Ŭ | |
| Control of Epilepsy | | | |
| CARBAMAZEPINE | | | |
| Tab 200 mg | | 100 | Tegretol |
| Tab long-acting 200 mg | | 100 | Tegretol CR |
| Tab 400 mg | | 100 | Tegretol |
| Tab long-acting 400 mg | | 100 | Tegretol CR |
| Oral liq 20 mg per ml | | 250 ml | Tegretol |
| CLOBAZAM | | | |
| Tab 10 mg | | | |
| CLONAZEPAM | | | |
| Oral drops 2.5 mg per ml | | | |
| ETHOSUXIMIDE | | | |
| Cap 250 mg | | 100 | Zarontin |
| Oral lig 50 mg per ml | | 200 ml | Zarontin |
| GABAPENTIN | | | |
| Note: Gabapentin not to be given in combination with pregabalin | | | |
| Cap 100 mg – 1% DV Aug-18 to 2021 | 2.65 | 100 | Apo-Gabapentin |
| Cap 300 mg – 1% DV Aug-18 to 2021 | | 100 | Apo-Gabapentin |
| Cap 400 mg - 1% DV Aug-18 to 2021 | 5.64 | 100 | Apo-Gabapentin |
| LACOSAMIDE - Restricted see terms below | | | |
| ↓ Tab 50 mg | | 14 | Vimpat |
| ↓ Tab 100 mg | | 14 | Vimpat |
| - | 200.24 | 56 | Vimpat |
| ↓ Tab 150 mg | 75.10 | 14 | Vimpat |
| _ | 300.40 | 56 | Vimpat |
| Tab 200 mg | | 56 | Vimpat |
| Inj 10 mg per ml, 20 ml vial | | | |
| → Restricted (RS1151) | | | |
| Initiation | | | |
| Re-assessment required after 15 months | | | |

Both:

- 1 Patient has partial-onset epilepsy; and
- 2 Seizures are not adequately controlled by, or patient has experienced unacceptable side effects from, optimal treatment

continued...

NERVOUS SYSTEM

| | Price (ex man. excl. GS \$ | ST) Per | Brand or Generic Manufacturer |
|--|----------------------------------|---------------|-------------------------------------|
| ontinued | | | |
| with all of the following: sodium valproate, topiramate, level phenytoin sodium (see Note). | - | | |
| ote: "Optimal treatment" is defined as treatment which is indicate | | | |
| oses for the patient's age, weight and other features affecting the | | | good evidence of |
| ompliance. Women of childbearing age are not required to have a | a trial of sodium valpro | ate. | |
| continuation atient has demonstrated a significant and sustained improvement | in anizura rata ar any | ority and/or | ruality of life compared w |
| hat prior to starting lacosamide treatment (see Note). | III Seizure rate of Sev | enty anu/or o | quality of the compared w |
| lote: As a guideline, clinical trials have referred to a notional 50% | reduction in seizure fi | equency as | an indicator of success w |
| nticonvulsant therapy and have assessed quality of life from the p | | equency as | |
| AMOTRIGINE | | | |
| Tab dispersible 2 mg | 6 74 | 30 | Lamictal |
| Tab dispersible 5 mg | | 56 | Arrow-Lamotrigine |
| | 9.64 | 30 | Lamictal |
| Tab dispersible 25 mg - 5% DV Oct-19 to 2022 | •••• | 56 | Logem |
| Tab dispersible 50 mg - 5% DV Oct-19 to 2022 | | 56 | Logem |
| Tab dispersible 100 mg - 5% DV Oct-19 to 2022 | | 56 | Logem |
| Arrow-Lamotrigine Tab dispersible 5 mg to be delisted 1 October 2 | | 50 | Logeni |
| | -020) | | |
| EVETIRACETAM | 4.00 | <u></u> | French |
| Tab 250 mg – 1% DV Aug-19 to 2022 Tab 500 mg – 1% DV Aug-19 to 2022 | | 60 60 | Everet Everet |
| | | 60 60 | |
| Tab 750 mg - 1% DV Aug-19 to 2022 Tab 1,000 mg - 1% DV Aug-19 to 2022 | | 60 60 | Everet Everet |
| | | 60 200 ml | Levetiracetam-AFT |
| Oral liq 100 mg per ml Inj 100 mg per ml, 5 ml vial – 1% DV Oct-19 to 2022 | | 300 ml 10 | Levelinacetam-AFT |
| | | 10 | Levelinacelain-AFT |
| HENOBARBITONE | | | |
| Tab 15 mg - 1% DV Oct-18 to 2021 | | 500 | PSM |
| Tab 30 mg - 1% DV Oct-18 to 2021 | | 500 | PSM |
| HENYTOIN | | | |
| Tab 50 mg | | | |
| HENYTOIN SODIUM | | | |
| Cap 30 mg | | | |
| Cap 100 mg | | | |
| Oral liq 6 mg per ml | | | |
| REGABALIN | | | |
| Note: Pregabalin not to be given in combination with gabapen | tin | | |
| Cap 25 mg - 1% DV Jul-18 to 2021 | | 56 | Pregabalin Pfizer |
| Cap 75 mg - 1% DV Jul-18 to 2021 | 2.65 | 56 | Pregabalin Pfizer |
| Cap 150 mg - 1% DV Jul-18 to 2021 | 4.01 | 56 | Pregabalin Pfizer |
| Cap 300 mg - 1% DV Jul-18 to 2021 | 7.38 | 56 | Pregabalin Pfizer |
| RIMIDONE | | | |
| Tab 250 mg | | | |
| ODIUM VALPROATE | | | |
| Tab 100 mg | | | |
| Tab EC 200 mg | | | |
| | | | |
| | | | |
| Tab EC 500 mg Oral lig 40 mg per ml | | | |

e.g. Brand indicates brand example only. It is not a contracted product.

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| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|---|-----------------------------------|----------|-------------------------------------|
| STIRIPENTOL – Restricted see terms below Cap 250 mg | 509.29 | 60 | Diacomit |
| Powder for oral liq 250 mg sachet | | 60 | Diacomit |

→ Restricted (RS1152) Initiation

Paediatric neurologist

Re-assessment required after 6 months

Both:

- 1 Patient has confirmed diagnosis of Dravet syndrome; and
- 2 Seizures have been inadequately controlled by appropriate courses of sodium valproate, clobazam and at least two of the following: topiramate. levetiracetam. ketogenic diet.

Continuation

Paediatric neurologist

Patient continues to benefit from treatment as measured by reduced seizure frequency from baseline.

TOPIRAMATE

| Tab 25 mg | 11.07 | 60 | Arrow-Topiramate |
|--------------------|--------|----|--------------------|
| | 26.04 | | Topamax |
| | 11.07 | | Topiramate Actavis |
| Tab 50 mg | | 60 | Arrow-Topiramate |
| ů – | 44.26 | | Topamax |
| | 18.81 | | Topiramate Actavis |
| Tab 100 mg | | 60 | Arrow-Topiramate |
| | 75.25 | | Topamax |
| | 31.99 | | Topiramate Actavis |
| Tab 200 mg | 55.19 | 60 | Arrow-Topiramate |
| ů – | 129.85 | | Topamax |
| | 55.19 | | Topiramate Actavis |
| Cap sprinkle 15 mg | 20.84 | 60 | Topamax |
| Cap sprinkle 25 mg | | 60 | Topamax |
| | | | • |

VIGABATRIN - Restricted see terms below

Tab 500 mg

→ Restricted (RS1739)

Initiation

Re-assessment required after 15 months

Both:

1 Either:

1.1 Patient has infantile spasms; or

- 1.2 Both:
 - 1.2.1 Patient has epilepsy; and
 - 1.2.2 Fither:
 - 1.2.2.1 Seizures are not adequately controlled with optimal treatment with other antiepilepsy agents; or
 - 1.2.2.2 Seizures are controlled adequately but the patient has experienced unacceptable side effects from optimal treatment with other antiepilepsy agents; and

2 Either:

- 2.1 Patient is, or will be, receiving regular automated visual field testing (ideally before starting therapy and on a 6-monthly basis thereafter); or
- 2.2 It is impractical or impossible (due to comorbid conditions, or health system capacity constraints) to monitor the patient's visual fields.

| | Price | | Brand or |
|-----|---------------|-----|--------------|
| (ex | man. excl. GS | | Generic |
| | \$ | Per | Manufacturer |

continued...

Notes: "Optimal treatment with other antiepilepsy agents" is defined as treatment with other antiepilepsy agents which are indicated and clinically appropriate for the patient, given in adequate doses for the patient's age, weight, and other features affecting the pharmacokinetics of the drug with good evidence of compliance.

Vigabatrin is associated with a risk of irreversible visual field defects, which may be asymptomatic in the early stages. Continuation

Both:

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- 1 The patient has demonstrated a significant and sustained improvement in seizure rate or severity and or quality of life; and
- 2 Either:
 - 2.1 Patient is receiving regular automated visual field testing (ideally every 6 months) on an ongoing basis for duration of treatment with vigabatrin; or
 - 2.2 It is impractical or impossible (due to comorbid conditions, or health system capacity constraints) to monitor the patient's visual fields.

Notes: As a guideline, clinical trials have referred to a notional 50% reduction in seizure frequency as an indicator of success with anticonvulsant therapy and have assessed quality of life from the patient's perspective.

Vigabatrin is associated with a risk of irreversible visual field defects, which may be asymptomatic in the early stages.

Antimigraine Preparations

Acute Migraine Treatment

| • | | |
|--|------------|--------------------|
| DIHYDROERGOTAMINE MESYLATE | | |
| Inj 1 mg per ml, 1 ml ampoule | | |
| METOCLOPRAMIDE HYDROCHLORIDE WITH PARACETAMOL | | |
| Tab 5 mg with paracetamol 500 mg | | |
| RIZATRIPTAN | | |
| Tab orodispersible 10 mg – 1% DV Oct-20 to 2023 | 30 | Rizamelt |
| SUMATRIPTAN | | |
| Tab 50 mg – 1% DV Oct-19 to 2022 | 100 | Apo-Sumatriptan |
| Tab 100 mg - 1% DV Oct-19 to 2022 | 100 | Apo-Sumatriptan |
| Inj 12 mg per ml, 0.5 ml prefilled pen – 1% DV Sep-20 to 2022 | 2 | Clustran |
| 11 12 mg per mi, 0.5 mi premied per - 176 DV Sep-20 to 2022 | 2 | Imigran |
| (Clustran Inj 12 mg per ml, 0.5 ml prefilled pen to be delisted 1 September 2020) | | inngran |
| | | |
| Prophylaxis of Migraine | | |
| PIZOTIFEN | | |
| Tab 500 mcg | 100 | Sandomigran |
| | 100 | Gandonngran |
| Antinausea and Vertigo Agents | | |
| | | |
| APREPITANT – Restricted see terms below | | |
| Cap 2 × 80 mg and 1 × 125 mg – 1% DV Jul-18 to 2021 | 3 | Emend Tri-Pack |
| → Restricted (RS1154) | | |
| Initiation | harany fa | r the treatment of |
| Patient is undergoing highly emetogenic chemotherapy and/or anthracycline-based chemot | inerapy io | r the treatment of |
| malignancy. | | |
| BETAHISTINE DIHYDROCHLORIDE | | |
| Tab 16 mg - 1% DV Nov-20 to 2023 | 84 | Vergo 16 |
| CYCLIZINE HYDROCHLORIDE | | |
| Tab 50 mg – 1% DV Jan-19 to 2021 0.55 | 10 | Nausicalm |
| | | |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|--|----------------------|-------------------------------------|
| CYCLIZINE LACTATE | | | |
| Inj 50 mg per ml, 1 ml ampoule | 14.95 | 5 | Nausicalm |
| DOMPERIDONE | | | |
| Tab 10 mg - 1% DV Mar-19 to 2021 | 2.25 | 100 | Pharmacy Health |
| DROPERIDOL | | | |
| Inj 2.5 mg per ml, 1 ml ampoule - 1% DV May-20 to 2022 | | 10 | Droleptan |
| GRANISETRON | | | - |
| Inj 1 mg per ml, 3 ml ampoule | 0.40 | 1 | Deva |
| HYOSCINE HYDROBROMIDE | | | |
| Inj 400 mcg per ml, 1 ml ampoule | | 5 | Hospira |
| Fatch 1.5 mg | | 2 | Scopoderm TTS |
| ➡ Restricted (RS1155) | | | |
| Initiation | | | |
| Any of the following: | | | |
| Control of intractable nausea, vomiting, or inability to swallow where the patient cannot tolerate or does not adequately resp Control of clozapine-induced hypersalivation where trials of a ineffective; or For treatment of post-operative nausea and vomiting where c ineffective, are not tolerated or are contraindicated. | oond to oral anti-nausea t least two other alternat | agents; ive treat | or ments have proven |
| | 0000 | | |
| (Hospira Inj 400 mcg per ml, 1 ml ampoule to be delisted 1 Septemb | er 2020) | | |
| METOCLOPRAMIDE HYDROCHLORIDE | 4.00 | 400 | |
| Tab 10 mg – 1% DV Oct-20 to 2023 | 1.30 | 100 | Metoclopramide Actavis 10 |
| Oral lig 5 mg per 5 ml | | | ACIAVIS TO |
| Inj 5 mg per ml, 2 ml ampoule – 1% DV Jan-20 to 2022 | 9.50 | 10 | Pfizer |
| ONDANSETRON | | | |
| Tab 4 mg – 1% DV Apr-20 to 2022 | | 50 | Onrex |
| Tab dispersible 4 mg - 1% DV Oct-20 to 2023 | | 10 | Ondansetron |
| | | | ODT-DRLA |
| Tab 8 mg - 1% DV Apr-20 to 2022 | | 50 | Onrex |
| Tab dispersible 8 mg - 1% DV Oct-20 to 2023 | 1.13 | 10 | Ondansetron |
| Inj 2 mg per ml, 2 ml ampoule | 1 50 | 5 | ODT-DRLA Ondansetron-Claris |
| Inj 2 mg per ml, 4 ml ampoule | | 5 | Ondansetron Kabi |
| PROCHLORPERAZINE | | Ũ | ondanoonon nabi |
| Tab buccal 3 mg | | | |
| Tab 5 mg | 6.35 | 250 | Nausafix |
| Inj 12.5 mg per ml, 1 ml ampoule | | 200 | |
| Suppos 25 mg | | | |
| TROPISETRON | | | |
| Inj 1 mg per ml, 2 ml ampoule – 1% DV Sep-18 to 2021 | 8.95 | 1 | Tropisetron-AFT |
| Inj 1 mg per ml, 5 ml ampoule | | 1 | Tropisetron-AFT |
| | | | |

| | rice excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|--------------------------|-----------|-------------------------------------|
| Antipsychotic Agents | | | |
| General | | | |
| AMISULPRIDE | | | |
| Tab 100 mg - 1% DV Nov-19 to 2022 | 5.15 | 30 | Sulprix |
| Tab 200 mg - 1% DV Nov-19 to 2022 | | 60 | Sulprix |
| Tab 400 mg – 1% DV Feb-20 to 2022 Oral liq 100 mg per ml | 29.78 | 60 | Sulprix |
| ARIPIPRAZOLE | | | |
| Tab 5 mg - 1% DV Aug-18 to 2021 | 17.50 | 30 | Aripiprazole Sandoz |
| Tab 10 mg - 1% DV Aug-18 to 2021 | | 30 | Aripiprazole Sandoz |
| Tab 15 mg - 1% DV Aug-18 to 2021 | | 30 | Aripiprazole Sandoz |
| Tab 20 mg - 1% DV Aug-18 to 2021 | | 30 | Aripiprazole Sandoz |
| Tab 30 mg – 1% DV Aug-18 to 2021 | 17.50 | 30 | Aripiprazole Sandoz |
| CHLORPROMAZINE HYDROCHLORIDE | | | |
| Tab 10 mg - 1% DV Jan-20 to 2022 | | 100 | Largactil |
| Tab 25 mg – 1% DV Jan-20 to 2022 | | 100 | Largactil |
| Tab 100 mg - 1% DV Jan-20 to 2022 | 36.73 | 100 | Largactil |
| Oral liq 10 mg per ml | | | |
| Oral liq 20 mg per ml | ~~ ~~ | | |
| Inj 25 mg per ml, 2 ml ampoule – 1% DV Jan-20 to 2022 | 30.79 | 10 | Largactil |
| CLOZAPINE | | | |
| Tab 25 mg | | 50 | Clopine |
| | 13.37 | 100 | Clopine |
| | 5.69 | 50 | Clozaril |
| Tab 50 mg | 11.36 | 100 50 | Clozaril |
| Tab 50 mg | 8.07 17.33 | 50 100 | Clopine Clopine |
| Tab 100 mg | | 50 | Clopine |
| Tab 100 mg | 34.65 | 100 | Clopine |
| | 14.73 | 50 | Clozaril |
| | 29.45 | 100 | Clozaril |
| Tab 200 mg | | 50 | Clopine |
| · | 69.30 | 100 | Clopine |
| Oral liq 50 mg per ml | | 100 ml | Clopine |
| HALOPERIDOL | | | - |
| Tab 500 mcg – 1% DV Oct-19 to 2022 | 6.23 | 100 | Serenace |
| Tab 1.5 mg – 1% DV Oct-19 to 2022 | | 100 | Serenace |
| Tab 5 mg – 1% DV Oct-19 to 2022 | | 100 | Serenace |
| Oral liq 2 mg per ml - 1% DV Oct-19 to 2022 | | 100 ml | Serenace |
| Inj 5 mg per ml, 1ml ampoule - 1% DV Oct-19 to 2022 | 21.55 | 10 | Serenace |
| LEVOMEPROMAZINE | | | |
| Tab 25 mg – 1% DV Sep-19 to 2022 | 16.10 | 100 | Nozinan |
| Tab 100 mg - 1% DV Sep-19 to 2022 | | 100 | Nozinan |
| LEVOMEPROMAZINE HYDROCHLORIDE | | | |
| Inj 25 mg per ml, 1 ml ampoule – 1% DV Apr-20 to 2022 | 33.50 | 10 | Nozinan |

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|--|-----------------------------------|----------|-------------------------------------|
| LITHIUM CARBONATE | | | |
| Tab long-acting 400 mg | | | |
| Tab 250 mg | | 500 | Lithicarb FC |
| Cap 250 mg | | 100 | Douglas |
| (Lithicarb FC Tab 250 mg to be delisted 1 November 2020) | | | 0 |
| OLANZAPINE | | | |
| Tab 2.5 mg – 1% DV Nov-20 to 2023 | 1 35 | 28 | Zypine |
| Tab 5 mg - 1% DV Nov-20 to 2023 | | 28 | Zypine |
| Tab orodispersible 5 mg - 1% DV Nov-20 to 2023 | | 28 | Zypine ODT |
| Tab 10 mg - 1% DV Nov-20 to 2023 | | 28 | Zypine |
| Tab orodispersible 10 mg - 1% DV Nov-20 to 2023 | | 28 | Zypine ODT |
| Inj 10 mg vial | 2.00 | 20 | Lypine OD1 |
| , , | | | |
| PERICYAZINE | | | |
| Tab 2.5 mg | | | |
| Tab 10 mg | | | |
| QUETIAPINE | | | |
| Tab 25 mg - 1% DV Nov-20 to 2023 | 2.15 | 90 | Quetapel |
| Tab 100 mg - 1% DV Nov-20 to 2023 | | 90 | Quetapel |
| Tab 200 mg - 1% DV Nov-20 to 2023 | | 90 | Quetapel |
| Tab 300 mg - 1% DV Nov-20 to 2023 | | 90 | Quetapel |
| RISPERIDONE | | | anompo. |
| | 1 00 | 60 | Actovia |
| Tab 0.5 mg | | 60 | Actavis |
| Tab 1 mg | | 60 | Actavis |
| Tab 2 mg | | 60 | Actavis |
| Tab 3 mg | | 60 | Actavis |
| Tab 4 mg | | 60 | Actavis |
| Oral liq 1 mg per ml – 1% DV Nov-20 to 2023 | 8.90 | 30 ml | Risperon |
| ZIPRASIDONE | | | |
| Cap 20 mg - 1% DV Dec-18 to 2021 | | 60 | Zusdone |
| Cap 40 mg - 1% DV Sep-18 to 2021 | 24.70 | 60 | Zusdone |
| Cap 60 mg - 1% DV Sep-18 to 2021 | | 60 | Zusdone |
| Cap 80 mg - 1% DV Sep-18 to 2021 | | 60 | Zusdone |
| ZUCLOPENTHIXOL ACETATE | | | |
| Inj 50 mg per ml, 1 ml ampoule | | | |
| Inj 50 mg per ml, 2 ml ampoule | | | |
| | | | |
| | 04.45 | 100 | 0 |
| Tab 10 mg | | 100 | Clopixol |
| Depot Injections | | | |
| FLUPENTHIXOL DECANOATE | | | |
| Inj 20 mg per ml, 1 ml ampoule | 12 1/ | 5 | Fluanxol |
| Inj 20 mg per ml, 2 ml ampoule | | 5 | Fluanxol |
| Inj 100 mg per ml, 1 ml ampoule | | 5 | Fluanxol |
| | | 0 | i lualikui |
| HALOPERIDOL DECANOATE | | | |
| Inj 50 mg per ml, 1 ml ampoule | | 5 | Haldol |
| Inj 100 mg per ml, 1 ml ampoule | 55.90 | 5 | Haldol Concentrate |
| OLANZAPINE - Restricted see terms on the next page | | | |
| Inj 210 mg vial – 1% DV Oct-18 to 2021 | | 1 | Zyprexa Relprevv |
| ↓ Inj 300 mg vial - 1% DV Oct-18 to 2021 | | 1 | Zyprexa Relprevv |
| Inj 000 mg vial − 1% DV Oct-18 to 2021 | | 1 | Zyprexa Relprevv |
| | | | -1hiova Heihiott |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| Price | | | Brand or |
|--------------|---------|-----|--------------|
| (ex man. exc | I. GST) | _ | Generic |
| \$ | | Per | Manufacturer |

➡ Restricted (RS1379)

Initiation

Re-assessment required after 12 months

Either:

- 1 The patient has had an initial Special Authority approval for risperidone depot injection or paliperidone depot injection; or
- 2 All of the following:
 - 2.1 The patient has schizophrenia; and
 - 2.2 The patient has tried but failed to comply with treatment using oral atypical antipsychotic agents; and
 - 2.3 The patient has been admitted to hospital or treated in respite care, or intensive outpatient or home-based treatment for 30 days or more in the last 12 months.

Continuation

Re-assessment required after 12 months

The initiation of olanzapine depot injection has been associated with fewer days of intensive intervention than was the case during a corresponding period of time prior to the initiation of an atypical antipsychotic depot injection.

PALIPERIDONE - Restricted see terms below

| t | Inj 25 mg syringe | 1 | Invega Sustenna |
|---|---------------------|-------|-----------------|
| t | Inj 50 mg syringe | 1 | Invega Sustenna |
| t | lnj 75 mg syringe | 1 | Invega Sustenna |
| | Inj 100 mg syringe | 1 | Invega Sustenna |
| Í | Inj 150 mg syringe | 1 | Invega Sustenna |
| | Destricted (DC1001) | - | |

➡ Restricted (RS1381)

Initiation

Re-assessment required after 12 months Fither:

- Either:
 - 1 The patient has had an initial Special Authority approval for risperidone depot injection or olanzapine depot injection; or
 - 2 All of the following:
 - 2.1 The patient has schizophrenia or other psychotic disorder; and
 - 2.2 The patient has tried but failed to comply with treatment using oral atypical antipsychotic agents; and
 - 2.3 The patient has been admitted to hospital or treated in respite care, or intensive outpatient or home-based treatment for 30 days or more in the last 12 months.

Continuation

Re-assessment required after 12 months

The initiation of paliperidone depot injection has been associated with fewer days of intensive intervention than was the case during a corresponding period of time prior to the initiation of an atypical antipsychotic depot injection.

PIPOTHIAZINE PALMITATE - Restricted: For continuation only

- ➡ Inj 50 mg per ml, 1 ml ampoule
- ➡ Inj 50 mg per ml, 2 ml ampoule

RISPERIDONE - Restricted see terms below

| t | Inj 25 mg vial | 98 1 | Risperdal Consta |
|---|------------------|------|------------------|
| t | Inj 37.5 mg vial | 71 1 | Risperdal Consta |
| t | Inj 50 mg vial | 56 1 | Risperdal Consta |
| | | | |

➡ Restricted (RS1380)

Initiation

Re-assessment required after 12 months

Either:

- 1 The patient has had an initial Special Authority approval for paliperidone depot injection or olanzapine depot injection; or
- 2 All of the following:
 - 2.1 The patient has schizophrenia or other psychotic disorder; and

| | ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|---|---------------|----------------------|---------|------------|-------------------------------------|
| ontinued | | | | | |
| 2.2 The patient has tried but failed to comply with treatme2.3 The patient has been admitted to hospital or treated in treatment for 30 days or more in the last 12 months. | | | | | |
| continuation | | | | | |
| Re-assessment required after 12 months he initiation of risperidone depot injection has been associated with uring a corresponding period of time prior to the initiation of an atyp UCLOPENTHIXOL DECANOATE | bical antipsy | chotic | depot i | | |
| Inj 200 mg per ml, 1 ml ampoule Inj 500 mg per ml, 1 ml ampoule | | . 19.8 | 0 | 5 | Clopixol e.g. Clopixol Conc |
| Anxiolytics | | | | | |
| USPIRONE HYDROCHLORIDE | | | | | |
| Tab 5 mg - 1% DV Sep-18 to 2021 | | | | 100 | Orion |
| Tab 10 mg - 1% DV Sep-18 to 2021 | | .13.1 | 6 | 100 | Orion |
| | | | | 400 | B |
| Tab 500 mcg – 1% DV Jun-18 to 2021 | | | | 100 100 | Paxam Paxam |
| Tab 2 mg – 1% DV Jun-18 to 2021 | | . 10.7 | 0 | 100 | Faxalli |
| | | 15.0 | - | 500 | Arrow Diazonom |
| Tab 2 mg Tab 5 mg | | | | 500 500 | Arrow-Diazepam Arrow-Diazepam |
| C C | | . 10.1 | 0 | 500 | Allow-Diazepaili |
| DRAZEPAM | | 0.7 | 0 | 250 | Ativan |
| Tab 1 mg - 1% DV Sep-18 to 2021 Tab 2.5 mg - 1% DV Sep-18 to 2021 | | | | 100 | Ativan |
| XAZEPAM | | . 12.0 | 0 | 100 | Auvan |
| Tab 10 mg | | 61 | 7 | 100 | Ox-Pam |
| Tab 15 mg | | | | 100 | Ox-Pam |
| č | | | • | | |
| Multiple Sclerosis Treatments | | | | | |
| Cap 120 mg | | 520 0 | 0 | 14 | Tecfidera |
| Cap 240 mg | | | | 56 | Tecfidera |
| ▶ Restricted (RS1504) | _ , | | • | 00 | loondord |
| itiation | | | | | |
| Inly for use in patients with approval by the Multiple Sclerosis Treat onsidered by MSTAC at its regular meetings and approved subject ut in Section B of the Pharmaceutical Schedule). | | | | | |
| INGOLIMOD – Restricted see terms below | 0.0 | | • | 00 | Cilonus |
| Cap 0.5 mg | 2,2 | 200.0 | U | 28 | Gilenya |
| ▶ Restricted (RS1433) itiation | | | | | |
| inly for use in patients with approval by the Multiple Sclerosis Treat | ment Asses | smen | t Comm | nittee (M | STAC), Applications will b |
| onsidered by MSTAC at its regular meetings and approved subject ut in Section B of the Pharmaceutical Schedule). | | | | | |
| | | | | | |
| ATALIZUMAB - Restricted see terms on the next page | | | | | |

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST | | Generic |
| \$ | Per | Manufacturer |

→ Restricted (RS1447)

Initiation

Only for use in patients with approval by the Multiple Sclerosis Treatment Assessment Committee (MSTAC). Applications will be considered by MSTAC at its regular meetings and approved subject to eligibility according to the Entry and Stopping criteria (set out in Section B of the Pharmaceutical Schedule).

| OCRELIZUMAB – Restricted see terms below ↓ Inj 30 mg per ml, 10 ml vial → Restricted (RS1711) | .9,346.00 | 1 | Ocrevus |
|--|-----------|-----|---------|
| Initiation Only for use in patients with approval by the Multiple Sclerosis Treatment As: considered by MSTAC at its regular meetings and approved subject to eligibi out in Section B of the Pharmaceutical Schedule). | | · · | / 11 |
| TERIFLUNOMIDE – Restricted see terms below ↓ Tab 14 mg | .1,582.62 | 28 | Aubagio |

Initiation

Only for use in patients with approval by the Multiple Sclerosis Treatment Assessment Committee (MSTAC). Applications will be considered by MSTAC at its regular meetings and approved subject to eligibility according to the Entry and Stopping criteria (set out in Section B of the Pharmaceutical Schedule).

Other Multiple Sclerosis Treatments

→ Restricted (RS1434)

Initiation

Only for use in patients with approval by the Multiple Sclerosis Treatment Assessment Committee (MSTAC). Applications will be considered by MSTAC at its regular meetings and approved subject to eligibility according to the Entry and Stopping criteria (set out in Section B of the Pharmaceutical Schedule).

| GLATIRAMER ACETATE – Restricted see terms above t Inj 40 mg prefilled syringe | 2,275.00 | 12 | Copaxone |
|---|----------|----|------------|
| INTERFERON BETA-1-ALPHA – Restricted see terms above Inj 6 million iu in 0.5 ml pen injector | 1,170.00 | 4 | Avonex Pen |
| t Inj 6 million iu in 0.5 ml syringe | | 4 | Avonex |
| INTERFERON BETA-1-BETA - Restricted see terms above | | | |

1 Inj 8 million iu per ml, 1 ml vial

Sedatives and Hypnotics

CHLORAL HYDRATE

Oral liq 100 mg per ml Oral liq 200 mg per ml LORMETAZEPAM – **Restricted:** For continuation only

🛏 Tab 1 mg

MELATONIN - Restricted see terms on the next page

I Tab 3 mg

122

Note: Only for use in compounding an oral liquid formulation, for in-hospital use only.

| Price | | Brand or |
|-------------------|-----|--------------|
| (ex man. excl. GS | ST) | Generic |
| \$ | Per | Manufacturer |

⇒ Restricted (RS1576)

Initiation - insomnia secondary to neurodevelopmental disorder

Psychiatrist, paediatrician, neurologist or respiratory specialist

Re-assessment required after 12 months

All of the following:

- 1 Patient has been diagnosed with persistent and distressing insomnia secondary to a neurodevelopmental disorder (including, but not limited to, autism spectrum disorder or attention deficit hyperactivity disorder); and
- 2 Behavioural and environmental approaches have been tried or are inappropriate; and
- 3 Funded modified-release melatonin is to be given at doses no greater than 10 mg per day; and
- 4 Patient is aged 18 years or under.

Continuation - insomnia secondary to neurodevelopmental disorder

Psychiatrist, paediatrician, neurologist or respiratory specialist

Re-assessment required after 12 months

All of the following:

- 1 Patient is aged 18 years or under; and
- 2 Patient has demonstrated clinically meaningful benefit from funded modified-release melatonin (clinician determined); and
- 3 Patient has had a trial of funded modified-release melatonin discontinuation within the past 12 months and has had a recurrence of persistent and distressing insomnia; and
- 4 Funded modified-release melatonin is to be given at doses no greater than 10 mg per day.

Initiation - insomnia where benzodiazepines and zopiclone are contraindicated

Both:

- 1 Patient has insomnia and benzodiazepines and zopiclone are contraindicated; and
- 2 For in-hospital use only.

MIDAZOLAM

| Tab 7.5 mg | | | |
|--|------|-----|-----------------|
| Oral liq 2 mg per ml | | | |
| Inj 1 mg per ml, 5 ml ampoule - 1% DV Jan-19 to 2021 | 2.98 | 10 | Mylan Midazolam |
| Inj 5 mg per ml, 3 ml ampoule - 1% DV Jan-19 to 2021 | 2.36 | 5 | Mylan Midazolam |
| NITRAZEPAM – Restricted: For continuation only | | | |
| → Tab 5 mg | 5.22 | 100 | Nitrados |
| (Nitrados Tab 5 mg to be delisted 1 September 2020) | | | |
| PHENOBARBITONE | | | |
| Inj 200 mg per ml, 1 ml ampoule | | | |
| TEMAZEPAM | | | |
| Tab 10 mg - 1% DV Nov-20 to 2023 | 1.33 | 25 | Normison |
| TRIAZOLAM – Restricted: For continuation only | | | |
| ➡ Tab 125 mcg | | | |
| ➡ Tab 250 mcg | | | |
| ZOPICLONE | | | |
| | | | |

Tab 7.5 mg

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|----------|-------------------------------------|
| Stimulants / ADHD Treatments | | | |
| TOMOXETINE | | | |
| Cap 10 mg - 1% DV Sep-20 to 2022 | | 28 | Generic Partners |
| | 107.03 | | Strattera |
| Cap 18 mg - 1% DV Sep-20 to 2022 | | 28 | Generic Partners |
| | 107.03 | | Strattera |
| Cap 25 mg - 1% DV Sep-20 to 2022 | | 28 | Generic Partners |
| | 107.03 | | Strattera |
| Cap 40 mg - 1% DV Sep-20 to 2022 | | 28 | Generic Partners |
| | 107.03 | | Strattera |
| Cap 60 mg - 1% DV Sep-20 to 2022 | 46.51 | 28 | Generic Partners |
| | 107.03 | | Strattera |
| Cap 80 mg - 1% DV Sep-20 to 2022 | | 28 | Generic Partners |
| | 139.11 | | Strattera |
| Cap 100 mg - 1% DV Sep-20 to 2022 | | 28 | Generic Partners |
| | 139.11 | | Strattera |
| Strattera Cap 10 mg to be delisted 1 September 2020) Strattera Cap 18 mg to be delisted 1 September 2020) Strattera Cap 25 mg to be delisted 1 September 2020) Strattera Cap 40 mg to be delisted 1 September 2020) Strattera Cap 60 mg to be delisted 1 September 2020) Strattera Cap 80 mg to be delisted 1 September 2020) Strattera Cap 100 mg to be delisted 1 September 2020) | | | |
| AFFEINE Tab 100 mg | | | |
| EXAMFETAMINE SULFATE - Restricted see terms below Tab 5 mg - 1% DV Oct-18 to 2021 Restricted (RS1169) hitiation - ADHD | 20.00 | 100 | PSM |
| aediatrician or psychiatrist atient has ADHD (Attention Deficit and Hyperactivity Disorder), dia itiation – Narcolepsy eurologist or respiratory specialist <i>te-assessment required after 24 months</i> atient suffers from narcolepsy. | gnosed according to DS | SM-IV or | ICD 10 criteria. |
| ontinuation – Narcolepsy eurologist or respiratory specialist <i>e-assessment required after 24 months</i> ne treatment remains appropriate and the patient is benefiting from | n treatment. | | |

| _ | | Price | | Brand or |
|-----|--|---------------------------|----------|--------------------------------|
| | | (ex man. excl. GST) \$ | Per | Generic Manufacturer |
| _ | | • | FEI | Waltulaclurei |
| - | THYLPHENIDATE HYDROCHLORIDE – Restricted see terms be | | 00 | Concerto |
| ŧ | Tab extended-release 18 mg | | 30 | Concerta Mathulahanidata ED |
| | | 18.20 | | Methylphenidate ER - |
| t | Tab extended-release 27 mg | 65 11 | 30 | Teva Concerta |
| • | Tab exterided-release 27 mg | 22.00 | 30 | Methylphenidate ER - |
| | | 22.00 | | Teva |
| ſ | Tab extended-release 36 mg | 71 93 | 30 | Concerta |
| • | | 22.40 | 00 | Methylphenidate ER - |
| | | 22.10 | | Teva |
| t | Tab extended-release 54 mg | | 30 | Concerta |
| | · · · · · · · · · · · · · · · · · · · | 26.40 | | Methylphenidate ER - |
| | | | | Teva |
| t | Tab immediate-release 5 mg | 3.20 | 30 | Rubifen |
| t | Tab immediate-release 10 mg | 3.00 | 30 | Ritalin |
| | ° | | | Rubifen |
| t | Tab immediate-release 20 mg | 7.85 | 30 | Rubifen |
| t | Tab sustained-release 20 mg | | 100 | Ritalin SR |
| | J J | 10.95 | 30 | Rubifen SR |
| t | Cap modified-release 10 mg | | 30 | Ritalin LA |
| t | Cap modified-release 20 mg | | 30 | Ritalin LA |
| t | Cap modified-release 30 mg | | 30 | Ritalin LA |
| t | Cap modified-release 40 mg | | 30 | Ritalin LA |
| | Restricted (RS1294) | | | |
| | iation – ADHD (immediate-release and sustained-release form | ulations) | | |
| | ediatrician or psychiatrist | | | |
| | ient has ADHD (Attention Deficit and Hyperactivity Disorder), diagr | losed according to DS | M-IV or | ICD 10 criteria. |
| | iation – Narcolepsy (immediate-release and sustained-release | | | |
| | urologist or respiratory specialist | , | | |
| | assessment required after 24 months | | | |
| | ient suffers from narcolepsy. | | | |
| | ntinuation – Narcolepsy (immediate-release and sustained-rele | ease formulations) | | |
| | urologist or respiratory specialist | | | |
| | assessment required after 24 months | | | |
| | e treatment remains appropriate and the patient is benefiting from t | reatment | | |
| | iation – Extended-release and modified-release formulations | | | |
| | ediatrician or psychiatrist | | | |
| Bo | | | | |
| DU | Patient has ADHD (Attention Deficit and Hyperactivity Disorder Either: |), diagnosed accordin | g to DSN | IIV or ICD 10 criteria; and |
| | 2.1 Patient is taking a currently listed formulation of methyl | henidate hydrochloric | le (imme | diate-release or |
| | sustained-release) which has not been effective due to | | | |
| | 2.2 There is significant concern regarding the risk of diversi | | | |
| | hydrochloride. | | 1010 | add mothylphonidate |
| | - | | | |
| | DAFINIL – Restricted see terms below | | | |
| | Tab 100 mg | 64.00 | 60 | Modavigil |
| | Restricted (RS1761) | | | |
| | iation – Narcolepsy | | | |
| | urologist or respiratory specialist | | | |
| Re | assessment required after 24 months | | | |
| All | of the following: | | | |
| | | | | |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST | | Generic |
| \$ | Per | Manufacturer |

continued...

- 1 The patient has a diagnosis of narcolepsy and has excessive daytime sleepiness associated with narcolepsy occurring almost daily for three months or more; and
- 2 Any of the following:
 - 2.1 The patient has a multiple sleep latency test with a mean sleep latency of less than or equal to 10 minutes and 2 or more sleep onset rapid eye movement periods; or
 - 2.2 A multiple sleep latency test is not possible due to COVID-19 constraints on the health sector; or
 - 2.3 The patient has at least one of: cataplexy, sleep paralysis or hypnagogic hallucinations; and

3 Either:

- 3.1 An effective dose of a listed formulation of methylphenidate or dexamphetamine has been trialled and discontinued because of intolerable side effects; or
- 3.2 Methylphenidate and dexamphetamine are contraindicated.

Continuation - Narcolepsy

Neurologist or respiratory specialist

Re-assessment required after 24 months

The treatment remains appropriate and the patient is benefiting from treatment.

Treatments for Dementia

DONEPEZIL HYDROCHLORIDE

| Tab 5 mg4.34 | 90 | Donepezil-Rex |
|---|----|------------------|
| Tab 10 mg6.64 | 90 | Donepezil-Rex |
| RIVASTIGMINE – Restricted see terms below | | |
| Patch 4.6 mg per 24 hour – 1% DV Apr-20 to 2021 | 30 | Generic Partners |
| Patch 9.5 mg per 24 hour – 1% DV Apr-20 to 2021 | 30 | Generic Partners |
| → Restricted (RS1436) | | |

Initiation

Re-assessment required after 6 months

Both:

- 1 The patient has been diagnosed with dementia; and
- 2 The patient has experienced intolerable nausea and/or vomiting from donepezil tablets.

Continuation

Re-assessment required after 12 months

Both:

- 1 The treatment remains appropriate; and
- 2 The patient has demonstrated a significant and sustained benefit from treatment.

| Treatments for Substance Dependence | | |
|--|----|-------------------------------|
| BUPRENORPHINE WITH NALOXONE - Restricted see terms below | | |
| ■ Tab 2 mg with naloxone 0.5 mg - 1% DV Apr-20 to 2022 | 28 | Buprenorphine Naloxone BNM |
| I Tab 8 mg with naloxone 2 mg – 1% DV Apr-20 to 2022 | 28 | Buprenorphine Naloxone BNM |
| → Restricted (RS1172) Initiation – Detoxification | | |

All of the following:

1 Patient is opioid dependent; and

| | | Price (ex man. excl. GST) | | Brand or Generic |
|--|-----------------|------------------------------|-----------|-----------------------------------|
| | | \$ | Per | Manufacturer |
| continued | | | | |
| 2 Patient is currently engaged with an opioid treatment service3 Prescriber works in an opioid treatment service approved b | | | y of Hea | lth; and |
| nitiation – Maintenance treatment | | | | |
| All of the following: | | | | |
| 1 Patient is opioid dependent; and | | | | |
| 2 Patient will not be receiving methadone; and | | | | handler Minister of Line Min |
| Patient is currently enrolled in an opioid substitution treatm and | ent program ir | h a service a | pproved | by the Ministry of Health; |
| 4 Prescriber works in an opioid treatment service approved b | w the Ministry | of Health | | |
| | y uic wiinisuy | or ricalar. | | |
| SUPROPION HYDROCHLORIDE | | 11.00 | 20 | Zuban |
| Tab modified-release 150 mg | | . 11.00 | 30 | Zyban |
| DISULFIRAM | | 152.00 | 100 | Antohuoo |
| Tab 200 mg | | 103.00 | 100 | Antabuse |
| VALTREXONE HYDROCHLORIDE – Restricted see terms below | | 110 55 | 00 | N altura a a sual |
| J Tab 50 mg | | 112.55 | 30 | Naltraccord |
| nitiation – Alcohol dependence | | | | |
| Both: | | | | |
| 1 Patient is currently enrolled, or is planned to be enrolled, in | a recognised | comprehen | sive trea | tment programme for alco |
| dependence; and | u roooginoou | | | and programme for allo |
| 2 Naltrexone is to be prescribed by, or on the recommendation | on of, a physic | ian working | in an Ale | cohol and Drug Service. |
| nitiation – Constipation | | | | |
| or the treatment of opioid-induced constipation. | | | | |
| NICOTINE - Some items restricted see terms below | | | | |
| Patch 7 mg per 24 hours | | | 28 | Habitrol |
| Patch 14 mg per 24 hours | | | 28 28 | Habitrol |
| Patch 21 mg per 24 hours Oral spray 1 mg per dose | | .21.77 | 28 | Habitrol e.g. Nicorette QuickM |
| | | | | Mouth Spray |
| Lozenge 1 mg | | .18.27 | 216 | Habitrol |
| Lozenge 2 mg | | | 216 | Habitrol |
| Soln for inhalation 15 mg cartridge | | | | e.g. Nicorette Inhalato |
| Gum 2 mg | | .36.39 | 384 | Habitrol (Fruit) |
| | | | | Habitrol (Mint) |
| Gum 4 mg | | .42.07 | 384 | Habitrol (Fruit) |
| - Bestricted (BS1210) | | | | Habitrol (Mint) |
| → Restricted (RS1310) nitiation | | | | |
| ny of the following: | | | | |
| 1 For perioperative use in patients who have a 'nil by mouth' | instruction: or | | | |
| 2 For use within mental health inpatient units; or | | | | |
| 3 For acute use in agitated patients who are unable to leave | the hospital fa | cilities. | | |
| ARENICLINE – Restricted see terms below | | | | |
| Tab 0.5 mg × 11 and 1 mg × 42 – 1% DV Mar-19 to 2021 | | .25.64 | 53 | Varenicline Pfizer |
| Tab 1 mg – 1% DV Mar-19 to 2021 | | | 56 | Varenicline Pfizer |
| Restricted (RS1702) | | - | | |
| | | | | |
| nitiation | | | | |

All of the following:

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST |) | Generic |
| \$ | Per | Manufacturer |

- 1 Short-term therapy as an aid to achieving abstinence in a patient who has indicated that they are ready to cease smoking; and
- 2 The patient is part of, or is about to enrol in, a comprehensive support and counselling smoking cessation programme, which includes prescriber or nurse monitoring; and
- 3 Either:
 - 3.1 The patient has tried but failed to quit smoking after at least two separate trials of nicotine replacement therapy, at least one of which included the patient receiving comprehensive advice on the optimal use of nicotine replacement therapy; or
 - 3.2 The patient has tried but failed to quit smoking using bupropion or nortriptyline; and
- 4 The patient has not had a Special Authority for varenicline approved in the last 6 months; and
- 5 Varenicline is not to be used in combination with other pharmacological smoking cessation treatments and the patient has agreed to this; and
- 6 The patient is not pregnant; and
- 7 The patient will not be prescribed more than 12 weeks' funded varenicline in a 12 month period.

| | (ex man | Price . excl. | GST) | | Brand or Generic |
|---|---|---|---------------------------------|--------------------------------|----------------------------------|
| | | \$ | | Per | Manufacturer |
| Chemotherapeutic Agents | | | | | |
| Alkylating Agents | | | | | |
| BENDAMUSTINE HYDROCHLORIDE - Restricted see terms belo ↓ Inj 25 mg vial ↓ Inj 100 mg vial → Restricted (RS1578) Initiation - treatment naive CLL All of the following: 1 The patient has Binet stage B or C, or progressive stage A cl 2 The patient is chemotherapy treatment naive; and 3 The patient is unable to tolerate toxicity of full-dose FCR; and 4 Patient has ECOG performance status 0-2; and 5 Patient has a Cumulative Illness Rating Scale (CIRS) score of 2 Patient has a Cumulative Illness Rating Scale (CIRS) score of 2 Patient has a Cumulative Illness Rating Scale (CIRS) score of 3 Patient has a Cumulative Illness Rating Scale (CIRS) score of 3 Patient has a Cumulative Illness Rating Scale (CIRS) score of 3 Patient has a Cumulative Illness Rating Scale (CIRS) score of 3 Patient has a Cumulative Illness Rating Scale (CIRS) score of 3 Patient has a Cumulative Illness Rating Scale (CIRS) score of 4 Patient has a Cumulative Illness Rating Scale (CIRS) score of 4 Patient has a Cumulative Illness Rating Scale (CIRS) score of 4 Patient has a Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Patient has A Cumulative Illness Rating Scale (CIRS) score of 4 Pa | hronic lympl d of < 6; and | 085.3 | 8 c leuka | | - |
| 6 Bendamustine is to be administered at a maximum dose of 1 6 cycles. Note: 'Chronic lymphocytic leukaemia (CLL)' includes small lympho to comprise a known standard therapeutic chemotherapy regimen a Initiation – Indolent, Low-grade lymphomas <i>Re-assessment required after 9 months</i> All of the following: The patient has indolent low grade NHL requiring treatment; Patient has a WHO performance status of 0-2; and | ocytic lympho nd supportiv | oma (S | SLL). C | hemothe | |
| 3 Either: 3.1 Both: 3.1.1 Patient is treatment naive; and 3.1.2 Bendamustine is to be administered for a max CD20+); or 3.2 All of the following: 3.2.1 Patient has relapsed refractory disease followi 3.2.2 The patient has not received prior bendamusti 3.2.3 Either: 3.2.3.1 Both: 3.2.3.1.1 Bendamustine is to be administer combination with rituximab when 3.2.3.1.2 Patient has had a rituximab treati 3.2.3.2 Bendamustine is to be administered as refractory patients. | ing prior che ine therapy; red for a ma CD20+); an ment-free in | emothe and aximun id terval | erapy; a n of 6 c of 12 m | and ycles in r nonths or | elapsed patients (in more; or |
| Continuation – Indolent, Low-grade lymphomas Re-assessment required after 9 months Both: | | | | | |

- 1 Patients have not received a bendamustine regimen within the last 12 months; and
- 2 Either:
 - 2.1 Both:
 - 2.1.1 Bendamustine is to be administered for a maximum of 6 cycles in relapsed patients (in combination with rituximab when CD20+); and
 - 2.1.2 Patient has had a rituximab treatment-free interval of 12 months or more; or

| | Price (ex man. excl. GS \$ | Г) Per | Brand or Generic Manufacturer |
|---|----------------------------------|---------------|---|
| continued | | | |
| 2.2 Bendamustine is to be administered as a monothera | apy for a maximum of 6 | cycles in r | ituximab refractory patients. |
| Note: 'indolent, low-grade lymphomas' includes follicular, mantle o | | • | • • |
| macroglobulinaemia. | , J | 7 F -F | · ··· , ································ |
| BUSULFAN | | | |
| Tab 2 mg | 89 25 | 100 | Myleran |
| Inj 6 mg per ml, 10 ml ampoule | | 100 | myloran |
| CARMUSTINE | | | |
| Inj 100 mg vial | 1 297 00 | 1 | BICNU |
| III Too IIIg viai | 1,307.00 | I | Bicnu Heritage |
| | | | Dichu Heillage |
| | | | |
| Tab 2 mg | | | |
| CYCLOPHOSPHAMIDE | | | |
| Tab 50 mg | | 50 | Endoxan |
| | 158.00 | 100 | Procytox |
| Inj 1 g vial – 1% DV Oct-18 to 2021 | | 1 | Endoxan |
| Inj 2 g vial – 1% DV Oct-18 to 2021 | 71.25 | 1 | Endoxan |
| IFOSFAMIDE | | | |
| Inj 1 g vial | | 1 | Holoxan |
| Inj 2 g vial | | 1 | Holoxan |
| LOMUSTINE | | | |
| Cap 10 mg | | 20 | Ceenu |
| Cap 40 mg | | 20 | Ceenu |
| MELPHALAN | | | |
| Tab 2 mg | | | |
| Inj 50 mg vial | | | |
| | | | |
| THIOTEPA | | | |
| Inj 15 mg vial | | | |
| Inj 100 mg vial | | | |
| Anthracyclines and Other Cytotoxic Antibiotics | | | |
| BLEOMYCIN SULPHATE | | | |
| Inj 15,000 iu vial – 1% DV Dec-18 to 2021 | 161.01 | 1 | DBL Bleomycin Sulfate |
| DACTINOMYCIN [ACTINOMYCIN D] | | | |
| Inj 0.5 mg vial | 255.00 | 1 | Cosmegen |
| DAUNORUBICIN | | | |
| Inj 2 mg per ml, 10 ml vial | | 1 | Pfizer |
| DOXORUBICIN HYDROCHLORIDE | | · | |
| | | | |
| Inj 2 mg per ml, 5 ml vial | | 1 | Doxorubicin Ebewe |
| Inj 2 mg per ml, 25 ml vial Note: DV limit applies to all 50 mg presentations of doxor | | I | DOXULUDICITI EDEWE |
| Inj 50 mg vial | abient nyurochionue. | | |
| Inj 2 mg per ml, 50 ml vial | 23.00 | 1 | Doxorubicin Ebewe |
| Inj 2 mg per ml, 100 ml vial – 1% DV Jan-19 to 2021 | | 1 | Doxorubicin Ebewe |
| | | | PANA ANALIN FRAME |
| | 05.00 | | Entrophiain Electro |
| Inj 2 mg per ml, 5 ml vial Inj 2 mg per ml, 25 ml vial | | 1 | Epirubicin Ebewe |
| | 30.00 | 1 | Epirubicin Ebewe |
| Inj 2 mg per ml, 100 ml vial – 1% DV Apr-19 to 2021 | | 1 | Epirubicin Ebewe |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|-------------|-------------------------------------|
| IDARUBICIN HYDROCHLORIDE | | | |
| Inj 5 mg vial – 1% DV Sep-18 to 2021 Inj 10 mg vial – 1% DV Sep-18 to 2021 | | 1 1 | Zavedos Zavedos |
| MITOMYCIN C | | | |
| Inj 5 mg vial | | 1 | Teva |
| Inj 20 mg vial (Omegapharm Inj 20 mg vial to be delisted 1 November 2020) | 816.32 | 1 | Omegapharm |
| MITOZANTRONE Inj 2 mg per ml, 10 ml vial | | 1 | Mitozantrone Ebewe |
| Antimetabolites | | | |
| AZACITIDINE – Restricted see terms below | | | |
| Inj 100 mg vial - 1% DV Dec-18 to 2021 → Restricted (RS1418) Initiation | | 1 | Azacitidine Dr Reddy's |
| Haematologist | | | |
| <i>Re-assessment required after 12 months</i> All of the following: | | | |
| 1 Any of the following: | | | |
| 1.1 The patient has International Prognostic Scoring Sys | stem (IPSS) intermediate | -2 or high | n risk myelodysplastic |
| syndrome; or 1.2 The patient has chronic myelomonocytic leukaemia | (10%-29% marrow blast | s without | myeloproliferative disorder); |
| or | | | |
| The patient has acute myeloid leukaemia with 20-30 Health Organisation Classification (WHO); and | | ge dyspla | sia, according to World |
| The patient has performance status (WHO/ECOG) grade 0- The patient does not have secondary myelodysplastic synd chemotherapy and/or radiation for other diseases; and The patient has an estimated life expectancy of at least 3 m | rome resulting from cher | nical injur | y or prior treatment with |
| Continuation | | | |
| Haematologist | | | |
| Re-assessment required after 12 months Both: | | | |
| No evidence of disease progression, and; and The treatment remains appropriate and patient is benefitting | g from treatment. | | |
| CAPECITABINE | | | |
| Tab 150 mg – 1% DV Jul-20 to 2022 Tab 500 mg – 1% DV Jul-20 to 2022 | | 60 120 | Capercit Capercit |
| CLADRIBINE | | | |
| Inj 2 mg per ml, 5 ml vial | | | |
| Inj 1 mg per ml, 10 ml vial | 749.96 | 1 | Leustatin |
| CYTARABINE | 100.00 | _ | D. |
| Inj 20 mg per ml, 5 ml vial Inj 100 mg per ml, 20 ml vial – 1% DV Dec-18 to 2021 | | 5 1 | Pfizer Pfizer |
| FLUDARABINE PHOSPHATE | | •- | . |
| Tab 10 mg - 1% DV Sep-18 to 2021 | | 20 | Fludara Oral |
| Inj 50 mg vial – 1% DV Nov-19 to 2022 | | 5 | Fludarabine Ebewe |

| | (ex man. | rice excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|----------|--------------------------|--------|-------------------------------------|
| FLUOROURACIL | | | | |
| Inj 50 mg per ml, 20 ml vial - 1% DV Oct-18 to 2021 | | 12.00 | 1 | Fluorouracil Ebewe |
| Inj 50 mg per ml, 100 ml vial - 1% DV Oct-18 to 2021 | | 30.00 | 1 | Fluorouracil Ebewe |
| GEMCITABINE | | | | |
| Inj 10 mg per ml, 100 ml vial – 1% DV Jul-20 to 2023 | | 15.89 | 1 | Gemcitabine Ebewe |
| MERCAPTOPURINE | | | | |
| Tab 50 mg – 1% DV Jul-19 to 2022 | | 37.00 | 25 | Puri-nethol |
| Oral suspension 20 mg per ml | | | 100 ml | Allmercap |
| → Restricted (RS1635) | | | | , anno eup |
| nitiation | | | | |
| Paediatric haematologist or paediatric oncologist | | | | |
| Re-assessment required after 12 months | | | | |
| The patient requires a total dose of less than one full 50 mg tablet per da | ay. | | | |
| Continuation | | | | |
| Paediatric haematologist or paediatric oncologist | | | | |
| Re-assessment required after 12 months | | | | |
| The patient requires a total dose of less than one full 50 mg tablet per da | iy. | | | |
| METHOTREXATE | | | | |
| Tab 2.5 mg - 1% DV Jan-19 to 2021 | | .8.05 | 90 | Trexate |
| Tab 10 mg - 1% DV Jan-19 to 2021 | | 31.75 | 90 | Trexate |
| Inj 2.5 mg per ml, 2 ml vial | | | | |
| Inj 7.5 mg prefilled syringe | | | 1 | Methotrexate Sandoz |
| Inj 10 mg prefilled syringe | | | 1 | Methotrexate Sandoz |
| Inj 15 mg prefilled syringe | | | 1 | Methotrexate Sandoz |
| Inj 20 mg prefilled syringe | | | 1 | Methotrexate Sandoz |
| Inj 25 mg prefilled syringe | | | 1 | Methotrexate Sandoz |
| Inj 30 mg prefilled syringe | | | 1 5 | Methotrexate Sandoz |
| Inj 25 mg per ml, 2 ml vial | | 50.00 | э | DBL Methotrexate Onco-Vial |
| Inj 25 mg per ml, 20 ml vial | | 45.00 | 1 | DBL Methotrexate |
| | | 10.00 | | Onco-Vial |
| Inj 100 mg per ml, 10 ml vial | | 25.00 | 1 | Methotrexate Ebewe |
| Inj 100 mg per ml, 50 ml vial - 1% DV Oct-20 to 2023 | | | 1 | Methotrexate Ebewe |
| PEMETREXED – Restricted see terms below | | | | |
| Ini 100 mg vial | | 60.89 | 1 | Juno Pemetrexed |
| Inj 500 mg vial | | | 1 | Juno Pemetrexed |
| → Restricted (RS1596) | . – | | | |
| nitiation Monothaliama | | | | |

Initiation - Mesothelioma

Re-assessment required after 8 months

Both:

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- 1 Patient has been diagnosed with mesothelioma; and
- 2 Pemetrexed to be administered at a dose of 500 mg/m² every 21 days in combination with cisplatin or carboplatin for a maximum of 6 cycles.

Continuation – Mesothelioma

Re-assessment required after 8 months

All of the following:

- 1 No evidence of disease progression; and
- 2 The treatment remains appropriate and the patient is benefitting from treatment; and

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price Brand or (ex man. excl. GST) Generic \$ Per Manufacturer |
|--|---|
| continued | |
| 3 Pemetrexed to be administered at a dose of | g/m ² every 21 days for a maximum of 6 cycles. |
| Initiation – Non small cell lung cancer | |
| Re-assessment required after 8 months Both: | |
| 1 Patient has locally advanced or metastatic n | amous non-small cell lung carcinoma; and |
| 2 Either: | , , , , , , , , , , , , , , , , , , , |
| 2.1 Both: | |
| 2.1.1 Patient has chemotherapy-na | |
| 2.1.2 Pemetrexed is to be administe carboplatin for a maximum of | a dose of 500 mg/m ² every 21 days in combination with cisplatin o |
| 2.2 All of the following: | 55, 01 |
| 5 | vith platinum based chemotherapy; and |
| 2.2.2 Patient has not received prior | |
| | a dose of 500 mg/m ² every 21 days for a maximum of 6 cycles. |
| Continuation – Non small cell lung cancer Re-assessment required after 8 months | |
| All of the following: | |
| 1 No evidence of disease progression; and | |
| 2 The treatment remains appropriate and the | |
| 3 Pemetrexed is to be administered at a dose | mg/m² every 21 days. |
| THIOGUANINE | |
| Tab 40 mg | |
| Other Cytotoxic Agents | |
| AMSACRINE | |
| Inj 50 mg per ml, 1.5 ml ampoule Inj 75 mg | |
| ANAGRELIDE HYDROCHLORIDE Cap 0.5 mg | |
| | |

| ARSENIC TRIOXIDE Inj 1 mg per ml, 10 ml vial4,817.00 | 10 | Phenasen |
|---|----|-----------------------|
| BORTEZOMIB - Restricted see terms below ↓ Inj 3.5 mg vial - 1% DV Aug-20 to 2022 | 1 | Bortezomib Dr-Reddy's |
| 2 The patient has symptomatic systemic AL amyloidosis. | | |
| COLASPASE [L-ASPARAGINASE] Inj 10,000 iu vial | 1 | Leunase |
| DACARBAZINE | | |
| Inj 200 mg vial62.70 | 1 | DBL Dacarbazine |
| ETOPOSIDE | | |
| Cap 50 mg - 1% DV Jul-19 to 2022 | 20 | Vepesid |
| Cap 100 mg - 1% DV Jul-19 to 2022 | 10 | Vepesid |
| Inj 20 mg per ml, 5 ml vial7.90 | 1 | Rex Medical |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|--|-----------------------------------|----------|-------------------------------------|
| ETOPOSIDE (AS PHOSPHATE) | | | |
| Inj 100 mg vial | 40.00 | 1 | Etopophos |
| HYDROXYUREA | | | |
| Cap 500 mg | 31.76 | 100 | Hydrea |
| IRINOTECAN HYDROCHLORIDE | | | |
| Inj 20 mg per ml, 5 ml vial – 1% DV Apr-19 to 2021 | 71.44 | 1 | Irinotecan Actavis 100 |
| LENALIDOMIDE - Restricted see terms below | | | |
| Cap 5 mg | 5,122.76 | 28 | Revlimid |
| Cap 10 mg | 4,655.25 | 21 | Revlimid |
| | 6,207.00 | 28 | Revlimid |
| Cap 15 mg | 5,429.39 | 21 | Revlimid |
| | 7,239.18 | 28 | Revlimid |
| ↓ Cap 25 mg → Restricted (RS1730) | 7,627.00 | 21 | Revlimid |

Initiation – Relapsed/refractory disease

Haematologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has relapsed or refractory multiple myeloma with progressive disease; and
- 2 Patient has not previously been treated with lenalidomide; and
- 3 Either:
 - 3.1 Lenalidomide to be used as third line* treatment for multiple myeloma; or
 - 3.2 Both:
 - 3.2.1 Lenalidomide to be used as second line treatment for multiple myeloma; and
 - 3.2.2 The patient has experienced severe (grade 3 or higher), dose limiting, peripheral neuropathy with either bortezomib or thalidomide that precludes further treatment with either of these treatments; and
- 4 Lenalidomide to be administered at a maximum dose of 25 mg/day in combination with dexamethasone.

Continuation – Relapsed/refractory disease

Haematologist

Re-assessment required after 6 months

Both:

- 1 No evidence of disease progression; and
- 2 The treatment remains appropriate and patient is benefitting from treatment.

Initiation - Maintenance following first-line autologous stem cell transplant (SCT)

Haematologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has newly diagnosed symptomatic multiple myeloma and has undergone first-line treatment that included an autologous stem cell transplantation; and
- 2 Patient has at least a stable disease response in the first 100 days after transplantation; and
- 3 Lenalidomide maintenance is to be commenced within 6 months of transplantation; and
- 4 The patient has ECOG performance score of 0-1; and
- 5 Lenalidomide to be administered at a maximum dose of 15 mg/day.

Continuation – Maintenance following first-line autologous stem cell transplant (SCT)

Haematologist

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Re-assessment required after 6 months Both:

1 No evidence of disease progression; and

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

2 The treatment remains appropriate and patient is benefitting from treatment.

Note: Indication marked with * is an unapproved indication. A line of treatment is considered to comprise either: a) a known therapeutic chemotherapy regimen and supportive treatments or b) a transplant induction chemotherapy regimen, stem cell transplantation and supportive treatments. Prescriptions must be written by a registered prescriber in the lenalidomide risk management programme operated by the supplier.

OLAPARIB - Restricted see terms below

| t | Tab 100 mg3,701. | 00 56 | Lynparza |
|---|------------------|--------|------------|
| | Tab 150 mg | | Lynparza |
| t | Cap 50 mg7,402. | 00 448 | B Lynparza |

→ Restricted (RS1722)

Initiation

Medical oncologist

Re-assessment required after 12 months

All of the following:

- 1 Patient has a high-grade serous* epithelial ovarian, fallopian tube, or primary peritoneal cancer; and
- 2 There is documentation confirming pathogenic germline BRCA1 or BRCA2 gene mutation; and
- 3 Patient has received at least two lines of previous treatment with platinum-based chemotherapy; and
- 4 Patient has platinum sensitive disease defined as disease progression occurring at least 6 months after the last dose of the penultimate line of platinum-based chemotherapy; and
- 5 Patient's disease must have achieved partial or complete response to treatment with the immediately preceding platinum-based regimen; and
- 6 Patient's disease has not progressed following prior treatment with olaparib; and
- 7 Treatment will be commenced within 8 weeks of the patient's last dose of the immediately preceding platinum-based regimen; and
- 8 Treatment to be administered as maintenance treatment; and
- 9 Treatment not to be administered in combination with other chemotherapy.

Continuation

Medical oncologist

Re-assessment required after 12 months

All of the following:

- 1 Treatment remains clinically appropriate and patient is benefitting from treatment; and
- 2 No evidence of progressive disease; and
- 3 Treatment to be administered as maintenance treatment; and
- 4 Treatment not to be administered in combination with other chemotherapy.

Note: *Note "high-grade serous" includes tumours with high-grade serous features or a high-grade serous component.

PEGASPARGASE - Restricted see terms below

➡ Restricted (RS1190)

Initiation – Newly diagnosed ALL

Limited to 12 months treatment

All of the following:

- 1 The patient has newly diagnosed acute lymphoblastic leukaemia; and
- 2 Pegaspargase to be used with a contemporary intensive multi-agent chemotherapy treatment protocol; and

3 Treatment is with curative intent.

Initiation - Relapsed ALL

Limited to 12 months treatment All of the following:

| | Price | | Brand or Generic |
|---|---------------------------|-----------|---------------------|
| | (ex man. excl. GST) \$ | Per | Manufacturer |
| continued | | | |
| The patient has relapsed acute lymphoblastic leukaemia; and Pegaspargase to be used with a contemporary intensive mult Treatment is with curative intent. | | treatment | t protocol; and |
| PENTOSTATIN [DEOXYCOFORMYCIN] Inj 10 mg vial | | | |
| PROCARBAZINE HYDROCHLORIDE Cap 50 mg | | 50 | Natulan |
| TEMOZOLOMIDE - Restricted see terms below | | | |
| Cap 5 mg – 1% DV May-20 to 2022 | 9.13 | 5 | Temaccord |
| Cap 20 mg - 1% DV May-20 to 2022 | 16.38 | 5 | Temaccord |
| Cap 100 mg – 1% DV May-20 to 2022 | | 5 | Temaccord |
| Cap 140 mg – 1% DV May-20 to 2022 | | 5 | Temaccord |
| Cap 250 mg - 1% DV May-20 to 2022 | | 5 | Temaccord |
| → Restricted (RS1645) | | | |
| Initiation – High grade gliomas | | | |
| Re-assessment required after 12 months | | | |
| All of the following: | | | |
| 1 Fither: | | | |

- 1 Either:
 - 1.1 Patient has newly diagnosed glioblastoma multiforme; or
 - 1.2 Patient has newly diagnosed anaplastic astrocytoma*; and
- 2 Temozolomide is to be (or has been) given concomitantly with radiotherapy; and
- 3 Following concomitant treatment temozolomide is to be used for a maximum of 5 days treatment per cycle at a maximum dose of 200 mg/m² per day.

Continuation - High grade gliomas

Re-assessment required after 12 months

Either:

- 1 Both:
 - 1.1 Patient has glioblastoma multiforme; and
 - 1.2 The treatment remains appropriate and the patient is benefitting from treatment; or
- 2 All of the following:
 - 2.1 Patient has anaplastic astrocytoma*; and
 - 2.2 The treatment remains appropriate and the patient is benefitting from treatment; and
 - 2.3 Adjuvant temozolomide is to be used for a maximum of 24 months.

Initiation - Neuroendocrine tumours

Re-assessment required after 9 months

All of the following:

- 1 Patient has been diagnosed with metastatic or unresectable well-differentiated neuroendocrine tumour*; and
- 2 Temozolomide is to be given in combination with capecitabine; and
- 3 Temozolomide is to be used in 28 day treatment cycles for a maximum of 5 days treatment per cycle at a maximum dose of 200 mg/m² per day; and
- 4 Temozolomide to be discontinued at disease progression.

Continuation – Neuroendocrine tumours

Re-assessment required after 6 months

Both:

- 1 No evidence of disease progression; and
- 2 The treatment remains appropriate and the patient is benefitting from treatment.

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| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|-----|-------------------------------------|
| continued Initiation – ewing's sarcoma | | | |
| Re-assessment required after 9 months | | | |

Patient has relapse or refractory Ewing's sarcoma.

Continuation - ewing's sarcoma

Re-assessment required after 6 months

Both:

1 No evidence of disease progression; and

2 The treatment remains appropriate and the patient is benefitting from treatment.

Note: Indication marked with a * is an unapproved indication. Temozolomide is not funded for the treatment of relapsed high grade glioma.

| THALIDOMIDE | - Restricted | see terms below |
|-------------|--------------|-----------------|
|-------------|--------------|-----------------|

| t | Cap 50 mg | 28 | Thalomid |
|---|------------------|----|----------|
| t | Cap 100 mg756.00 | 28 | Thalomid |

➡ Restricted (RS1192)

Initiation

Re-assessment required after 12 months Any of the following:

- 1 The patient has multiple myeloma; or
- 2 The patient has systemic AL amyloidosis*; or
- 3 The patient has erythema nodosum leprosum.

Continuation

Patient has obtained a response from treatment during the initial approval period.

Notes: Prescription must be written by a registered prescriber in the thalidomide risk management programme operated by the supplier

Maximum dose of 400 mg daily as monotherapy or in a combination therapy regimen

Indication marked with * is an unapproved indication

TRETINOIN

| Cap 10 mg | | 100 | Vesanoid |
|--|----------|-----|-----------|
| VENETOCLAX – Restricted see terms below | | | |
| ↓ Tab 14 × 10 mg, 7 × 50 mg, 21 × 100 mg | 1,771.86 | 42 | Venclexta |
| ↓ Tab 10 mg | | 14 | Venclexta |
| ↓ Tab 50 mg | | 7 | Venclexta |
| ↓ Tab 100 mg | 8,209.41 | 120 | Venclexta |

⇒ Restricted (RS1713)

Initiation – relapsed/refractory chronic lymphocytic leukaemia

Haematologist

Re-assessment required after 7 months All of the following:

- 1 Patient has chronic lymphocytic leukaemia requiring treatment; and
- 2 Patient has received at least one prior therapy for chronic lymphocytic leukaemia; and
- 3 Patient has not previously received funded venetoclax; and
- 4 The patient's disease has relapsed within 36 months of previous treatment; and
- 5 Venetoclax to be used in combination with six 28-day cycles of rituximab commencing after the 5-week dose titration schedule with venetoclax; and
- 6 Patient has an ECOG performance status of 0-2.

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

Continuation - relapsed/refractory chronic lymphocytic leukaemia

Haematologist

Re-assessment required after 6 months

Both:

- 1 Treatment remains clinically appropriate and the patient is benefitting from and tolerating treatment; and
- 2 Venetoclax is to be discontinued after a maximum of 24 months of treatment following the titration schedule unless earlier discontinuation is required due to disease progression or unacceptable toxicity.
- Initiation previously untreated chronic lymphocytic leukaemia with 17p deletion or TP53 mutation*

Haematologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has previously untreated chronic lymphocytic leukaemia; and
- 2 There is documentation confirming that patient has 17p deletion by FISH testing or TP53 mutation by sequencing; and
- 3 Patient has an ECOG performance status of 0-2.

Continuation – previously untreated chronic lymphocytic leukaemia with 17p deletion or TP53 mutation* Haematologist

Re-assessment required after 6 months

The treatment remains clinically appropriate and the patient is benefitting from and tolerating treatment.

Note: 'Chronic lymphocytic leukaemia (CLL)' includes small lymphocytic lymphoma (SLL)* and B-cell prolymphocytic leukaemia (B-PLL)*. Indications marked with * are unapproved indications.

Platinum Compounds

| CARBOPLATIN Inj 10 mg per ml, 45 ml vial – 1% DV Jun-19 to 2021 | 1 | Carboplatin Ebewe |
|---|---|--------------------|
| CISPLATIN | | |
| Inj 1 mg per ml, 50 ml vial12.29 | 1 | DBL Cisplatin |
| Inj 1 mg per ml, 100 ml vial – 1% DV Sep-18 to 2021 | 1 | DBL Cisplatin |
| OXALIPLATIN | | |
| Inj 5 mg per ml, 20 ml vial – 1% DV Feb-20 to 2021 | 1 | Oxaliplatin Accord |
| , , , | | • |
| Dratain Turacina Kinaca Inhibitara | | |

Protein-Tyrosine Kinase Inhibitors

ALECTINIB - Restricted see terms below

↓ Cap 150 mg......7,935.00 224 Alecensa

➡ Restricted (RS1712)

Initiation

Re-assessment required after 6 months

All of the following:

- 1 Patient has locally advanced, or metastatic, unresectable, non-small cell lung cancer; and
- 2 There is documentation confirming that the patient has an ALK tyrosine kinase gene rearrangement using an appropriate ALK test; and
- 3 Patient has an ECOG performance score of 0-2.

Continuation

Re-assessment required after 6 months

Both:

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- $1\,$ No evidence of progressive disease according to RECIST criteria; and
- 2 The patient is benefitting from and tolerating treatment.

| | | Price | | Brand or |
|---|-----------|------------------|-----------|------------------------------|
| | (ex man. | excl. GST) \$ | Per | Generic Manufacturer |
| DASATINIB - Restricted see terms below | | | | |
| ↓ Tab 20 mg | | 774.06 | 60 | Sprycel |
| ↓ Tab 50 mg | 6,2 | 214.20 | 60 | Sprycel |
| ↓ Tab 70 mg | 7,6 | 692.58 | 60 | Sprycel |
| → Restricted (RS1685) | | | | |
| Initiation | | | | |
| Haematologist or any relevant practitioner on the recommendation of a | haemato | ologist | | |
| Re-assessment required after 6 months | | | | |
| Any of the following: | | | | |
| 1 Both: | | | | |
| 1.1 The patient has a diagnosis of chronic myeloid leukaemia | a (CML) i | n blast crisis | or acce | lerated phase; and |
| 1.2 Maximum dose of 140 mg/day; or | | | | |
| 2 Both: | | | | |
| 2.1 The patient has a diagnosis of Philadelphia chromosome | -positive | acute lymph | noid leuk | aemia (Ph+ ALL); and |
| 2.2 Maximum dose of 140 mg/day; or | | | | |
| 3 All of the following: | | | | |
| 3.1 The patient has a diagnosis of CML in chronic phase; and | b | | | |
| 3.2 Maximum dose of 100 mg/day; and | | | | |
| 3.3 Any of the following: | | | | |
| 3.3.1 Patient has documented treatment failure* with im | | | | |
| 3.3.2 Patient has experienced treatment-limiting toxicity | | | | |
| 3.3.3 Patient has high-risk chronic-phase CML defined | | | | |
| 3.3.4 Patients is enrolled in the KISS study** and requir | es dasat | inib treatmer | nt accord | ling to the study protocol. |
| Continuation | | | | |
| Haematologist or any relevant practitioner on the recommendation of a | haemato | ologist | | |
| Re-assessment required after 6 months | | | | |
| All of the following: | | | | |
| 1 Lack of treatment failure while on dasatinib*; and | | | | |
| 2 Dasatinib treatment remains appropriate and the patient is bene | | | | al 100 man/alau fan alanamia |
| 3 Maximum dasatinib dose of 140 mg/day for accelerated or blast phase CML. | pnase C | IVIL and Ph+ | ALL, an | a 100 mg/day for chronic |
| Note: *treatment failure for CML as defined by Leukaemia Net Guidelin https://www.cancertrialsnz.ac.nz/kiss/ | es. **Ki | nase-Inhibiti | on Study | with Sprycel Start-up |
| ERLOTINIB – Restricted see terms below | | | | |
| ↓ Tab 100 mg | | 764.00 | 30 | Tarceva |
| Tab 150 mm | | | 00 | Tanaana |

| t | Tab 150 mg1,146.00 | 30 | Tarceva |
|---|--------------------|----|---------|
| ŧ | Tab 100 mg | 30 | Tarceva |

➡ Restricted (RS1747)

Initiation

Re-assessment required after 4 months

All of the following:

- 1 Patient has locally advanced or metastatic, unresectable, non-squamous Non Small Cell Lung Cancer (NSCLC); and
- 2 There is documentation confirming that the disease expresses activating mutations of EGFR tyrosine kinase; and
- 3 Either:
 - 3.1 Patient is treatment naive; or
 - 3.2 Both:
 - 3.2.1 The patient has discontinued getitinib due to intolerance; and
 - 3.2.2 The cancer did not progress while on gefitinib; and
- 4 Erlotinib is to be given for a maximum of 3 months.

| | (ex man | Price . excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------------|------------------------|----------|-----------|-------------------------------------|
| continued | | | | | |
| Continuation | | | | | |
| Re-assessment required after 6 months | | | | | |
| Both: | inatan NCCL | Chao | not nre | ~~~~~~ | l. and |
| Radiological assessment (preferably including CT scan) ind Erlotinib is to be given for a maximum of 3 months. | Icales NSCL | 6 nas | not pro | gressed | i, anu |
| Continuation – pandemic circumstances | | | | | |
| Re-assessment required after 6 months | | | | | |
| All of the following: | | | | | |
| 1 The patient is clinically benefiting from treatment and contin | ued treatmer | nt rema | ains ap | propriate | e; and |
| 2 Erlotinib to be discontinued at progression; and | | | | | |
| 3 The regular renewal requirements cannot be met due to CC | VID-19 cons | traints | on the | health s | sector. |
| GEFITINIB – Restricted see terms below | | | | | |
| I Tab 250 mg | 1, | 700.00 |) | 30 | Iressa |
| ➡ Restricted (RS1748) | | | | | |
| Initiation | | | | | |
| Re-assessment required after 4 months | | | | | |
| All of the following: | | o Non | Small | | a Consor (NECLC); and |
| Patient has locally advanced, or metastatic, unresectable, r Either: | ion-squamou | 5 11011 | Sillali | | g Cancer (NSCLC), and |
| 2.1 Patient is treatment naive; or2.2 Both: | | | | | |
| 2.2.1 The patient has discontinued erlotinib due to 2.2.2 The cancer did not progress whilst on erlotini | | and | | | |
| 3 There is documentation confirming that disease expresses4 Gefitinib is to be given for a maximum of 3 months. | activating mu | tations | s of EG | FR tyros | sine kinase; and |
| Continuation | | | | | |
| Re-assessment required after 6 months | | | | | |
| Both: | | | | | |
| Radiological assessment (preferably including CT scan) ind Gefitinib is to be given for a maximum of 3 months. | icates NSCL | C has | not pro | gressed | l; and |
| Continuation – pandemic circumstances | | | | | |
| Re-assessment required after 6 months | | | | | |
| All of the following: | | | | | |
| The patient is clinically benefiting from treatment and contin Gefitinib to be discontinued at progression; and | ued treatmer | nt rema | ains; ai | าต | |
| 3 The regular renewal requirements cannot be met due to CC | WID-19 cons | traints | on the | health o | sector |
| | | anto | | nearth | |
| IMATINIB MESILATE Imatinib-AFT is not a registered for the treatment of Gastro Into | natinal Strom | | ouro (| | be Clives brand of imatini |
| mesilate (supplied by Novartis) remains fully subsidised under | Special Auth | ority fo | or patie | , | |
| metastatic malignant GIST, see SA1460 in Section B of the Ph Tab 100 mg | | | | 60 | Glivec |
| Tab foo ing | ∠, | -00.00 | , | 00 | |
| Initiation | | | | | |
| Re-assessment required after 12 months | | | | | |
| Both [.] | | | | | |

Both:

1 Patient has diagnosis (confirmed by an oncologist) of unresectable and/or metastatic malignant gastrointestinal stromal

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. (\$ | | er | Brand or Generic Manufacturer |
|--|--|------------------------|----------|---|
| continued tumour (GIST); and 2 Maximum dose of 400 mg/day. Continuation <i>Re-assessment required after 12 months</i> Adequate clinical response to treatment with imatinib (prescriber deter Note: The Glivec brand of imatinib mesilate (supplied by Novartis) rer with unresectable and/or metastatic malignant GIST, see SA1460 in S | mains fully subsid Section B of the P | harmace | | Schedule. |
| Cap 100 mg Cap 400 mg | | | 30 30 | Imatinib-AFT Imatinib-AFT |
| LAPATINIB - Restricted see terms below ↓ Tab 250 mg | 1,899.00 | 7 | 70 | Tykerb |
| Re-assessment required after 12 months | | | | |
| Either: 1 All of the following: | | | | |
| 1.1 The patient has metastatic breast cancer expressing Hittechnology); and 1.2 The patient has not previously received trastuzumability for the patient has not previously received trastuzumability. 1.3 Lapatinib not to be given in combination with trastuzum 1.4 Lapatinib to be discontinued at disease progression; or 2 All of the following: 2.1 The patient has metastatic breast cancer expressing Hittechnology); and 2.2 The patient started trastuzumab for metastatic breast castarting treatment due to intolerance; and 2.3 The cancer did not progress whilst on trastuzumab; and 2.4 Lapatinib not to be given in combination with trastuzum 2.5 Lapatinib to be discontinued at disease progression. | eatment for HER 2 ab; and ER-2 IHC 3+ or IS ancer but disconti I | 2 positiv 6H+ (incl | e meta | astatic breast cancer; and FISH or other current |
| Continuation Re-assessment required after 12 months All of the following: | | | | |
| The patient has metastatic breast cancer expressing HER-2 IH and The cancer has not progressed at any time point during the pression at any time point during the pression at any time point during the pression. | evious 12 months | Ū | | 077 |
| | | | 20 20 | Tasigna Tasigna |

Haematologist

Re-assessment required after 6 months All of the following:

1 Patient has a diagnosis of chronic myeloid leukaemia (CML) in blast crisis, accelerated phase, or in chronic phase; and

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

2 Either:

- 2.1 Patient has documented CML treatment failure* with imatinib; or
- 2.2 Patient has experienced treatment limiting toxicity with imatinib precluding further treatment with imatinib; and
- 3 Maximum nilotinib dose of 800 mg/day; and
- 4 Subsidised for use as monotherapy only.

Note: *treatment failure as defined by Leukaemia Net Guidelines.

Continuation

Haematologist

Re-assessment required after 6 months

All of the following:

- 1 Lack of treatment failure while on nilotinib as defined by Leukaemia Net Guidelines; and
- 2 Nilotinib treatment remains appropriate and the patient is benefiting from treatment; and
- 3 Maximum nilotinib dose of 800 mg/day; and
- 4 Subsidised for use as monotherapy only.

PALBOCICLIB - Restricted see terms below

| t | Cap 75 mg4,000.00 | 21 | Ibrance |
|---|-------------------|----|---------|
| | | 21 | Ibrance |
| t | Cap 125 mg | 21 | Ibrance |

→ Restricted (RS1731)

Initiation

Medical oncologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has unresectable locally advanced or metastatic breast cancer; and
- 2 There is documentation confirming disease is hormone-receptor positive and HER2-negative; and
- 3 Patient has an ECOG performance score of 0-2; and
- 4 Either:

second or subsequent line setting

- 4.1 Disease has relapsed or progressed during prior endocrine therapy; or
- 4.2 Both:

first line setting

- 4.2.1 Patient is amenorrhoeic, either naturally or induced, with endocrine levels consistent with a postmenopausal state; and
- 4.2.2 Either:
 - 4.2.2.1 Patient has not received prior systemic treatment for metastatic disease; or
 - 4.2.2.2 All of the following:
 - 4.2.2.2.1 Patient commenced treatment with palbociclib in combination with an endocrine agent prior to 1 April 2020; and
 - 4.2.2.2.2 Patient has not received prior systemic endocrine treatment for metastatic disease; and
 - 4.2.2.2.3 There is no evidence of progressive disease; and
- 5 Treatment must be used in combination with an endocrine partner.

Continuation

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Medical oncologist

Re-assessment required after 12 months

All of the following:

- 1 Treatment must be used in combination with an endocrine partner; and
- 2 No evidence of progressive disease; and
- 3 The treatment remains appropriate and the patient is benefitting from treatment.

| | | - | | | |
|---|---------------|----------------------|----------|-----------|-------------------------------------|
| | F (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
| PAZOPANIB – Restricted see terms below | | | | | |
| Tab 200 mg | 1,3 | 334.70 |) | 30 | Votrient |
| ↓ Tab 400 mg | 2,6 | 669.40 |) | 30 | Votrient |
| ➡ Restricted (RS1198) | | | | | |
| Initiation | | | | | |
| Re-assessment required after 3 months | | | | | |
| All of the following: | | | | | |
| 1 The patient has metastatic renal cell carcinoma; and | | | | | |
| 2 Any of the following: | | | | | |
| 2.1 The patient is treatment naive; or | | | | | |
| 2.2 The patient has only received prior cytokine treatment | : or | | | | |
| 2.3 Both: | , - | | | | |
| 2.3.1 The patient has discontinued sunitinib within 3 | months of s | tarting | n treatr | nent du | e to intolerance: and |
| 2.3.2 The cancer did not progress whilst on sunitinib | | | , | | |
| 3 The patient has good performance status (WHO/ECOG grade | | | | | |
| 4 The disease is of predominant clear cell histology; and | o o ב), and | | | | |
| 5 All of the following: | | | | | |
| 5.1 Lactate dehydrogenase level > 1.5 times upper limit o | f normal: an | h | | | |
| 5.2 Haemoglobin level < lower limit of normal; and | i normai, an | ŭ | | | |
| 5.3 Corrected serum calcium level > 10 mg/dL (2.5 mmol/ | (L): and | | | | |
| 5.4 Interval of < 1 year from original diagnosis to the start | ,. | thera | nv: and | 4 | |
| 5.5 Karnofsky performance score of less than or equal to | | anora | py, and | | |
| 5.6 2 or more sites of organ metastasis. | , | | | | |
| Continuation | | | | | |
| Re-assessment required after 3 months | | | | | |
| Both: | | | | | |
| 1 No evidence of disease progression; and | | | | | |
| 2 The treatment remains appropriate and the patient is benefiti | na from trea | tmont | | | |
| Notes: Pazopanib treatment should be stopped if disease progresse | | unoni | • | | |
| Poor prognosis patients are defined as having at least 3 of criteria 5. | | modia | ito nroi | nnoeie r | atients are defined as having |
| 1 or 2 of criteria 5.1-5.6. | 1 0.0. Inter | mould | lie proj | griosis p | alients are defined as having |
| RUXOLITINIB – Restricted see terms below | | | | | |
| Tab 5 mg | 24 | 500.00 | ` | 56 | Jakavi |
| Tab 5 mg | | | | 56 | Jakavi |
| I Tab 20 mg | , | | | 56 | Jakavi |
| → Restricted (RS1726) | | | • | 00 | GUILUVI |
| Initiation | | | | | |
| Haematologist | | | | | |
| Re-assessment required after 12 months | | | | | |
| All of the following: | | | | | |
| 1 The patient has primary myelofibrosis or post-polycythemia v | ora mualafih | vroeic | or noc | t-accont | ial thrombooythemia |
| myelofibrosis: and | ora myelolik | 10313 | or pos | COSCIII | |

- myelofibrosis; and
- 2 Either:
 - 2.1 A classification of risk of intermediate-2 or high-risk myelofibrosis according to either the International Prognostic Scoring System (IPSS), Dynamic International Prognostic Scoring System (DIPSS), or the Age-Adjusted DIPSS; or
 - 2.2 Both:
 - 2.2.1 A classification of risk of intermediate-1 myelofibrosis according to either the International Prognostic Scoring System (IPSS), Dynamic International Prognostic Scoring System (DIPSS), or the Age-Adjusted

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

DIPSS; and

2.2.2 Patient has severe disease-related symptoms that are resistant, refractory or intolerant to available therapy; and

3 A maximum dose of 20 mg twice daily is to be given.

Continuation

Relevant specialist or medical practitioner on the recommendation of a Relevant specialist

Re-assessment required after 12 months

Both:

- 1 The treatment remains appropriate and the patient is benefiting from treatment; and
- 2 A maximum dose of 20 mg twice daily is to be given.

SUNITINIB - Restricted see terms below

| t | Cap 12.5 mg2,315.38 | 28 | Sutent |
|---|---------------------|----|--------|
| | Cap 25 mg | | Sutent |
| | Cap 50 mg | 28 | Sutent |
| | Destricted (DC1740) | | |

→ Restricted (RS1749)

Initiation – RCC

Re-assessment required after 3 months All of the following:

- 1 The patient has metastatic renal cell carcinoma; and
- 2 Any of the following:
 - 2.1 The patient is treatment naive; or
 - 2.2 The patient has only received prior cytokine treatment; or
 - 2.3 The patient has only received prior treatment with an investigational agent within the confines of a bona fide clinical trial which has Ethics Committee approval; or
 - 2.4 Both:

2.4.1 The patient has discontinued pazopanib within 3 months of starting treatment due to intolerance; and

- 2.4.2 The cancer did not progress whilst on pazopanib; and
- 3 The patient has good performance status (WHO/ECOG grade 0-2); and
- 4 The disease is of predominant clear cell histology; and
- 5 All of the following:
 - 5.1 Lactate dehydrogenase level > 1.5 times upper limit of normal; and
 - 5.2 Haemoglobin level < lower limit of normal; and
 - 5.3 Corrected serum calcium level > 10 mg/dL (2.5 mmol/L); and
 - 5.4 Interval of < 1 year from original diagnosis to the start of systemic therapy; and
 - 5.5 Karnofsky performance score of less than or equal to 70; and
 - 5.6 2 or more sites of organ metastasis; and
- 6 Sunitinib to be used for a maximum of 2 cycles.

Notes: RCC - Sunitinib treatment should be stopped if disease progresses.

Poor prognosis patients are defined as having at least 3 of criteria 5.1-5.6. Intermediate prognosis patients are defined as having 1 or 2 of criteria 5.1-5.6.

Continuation – RCC

Re-assessment required after 3 months

Both:

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- 1 No evidence of disease progression; and
- 2 The treatment remains appropriate and the patient is benefiting from treatment.

| Price | е | | Brand or |
|-------------|----------|-----|--------------|
| (ex man. ex | cl. GST) | | Generic |
| \$ | | Per | Manufacturer |

continued...

Initiation - GIST

Re-assessment required after 3 months

Both:

1 The patient has unresectable or metastatic malignant gastrointestinal stromal tumour (GIST); and

- 2 Either:
 - 2.1 The patient's disease has progressed following treatment with imatinib; or
 - 2.2 The patient has documented treatment-limiting intolerance, or toxicity to, imatinib.

Continuation – GIST

Re-assessment required after 6 months

Both:

The patient has responded to treatment or has stable disease as determined by Choi's modified CT response evaluation criteria as follows:

- 1 Any of the following:
 - 1.1 The patient has had a complete response (disappearance of all lesions and no new lesions); or
 - 1.2 The patient has had a partial response (a decrease in size of 10% or more or decrease in tumour density in Hounsfield Units (HU) of 15% or more on CT and no new lesions and no obvious progression of non-measurable disease); or
 - 1.3 The patient has stable disease (does not meet criteria the two above) and does not have progressive disease and no symptomatic deterioration attributed to tumour progression; and
- 2 The treatment remains appropriate and the patient is benefiting from treatment.

Continuation – GIST pandemic circumstances

Re-assessment required after 6 months

All of the following:

- 1 The patient has unresectable or metastatic malignant gastrointestinal stromal tumour (GIST); and
- 2 The patient is clinically benefiting from treatment and continued treatment remains appropriate; and
- 3 Sunitinib is to be discontinued at progression; and
- 4 The regular renewal requirements cannot be met due to COVID-19 constraints on the health sector.

Note: GIST - It is recommended that response to treatment be assessed using Choi's modified CT response evaluation criteria (J Clin Oncol, 2007, 25:1753-1759). Progressive disease is defined as either: an increase in tumour size of 10% or more and not meeting criteria of partial response (PR) by tumour density (HU) on CT; or: new lesions, or new intratumoral nodules, or increase in the size of the existing intratumoral nodules.

Taxanes

| Inj 6 mg per ml, 25 ml vial Inj 6 mg per ml, 50 ml vial – 1% DV Nov-20 to 2023 | | 1 1 | Paclitaxel Ebewe Paclitaxel Ebewe |
|--|-------|--------|--------------------------------------|
| Inj 6 mg per ml, 16.7 ml vial – 1% DV Nov-20 to 2023 | 24.00 | 1 | Paclitaxel Ebewe |
| Inj 6 mg per ml, 5 ml vial | 47.30 | 5 | Paclitaxel Ebewe |
| PACLITAXEL | | | |
| Inj 10 mg per ml, 8 ml vial | 26.95 | 1 | DBL Docetaxel |
| Inj 10 mg per ml, 2 ml vial | 12.40 | 1 | DBL Docetaxel |
| DOCETAXEL | | | |

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|------------|--|
| Treatment of Cytotoxic-Induced Side Effects | | | |
| CALCIUM FOLINATE | 114.00 | 10 | |
| Tab 15 mg Inj 3 mg per ml, 1 ml ampoule | | 10 | DBL Leucovorin Calcium |
| Inj 10 mg per ml, 5 ml ampoule Inj 10 mg per ml, 5 ml vial – 1% DV Jan-20 to 2022 | | 5 1 | Calcium Folinate Ebewe Calcium Folinate |
| Inj 10 mg per ml, 10 ml vial – 1% DV Jan-20 to 2022 | 9.49 | 1 | Sandoz Calcium Folinate |
| Inj 10 mg per ml, 30 ml vial | | 1 | Sandoz Calcium Folinate Ebewe |
| Inj 10 mg per ml, 35 ml vial - 1% DV Nov-19 to 2022 | | 1 | Calcium Folinate Sandoz |
| Inj 10 mg per ml, 100 ml vial – 1% DV Mar-20 to 2022 | 72.00 | 1 | Calcium Folinate Sandoz |
| DEXRAZOXANE – Restricted see terms below | | | o a Cardiaxana |
| → Restricted (RS1695) | | | e.g. Cardioxane |
| Initiation Medical oncologist, paediatric oncologist, haematologist or paediatric | haematologist | | |
| All of the following: | nacinatologist | | |
| 1 Patient is to receive treatment with high dose anthracycline gi | | | d 050mm/m0 daverytisia |
| 2 Based on current treatment plan, patient's cumulative lifetime equivalent or greater; and | dose of anthracycline | will excee | a 250mg/m2 aoxorubicin |
| 3 Dexrazoxane to be administered only whilst on anthracycline | treatment; and | | |
| 4 Either: 4.1 Treatment to be used as a cardioprotectant for a child | or young adult: or | | |
| 4.2 Treatment to be used as a cardioprotectant for a child | | | |
| MESNA | | | |
| Tab 400 mg - 1% DV Nov-19 to 2022 | | 50 50 | Uromitexan Uromitexan |
| Tab 600 mg - 1% DV Nov-19 to 2022 Inj 100 mg per ml, 4 ml ampoule - 1% DV Nov-19 to 2022 | | 50 15 | Uromitexan |
| Inj 100 mg per ml, 10 ml ampoule – 1% DV Nov-19 to 2022 | | 15 | Uromitexan |
| Vinca Alkaloids | | | |
| VINBLASTINE SULPHATE | 070.07 | - | l la calua |
| Inj 1 mg per ml, 10 ml vial | | 5 | Hospira |
| VINCRISTINE SULPHATE Inj 1 mg per ml, 1 ml vial | | 5 | DBL Vincristine Sulfate |
| lnj 1 mg per ml, 2 ml vial | | 5 | DBL Vincristine Sulfate |
| VINORELBINE | | | |
| Inj 10 mg per ml, 1 ml vial Inj 10 mg per ml, 5 ml vial | | 1 | Navelbine Navelbine |
| · • | | | |
| Endocrine Therapy | | | |
| ABIRATERONE ACETATE - Restricted see terms on the next page Tab 250 mg | | 120 | Zutian |
| ↓ Tab 250 mg | | 120 | Zytiga |

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | | | Brand or |
|---------|---------|--------|-----|--------------|
| (ex man | . excl. | . GST) | | Generic |
| | \$ | | Per | Manufacturer |

→ Restricted (RS1746)

Initiation

Medical oncologist, radiation oncologist or urologist *Re-assessment required after 6 months* All of the following:

- 1 Patient has prostate cancer; and
- 2 Patient has metastases; and
- 3 Patient's disease is castration resistant; and
- 4 Either:

4.1 All of the following:

- 4.1.1 Patient is symptomatic; and
- 4.1.2 Patient has disease progression (rising serum PSA) after second line anti-androgen therapy; and
- 4.1.3 Patient has ECOG performance score of 0-1; and
- 4.1.4 Patient has not had prior treatment with taxane chemotherapy; or
- 4.2 All of the following:
 - 4.2.1 Patient's disease has progressed following prior chemotherapy containing a taxane; and
 - 4.2.2 Patient has ECOG performance score of 0-2; and
 - 4.2.3 Patient has not had prior treatment with abiraterone.

Continuation

Medical oncologist, radiation oncologist or urologist

Re-assessment required after 6 months

All of the following:

- 1 No evidence of clinical disease progression; and
- 2 No initiation of taxane chemotherapy with abiraterone; and
- 3 The treatment remains appropriate and the patient is benefiting from treatment.

BICALUTAMIDE

| Tab 50 mg | 28 | Binarex |
|--|-----|----------|
| FLUTAMIDE | | |
| Tab 250 mg | 100 | Flutamin |
| FULVESTRANT – Restricted see terms below | | |
| Inj 50 mg per ml, 5 ml prefilled syringe | 2 | Faslodex |

→ Restricted (RS1732)

Initiation

Medical oncologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has oestrogen-receptor positive locally advanced or metastatic breast cancer; and
- 2 Patient has disease progression following prior treatment with an aromatase inhibitor or tamoxifen for their locally advanced or metastatic disease; and
- 3 Treatment to be given at a dose of 500 mg monthly following loading doses; and
- 4 Treatment to be discontinued at disease progression.

Continuation

Medical oncologist

Re-assessment required after 6 months

All of the following:

- 1 Treatment remains appropriate and patient is benefitting from treatment; and
- 2 Treatment to be given at a dose of 500 mg monthly; and
- 3 No evidence of disease progression.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|-----|-------------------------------------|
| MEGESTROL ACETATE | | | |
| Tab 160 mg - 1% DV Oct-18 to 2021 | 63.53 | 30 | Apo-Megestrol |
| OCTREOTIDE - Restricted see terms below | | | |
| Inj 50 mcg per ml, 1 ml ampoule | | 5 | DBL Octreotide |
| Inj 100 mcg per ml, 1 ml ampoule | | 5 | DBL Octreotide |
| Inj 500 mcg per ml, 1 ml ampoule | 72.50 | 5 | DBL Octreotide |
| Inj 10 mg vial | 1,772.50 | 1 | Sandostatin LAR |
| Inj 20 mg vial | | 1 | Sandostatin LAR |
| Inj 30 mg vial | 2,951.25 | 1 | Sandostatin LAR |
| Bootristed (PC1744) | | | |

Restricted (RS1744)

Initiation - Malignant bowel obstruction

All of the following:

- 1 The patient has nausea* and vomiting* due to malignant bowel obstruction*; and
- 2 Treatment with antiemetics, rehydration, antimuscarinic agents, corticosteroids and analgesics for at least 48 hours has failed; and
- 3 Octreotide to be given at a maximum dose 1500 mcg daily for up to 4 weeks.
- Note: Indications marked with * are unapproved indications

Initiation – acromegaly

Re-assessment required after 3 months

Both:

- 1 The patient has acromegaly; and
- 2 Any of the following:
 - 2.1 Treatment with surgery, radiotherapy and a dopamine agonist has failed; or
 - 2.2 Treatment with octreotide is for an interim period while awaiting the effects of radiotherapy and a dopamine agonist has failed; or
 - 2.3 The patient is unwilling, or unable, to undergo surgery and/or radiotherapy.

Continuation - acromegaly

Both:

- 1 IGF1 levels have decreased since starting octreotide; and
- 2 The treatment remains appropriate and the patient is benefiting from treatment.

Note: In patients with acromegaly octreotide treatment should be discontinued if IGF1 levels have not decreased after 3 months treatment. In patients treated with radiotherapy octreotide treatment should be withdrawn every 2 years, for 1 month, for assessment of remission. Octreotide treatment should be stopped where there is biochemical evidence of remission (normal IGF1 levels) following octreotide treatment withdrawal for at least 4 weeks.

Initiation - Other indications

Any of the following:

- 1 VIPomas and glucagonomas for patients who are seriously ill in order to improve their clinical state prior to definitive surgery; or
- 2 Both:
 - 2.1 Gastrinoma; and
 - 2.2 Either:
 - 2.2.1 Patient has failed surgery; or
 - 2.2.2 Patient in metastatic disease after H2 antagonists (or proton pump inhibitors) have failed; or
- 3 Both:
 - 3.1 Insulinomas; and
 - 3.2 Surgery is contraindicated or has failed; or
- 4 For pre-operative control of hypoglycaemia and for maintenance therapy; or
- 5 Both:

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e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. ex \$ | | Per | Brand or Generic Manufacturer |
|---|----------------------------|------------|-------------|-------------------------------------|
| continued | | | | |
| 5.1 Carcinoid syndrome (diagnosed by tissue pathology 5.2 Disabling symptoms not controlled by maximal medi | | HIAA ana | lysis); a | nd |
| Continuation – Acromegaly - pandemic circumstances | icai illeiapy. | | | |
| Re-assessment required after 6 months | | | | |
| All of the following: | | | | |
| 1 Patient has acromegaly; and | und tractment re | maina an | nunuint | a, and |
| 2 The patient is clinically benefiting from treatment and contir3 The regular renewal requirements cannot be met due to CC | | | | |
| Note: restriction applies only to the long-acting formulations of oct | | | noaint | |
| TAMOXIFEN CITRATE | | | | |
| Tab 10 mg – 1% DV Nov-20 to 2023 | | .00 | 60 | Tamoxifen Sandoz |
| Tab 20 mg - 1% DV Nov-20 to 2023 | 6 | .65 | 60 | Tamoxifen Sandoz |
| Aromatase Inhibitors | | | | |
| ANASTROZOLE | | | | |
| Tab 1 mg | 5 | .04 | 30 | Rolin |
| | | 50 | 00 | Dinar Europeatore |
| Tab 25 mg | 14 | .50 | 30 | Pfizer Exemestane |
| LETROZOLE Tab 2.5 mg - 1% DV Nov-18 to 2021 | 4 | 68 | 30 | Letrole |
| - | | .00 | 00 | Lettole |
| Imaging Agents | | | | |
| AMINOLEVULINIC ACID HYDROCHLORIDE - Restricted see te | | | | |
| Powder for oral soln, 30 mg per ml, 1.5 g vial | | | 1 | Gliolan |
| → Restricted (RS1565) | 44,000 | .00 | 10 | Gliolan |
| nitiation – high grade malignant glioma | | | | |
| All of the following: | | | | |
| 1 Patient has newly diagnosed, untreated, glioblastoma multi | | | | |
| 2 Treatment to be used as adjuvant to fluorescence-guided re | esection; and | | | |
| 3 Patient's tumour is amenable to complete resection. | | | | |
| Immunosuppressants | | | | |
| Calcineurin Inhibitors | | | | |
| CICLOSPORIN | | | | |
| Cap 25 mg | | | 50 | Neoral |
| Cap 50 mg Cap 100 mg | | | 50 50 | Neoral |
| Oral liq 100 mg per ml | | | 50 50 ml | Neoral Neoral |
| Inj 50 mg per ml, 5 ml ampoule | | | 10 | Sandimmun |
| | | | | |
| TACROLIMUS – Restricted see terms on the next page | | | | |
| Cap 0.5 mg | 49 | .60 | 100 | Tacrolimus Sandoz |
| Cap 0.5 mg Cap 0.75 mg | | .30 | 100 | Tacrolimus Sandoz |
| Cap 0.5 mg | 99 84 | .30 .30 | | |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

➡ Restricted (RS1651)

Initiation - organ transplant recipients

Any specialist

For use in organ transplant recipients.

Initiation - non-transplant indications*

Any specialist

Both:

- 1 Patient requires long-term systemic immunosuppression; and
- 2 Ciclosportin has been trialled and discontinued treatment because of unacceptable side effects or inadequate clinical response.

Note: Indications marked with * are unapproved indications

Fusion Proteins

ETANERCEPT - Restricted see terms below

| t | Inj 25 mg vial – 5% DV Sep-19 to 2024 690.00 | 4 | Enbrel |
|---|---|---|--------|
| t | Inj 50 mg autoinjector - 5% DV Sep-19 to 2024 | 4 | Enbrel |
| t | Inj 50 mg syringe – 5% DV Sep-19 to 20241,050.00 | 4 | Enbrel |

→ Restricted (RS1770)

Initiation - juvenile idiopathic arthritis

Rheumatologist or named specialist

Re-assessment required after 6 months Either:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for adalimumab for juvenile idiopathic arthritis (JIA); and 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for JIA; or
- 2 All of the following:
 - 2.1 Patient diagnosed with Juvenile Idiopathic Arthritis (JIA); and
 - 2.2 To be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
 - 2.3 Patient has had severe active polyarticular course JIA for 6 months duration or longer; and
 - 2.4 Patient has tried and not responded to at least three months of oral or parenteral methotrexate (at a dose of 10-20 mg/m² weekly or at the maximum tolerated dose) in combination with either oral corticosteroids (prednisone 0.25 mg/kg or at the maximum tolerated dose) or a full trial of serial intra-articular corticosteroid injections; and
 - 2.5 Both:
 - 2.5.1 Either:
 - 2.5.1.1 Patient has persistent symptoms of poorly-controlled and active disease in at least 20 swollen, tender joints; or
 - 2.5.1.2 Patient has persistent symptoms of poorly-controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, shoulder, cervical spine, hip; and
 - 2.5.2 Physician's global assessment indicating severe disease.

Continuation - juvenile idiopathic arthritis

Rheumatologist or named specialist

Re-assessment required after 6 months

Both:

1 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by

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e.g. Brand indicates brand example only. It is not a contracted product.

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

toxicity or intolerance; and

- 2 Either:
 - 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baseline; or
 - 2.2 On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline.

Initiation - rheumatoid arthritis

Rheumatologist

Re-assessment required after 6 months Either:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for adalimumab for rheumatoid arthritis; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for rheumatoid arthritis; or
- 2 All of the following:
 - 2.1 Patient has had severe and active erosive rheumatoid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP) antibody positive) for six months duration or longer; and
 - 2.2 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
 - 2.3 Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose; and
 - 2.4 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with sulfasalazine and hydroxychloroquine sulphate (at maximum tolerated doses); and
 - 2.5 Any of the following:
 - 2.5.1 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with the maximum tolerated dose of ciclosporin; or
 - 2.5.2 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with intramuscular gold; or
 - 2.5.3 Patient has tried and not responded to at least three months of therapy at the maximum tolerated dose of leflunomide alone or in combination with oral or parenteral methotrexate; and
 - 2.6 Either:
 - 2.6.1 Patient has persistent symptoms of poorly controlled and active disease in at least 20 swollen, tender joints; or
 - 2.6.2 Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
 - 2.7 Either:
 - 2.7.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 2.7.2 C-reactive protein levels not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months.

Continuation - rheumatoid arthritis

Rheumatologist

Re-assessment required after 6 months

All of the following:

1 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by

| Price | | Brand or |
|-------------------|-----|--------------|
| (ex man. excl. GS | Г) | Generic |
| \$ | Per | Manufacturer |

toxicity or intolerance; and

- 2 Either:
 - 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 2.2 On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; and
- 3 Etanercept to be administered at doses no greater than 50 mg every 7 days.

Initiation - ankylosing spondylitis

Rheumatologist

Re-assessment required after 6 months Either:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for adalimumab for ankylosing spondylitis; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for ankylosing spondylitis; or
- 2 All of the following:
 - 2.1 Patient has a confirmed diagnosis of ankylosing spondylitis present for more than six months; and
 - 2.2 Patient has low back pain and stiffness that is relieved by exercise but not by rest; and
 - 2.3 Patient has bilateral sacroiliitis demonstrated by plain radiographs, CT or MRI scan; and
 - 2.4 Patient's ankylosing spondylitis has not responded adequately to treatment with two or more non-steroidal anti-inflammatory drugs (NSAIDs), in combination with anti-ulcer therapy if indicated, while patient was undergoing at least 3 months of a regular exercise regimen for ankylosing spondylitis; and
 - 2.5 Either:
 - 2.5.1 Patient has limitation of motion of the lumbar spine in the sagittal and the frontal planes as determined by the following Bath Ankylosing Spondylitis Metrology Index (BASMI) measures: a modified Schober's test of less than or equal to 4 cm and lumbar side flexion measurement of less than or equal to 10 cm (mean of left and right); or
 - 2.5.2 Patient has limitation of chest expansion by at least 2.5 cm below the average normal values corrected for age and gender (see Notes); and
 - 2.6 Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) of at least 6 on a 0-10 scale.

Notes: The BASDAI must have been determined at the completion of the 3 month exercise trial, but prior to ceasing NSAID treatment. The BASDAI measure must be no more than 1 month old at the time of starting treatment. Average normal chest expansion corrected for age and gender:

| Age | Male | Female |
|-------|--------|--------|
| 18-24 | 7.0 cm | 5.5 cm |
| 25-34 | 7.5 cm | 5.5 cm |
| 35-44 | 6.5 cm | 4.5 cm |
| 45-54 | 6.0 cm | 5.0 cm |
| 55-64 | 5.5 cm | 4.0 cm |
| 65-74 | 4.0 cm | 4.0 cm |
| 75+ | 3.0 cm | 2.5 cm |
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Continuation - ankylosing spondylitis

Rheumatologist

Re-assessment required after 6 months

All of the following:

- 1 Following 12 weeks' initial treatment and for subsequent renewals, treatment has resulted in an improvement in BASDAI of 4 or more points from pre-treatment baseline on a 10 point scale, or an improvement in BASDAI of 50%, whichever is less; and
- 2 Physician considers that the patient has benefited from treatment and that continued treatment is appropriate; and
- 3 Etanercept to be administered at doses no greater than 50 mg every 7 days.

Initiation - psoriatic arthritis

Rheumatologist

Re-assessment required after 6 months Either:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for adalimumab for psoriatic arthritis; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for psoriatic arthritis; or
- 2 All of the following:
 - 2.1 Patient has had severe active psoriatic arthritis for six months duration or longer; and
 - 2.2 Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose; and
 - 2.3 Patient has tried and not responded to at least three months of sulfasalazine at a dose of at least 2 g per day or leflunomide at a dose of up to 20 mg daily (or maximum tolerated doses); and
 - 2.4 Either:
 - 2.4.1 Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen, tender joints; or
 - 2.4.2 Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
 - 2.5 Any of the following:
 - 2.5.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 2.5.2 Patient has an elevated erythrocyte sedimentation rate (ESR) greater than 25 mm per hour; or
 - 2.5.3 ESR and CRP not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months.

Continuation - psoriatic arthritis

Rheumatologist

Re-assessment required after 6 months Both:

- 1 Either:
 - 1.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 1.2 The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior etanercept treatment in the opinion of the treating physician; and
- 2 Etanercept to be administered at doses no greater than 50 mg every 7 days.

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Initiation - severe chronic plaque psoriasis, prior TNF use

Dermatologist

Limited to 4 months treatment

All of the following:

- 1 The patient has had an initial Special Authority approval for adalimumab for severe chronic plaque psoriasis; and
- 2 Either:
 - 2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for severe chronic plaque psoriasis; and
- 3 Patient must be reassessed for continuation after 3 doses.

Initiation - severe chronic plaque psoriasis, treatment-naive

Dermatologist

Limited to 4 months treatment

All of the following:

1 Either:

- 1.1 Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) score of greater than 10, where lesions have been present for at least 6 months from the time of initial diagnosis; or
- 1.2 Patient has severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot, where the plaque or plaques have been present for at least 6 months from the time of initial diagnosis; and
- 2 Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or acitretin; and
- 3 A PASI assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the most recent prior treatment course (but preferably all prior treatment courses), preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course; and
- 4 The most recent PASI or DLQI assessment is no more than 1 month old at the time of initiation.

Note: "Inadequate response" is defined as: for whole body severe chronic plaque psoriasis, a PASI score of greater than 10, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment; for severe chronic plaque psoriasis of the face, hand or foot, at least 2 of the 3 PASI symptom subscores for erythema, thickness and scaling are rated as severe or very severe, and the skin area affected is 30% or more of the face, palm of a hand or sole of a foot, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment. **Continuation – severe chronic plaque psoriasis**

Dermatologist

Re-assessment required after 6 months Both:

1 Either:

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1.1 Both:

- 1.1.1 Patient had "whole body" severe chronic plaque psoriasis at the start of treatment; and
- 1.1.2 Either:
 - 1.1.2.1 Following each prior etanercept treatment course the patient has a PASI score which is reduced by 75% or more, or is sustained at this level, when compared with the pre-etanercept treatment baseline value; or
 - 1.1.2.2 Following each prior etanercept treatment course the patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, when compared with the pre-treatment baseline value; or

1.2 Both:

1.2.1 Patient had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment; and

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- 1.2.2 Either:
 - 1.2.2.1 Following each prior etanercept treatment course the patient has a reduction in the PASI symptom subscores for all 3 of erythema, thickness and scaling, to slight or better, or sustained at this level, as compared to the treatment course baseline values; or
 - 1.2.2.2 Following each prior etanercept treatment course the patient has a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-etanercept treatment baseline value; and
- 2 Etanercept to be administered at doses no greater than 50 mg every 7 days.

Initiation – pyoderma gangrenosum

Dermatologist

All of the following:

- 1 Patient has pyoderma gangrenosum*; and
- 2 Patient has received three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisone, ciclosporin, azathioprine, or methotrexate) and not received an adequate response; and
- 3 A maximum of 8 doses.

Note: Indications marked with * are unapproved indications.

Continuation – pyoderma gangrenosum

Dermatologist

All of the following:

- 1 Patient has shown clinical improvement; and
- 2 Patient continues to require treatment; and
- 3 A maximum of 8 doses.

Initiation - adult-onset Still's disease

Rheumatologist

Re-assessment required after 6 months Either:

1 Both:

- 1.1 Either:
 - 1.1.1 The patient has had an initial Special Authority approval for etanercept for adult-onset Still's disease (AOSD); or
 - 1.1.2 The patient has been started on tocilizumab for AOSD in a DHB hospital in accordance with the Section H rules; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from etanercept and/or tocilizumab; or
 - 1.2.2 The patient has received insufficient benefit from at least a three-month trial of adalimumab and/or tocilizumab such that they do not meet the renewal criteria for AOSD; or
- 2 All of the following:
 - 2.1 Patient diagnosed with AOSD according to the Yamaguchi criteria (J Rheumatol 1992;19:424-430); and
 - 2.2 Patient has tried and not responded to at least 6 months of glucocorticosteroids at a dose of at least 0.5 mg/kg, non-steroidal antiinflammatory drugs (NSAIDs) and methotrexate; and
 - 2.3 Patient has persistent symptoms of disabling poorly controlled and active disease.

Continuation - adult-onset Still's disease

Rheumatologist

Re-assessment required after 6 months

The patient has a sustained improvement in inflammatory markers and functional status.

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Initiation - undifferentiated spondyloarthritis

Rheumatologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has undifferentiated peripheral spondyloarthritis* with active peripheral joint arthritis in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
- 2 Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose; and
- 3 Patient has tried and not responded to at least three months of sulfasalazine at a dose of at least 2 g per day (or maximum tolerated dose); and
- 4 Patient has tried and not responded to at least three months of leflunomide at a dose of up to 20 mg daily (or maximum tolerated dose); and
- 5 Any of the following:
 - 5.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 5.2 Patient has an elevated erythrocyte sedimentation rate (ESR) greater than 25 mm per hour measured no more than one month prior to the date of this application; or
 - 5.3 ESR and CRP not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months.

Note: Indications marked with * are unapproved indications.

Continuation - undifferentiated spondyloarthritis

Rheumatologist or medical practitioner on the recommendation of a Rheumatologist

Re-assessment required after 6 months

All of the following:

- 1 Either:
 - 1.1 Applicant is a rheumatologist; or
 - 1.2 Applicant is a Practitioner and confirms that a rheumatologist has provided a letter, email or fax recommending that the patient continues with etanercept treatment; and
- 2 Either:
 - 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 2.2 The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior etanercept treatment in the opinion of the treating physician; and
- 3 Etanercept to be administered at doses no greater than 50 mg dose every 7 days.

Monoclonal Antibodies

| ABCIXIMAB – Restricted see terms below ↓ Inj 2 mg per ml, 5 ml vial | 1 | ReoPro |
|--|-------------|-------------------------------|
| For use in patients with acute coronary syndromes undergoing percutaneous coronar For use in patients undergoing intra-cranial intervention. | y interve | ention; or |
| ADALIMUMAB – Restricted see terms on the next page Inj 20 mg per 0.4 ml syringe 1,599.96 Inj 40 mg per 0.8 ml pen 1,599.96 Inj 40 mg per 0.8 ml syringe 1,599.96 | 2 2 2 | Humira HumiraPen Humira |

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➡ Restricted (RS1771)

Initiation - juvenile idiopathic arthritis

Rheumatologist or named specialist *Re-assessment required after 6 months* Either:

1 Either:

- 1.1 Both:
 - 1.1.1 The patient has had an initial Special Authority approval for etanercept for juvenile idiopathic arthritis (JIA); and
 - 1.1.2 Either:
 - 1.1.2.1 The patient has experienced intolerable side effects from etanercept; or
 - 1.1.2.2 The patient has received insufficient benefit from etanercept to meet the renewal criteria for etanercept for JIA; or
- 2 All of the following:
 - 2.1 Patient diagnosed with Juvenile Idiopathic Arthritis (JIA); and
 - 2.2 To be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
 - 2.3 Patient has had severe active polyarticular course JIA for 6 months duration or longer; and
 - 2.4 Patient has tried and not responded to at least three months of oral or parenteral methotrexate (at a dose of 10-20 mg/m² weekly or at the maximum tolerated dose) in combination with either oral corticosteroids (prednisone 0.25 mg/kg or at the maximum tolerated dose) or a full trial of serial intra-articular corticosteroid injections; and
 - 2.5 Both:
 - 2.5.1 Either:
 - 2.5.1.1 Patient has persistent symptoms of poorly-controlled and active disease in at least 20 swollen, tender joints; or
 - 2.5.1.2 Patient has persistent symptoms of poorly-controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, shoulder, cervical spine, hip; and
 - 2.5.2 Physician's global assessment indicating severe disease.

Continuation - juvenile idiopathic arthritis

Rheumatologist or named specialist

Re-assessment required after 6 months

Both:

- 1 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
- 2 Either:
 - 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baseline; or
 - 2.2 On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline.

Initiation - fistulising Crohn's disease

Gastroenterologist

Re-assessment required after 4 months

All of the following:

- 1 Patient has confirmed Crohn's disease; and
- 2 Either:
 - 2.1 Patient has one or more complex externally draining enterocutaneous fistula(e); or
 - 2.2 Patient has one or more rectovaginal fistula(e); and

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3 A Baseline Fistula Assessment (a copy of which is available at

www.pharmac.govt.nz/latest/BaselineFistulaAssessment.pdf) has been completed and is no more than 1 month old at the time of application.

Continuation - fistulising Crohn's disease

Gastroenterologist

Re-assessment required after 6 months

Either:

- 1 The number of open draining fistulae have decreased from baseline by at least 50%; or
- 2 There has been a marked reduction in drainage of all fistula(e) from baseline as demonstrated by a reduction in the Fistula Assessment score, together with less induration and patient-reported pain.

Initiation - Crohn's disease - adults

Gastroenterologist

Re-assessment required after 3 months

All of the following:

- 1 Patient has severe active Crohn's disease; and
- 2 Any of the following:
 - 2.1 Patient has a Crohn's Disease Activity Index (CDAI) score of greater than or equal to 300; or
 - 2.2 Patient has extensive small intestine disease affecting more than 50 cm of the small intestine; or
 - 2.3 Patient has evidence of short gut syndrome or would be at risk of short gut syndrome with further bowel resection; or
 - 2.4 Patient has an ileostomy or colostomy, and has intestinal inflammation; and
- 3 Patient has tried but had an inadequate response to, or has experienced intolerable side effects from, prior systemic therapy with immunomodulators at maximum tolerated doses (unless contraindicated) and corticosteroids; and
- 4 Surgery (or further surgery) is considered to be clinically inappropriate.

Continuation - Crohn's disease - adults

Gastroenterologist

Re-assessment required after 3 months

Both:

- 1 Either:
 - 1.1 Either:
 - 1.1.1 CDAI score has reduced by 100 points from the CDAI score when the patient was initiated on adalimumab; or
 - 1.1.2 CDAI score is 150 or less; or
 - 1.2 Both:
 - 1.2.1 The patient has demonstrated an adequate response to treatment but CDAI score cannot be assessed; and
 - 1.2.2 Applicant to indicate the reason that CDAI score cannot be assessed; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - Crohn's disease - children

Gastroenterologist

Re-assessment required after 3 months

All of the following:

- 1 Paediatric patient has severe active Crohn's disease; and
- 2 Either:
 - 2.1 Patient has a Paediatric Crohn's Disease Activity Index (PCDAI) score of greater than or equal to 30; or
 - 2.2 Patient has extensive small intestine disease; and
- 3 Patient has tried but had an inadequate response to, or has experienced intolerable side effects from, prior systemic therapy with immunomodulators at maximum tolerated doses (unless contraindicated) and corticosteroids; and

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4 Surgery (or further surgery) is considered to be clinically inappropriate.

Continuation - Crohn's disease - children

Gastroenterologist

Re-assessment required after 3 months Both:

- 1 Any of the following:
 - 1.1 PCDAI score has reduced by 100 points from the PCDAI score when the patient was initiated on adalimumab; or
 - 1.2 PCDAI score is 150 or less; or
 - 1.3 The patient has demonstrated an adequate response to treatment but PCDAI score cannot be assessed; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - rheumatoid arthritis

Rheumatologist

Re-assessment required after 6 months Either:

- 1 Both:
 - 1.1 The patient has had an initial Special Authority approval for etanercept for rheumatoid arthritis; and
 - 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from etanercept; or
 - 1.2.2 The patient has received insufficient benefit from etanercept to meet the renewal criteria for etanercept for rheumatoid arthritis; or
- 2 All of the following:
 - 2.1 Patient has had severe and active erosive rheumatoid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP) antibody positive) for six months duration or longer; and
 - 2.2 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
 - 2.3 Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose; and
 - 2.4 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with sulfasalazine and hydroxychloroquine sulphate (at maximum tolerated doses); and
 - 2.5 Any of the following:
 - 2.5.1 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with the maximum tolerated dose of ciclosporin; or
 - 2.5.2 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with intramuscular gold; or
 - 2.5.3 Patient has tried and not responded to at least three months of therapy at the maximum tolerated dose of leflunomide alone or in combination with oral or parenteral methotrexate; and
 - 2.6 Either:
 - 2.6.1 Patient has persistent symptoms of poorly controlled and active disease in at least 20 swollen, tender joints; or
 - 2.6.2 Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
 - 2.7 Either:
 - 2.7.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 2.7.2 C-reactive protein levels not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months.

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Continuation - rheumatoid arthritis

Rheumatologist *Re-assessment required after 6 months* All of the following:

- 1 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
- 2 Either:
 - 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 2.2 On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; and
- 3 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - ankylosing spondylitis

Rheumatologist *Re-assessment required after 6 months* Either:

- 1 Both:
 - 1.1 The patient has had an initial Special Authority approval for etanercept for ankylosing spondylitis; and
 - 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from etanercept; or
 - 1.2.2 The patient has received insufficient benefit from etanercept to meet the renewal criteria for etanercept for ankylosing spondylitis; or
- 2 All of the following:
 - 2.1 Patient has a confirmed diagnosis of ankylosing spondylitis present for more than six months; and
 - 2.2 Patient has low back pain and stiffness that is relieved by exercise but not by rest; and
 - 2.3 Patient has bilateral sacroiliitis demonstrated by plain radiographs, CT or MRI scan; and
 - 2.4 Patient's ankylosing spondylitis has not responded adequately to treatment with two or more non-steroidal anti-inflammatory drugs (NSAIDs), in combination with anti-ulcer therapy if indicated, while patient was undergoing at least 3 months of a regular exercise regimen for ankylosing spondylitis; and
 - 2.5 Either:
 - 2.5.1 Patient has limitation of motion of the lumbar spine in the sagittal and the frontal planes as determined by the following Bath Ankylosing Spondylitis Metrology Index (BASMI) measures: a modified Schober's test of less than or equal to 4 cm and lumbar side flexion measurement of less than or equal to 10 cm (mean of left and right); or
 - 2.5.2 Patient has limitation of chest expansion by at least 2.5 cm below the average normal values corrected for age and gender (see Notes); and
 - 2.6 Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) of at least 6 on a 0-10 scale.

Notes: The BASDAI must have been determined at the completion of the 3 month exercise trial, but prior to ceasing NSAID treatment. The BASDAI measure must be no more than 1 month old at the time of starting treatment. Average normal chest expansion corrected for age and gender:

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| Age | Male | Female |
| 18-24 | 7.0 cm | 5.5 cm |
| 25-34 | 7.5 cm | 5.5 cm |
| 35-44 | 6.5 cm | 4.5 cm |
| 45-54 | 6.0 cm | 5.0 cm |
| 55-64 | 5.5 cm | 4.0 cm |
| 65-74 | 4.0 cm | 4.0 cm |
| 75+ | 3.0 cm | 2.5 cm |
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Continuation – ankylosing spondylitis

Rheumatologist

continued

Re-assessment required after 6 months

All of the following:

- 1 Following 12 weeks' initial treatment and subsequent renewals, treatment has resulted in an improvement in BASDAI of 4 or more points from pre-treatment baseline on a 10 point scale, or an improvement in BASDAI of 50%, whichever is less; and
- 2 Physician considers that the patient has benefited from treatment and that continued treatment is appropriate; and
- 3 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - psoriatic arthritis

Rheumatologist

Re-assessment required after 6 months Either

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- 1 Both:
 - 1.1 The patient has had an initial Special Authority approval for etanercept for psoriatic arthritis; and
 - 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from etanercept; or
 - 1.2.2 The patient has received insufficient benefit from etanercept to meet the renewal criteria for etanercept for psoriatic arthritis; or
- 2 All of the following:
 - 2.1 Patient has had severe active psoriatic arthritis for six months duration or longer; and
 - 2.2 Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose; and
 - 2.3 Patient has tried and not responded to at least three months of sulfasalazine at a dose of at least 2 g per day or leflunomide at a dose of up to 20 mg daily (or maximum tolerated doses); and
 - 2.4 Either:
 - 2.4.1 Patient has persistent symptoms of poorly controlled and active disease in at least 15 swollen, tender joints; or
 - 2.4.2 Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
 - 2.5 Any of the following:
 - 2.5.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 2.5.2 Patient has an elevated erythrocyte sedimentation rate (ESR) greater than 25 mm per hour; or
 - 2.5.3 ESR and CRP not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months.

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Continuation - psoriatic arthritis

Rheumatologist

Re-assessment required after 6 months Both:

1 Fither:

- 1.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
- 1.2 The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior adalimumab treatment in the opinion of the treating physician; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - plaque psoriasis, prior TNF use

Dermatologist

Limited to 4 months treatment

Both:

1 The patient has had an initial Special Authority approval for etanercept for severe chronic plaque psoriasis; and

2 Either:

- 2.1 The patient has experienced intolerable side effects from etanercept; or
- 2.2 The patient has received insufficient benefit from etanercept to meet the renewal criteria for etanercept for severe chronic plaque psoriasis.

Initiation - plaque psoriasis, treatment-naive

Dermatologist

Limited to 4 months treatment

All of the following:

- 1 Either:
 - 1.1 Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) score of greater than 10, where lesions have been present for at least 6 months from the time of initial diagnosis; or
 - 1.2 Patient has severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot, where the plaque or plaques have been present for at least 6 months from the time of initial diagnosis; and
- 2 Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or acitretin; and
- 3 A PASI assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the most recent prior treatment course (but preferably all prior treatment courses), preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course; and
- 4 The most recent PASI or DLQI assessment is no more than 1 month old at the time of initiation.

Note: "Inadequate response" is defined as: for whole body severe chronic plaque psoriasis, a PASI score of greater than 10, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment; for severe chronic plaque psoriasis of the face, hand or foot, at least 2 of the 3 PASI symptom subscores for erythema, thickness and scaling are rated as severe or very severe, and the skin area affected is 30% or more of the face, palm of a hand or sole of a foot, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment. **Continuation – plaque psoriasis**

Dermatologist

Re-assessment required after 6 months Both:

1 Either:

- 1.1 Both:
 - 1.1.1 Patient had "whole body" severe chronic plaque psoriasis at the start of treatment; and

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- 1.1.2 Either:
 - 1.1.2.1 Following each prior adalimumab treatment course the patient has a PASI score which is reduced by 75% or more, or is sustained at this level, when compared with the pre-adalimumab treatment baseline value; or
 - 1.1.2.2 Following each prior adalimumab treatment course the patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, when compared with the pre-treatment baseline value; or

1.2 Both:

- 1.2.1 Patient had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment; and
- 1.2.2 Either:
 - 1.2.2.1 Following each prior adalimumab treatment course the patient has a reduction in the PASI symptom subscores for all 3 of erythema, thickness and scaling, to slight or better, or sustained at this level, as compared to the treatment course baseline values; or
 - 1.2.2.2 Following each prior adalimumab treatment course the patient has a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-etanercept treatment baseline value; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - pyoderma gangrenosum

Dermatologist

All of the following:

- 1 Patient has pyoderma gangrenosum*; and
- 2 Patient has received three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisone, ciclosporin, azathioprine, or methotrexate) and not received an adequate response; and
- 3 A maximum of 8 doses.

Note: Indications marked with * are unapproved indications.

Continuation – pyoderma gangrenosum

Dermatologist

All of the following:

- 1 Patient has shown clinical improvement; and
- 2 Patient continues to require treatment; and
- 3 A maximum of 8 doses.

Initiation - adult-onset Still's disease

Rheumatologist

Re-assessment required after 6 months Either:

1 Both:

- 1.1 Either:
 - 1.1.1 The patient has had an initial Special Authority approval for etanercept for adult-onset Still's disease (AOSD); or
 - 1.1.2 The patient has been started on tocilizumab for AOSD in a DHB hospital in accordance with the Section H rules; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from etanercept and/or tocilizumab; or
 - 1.2.2 The patient has received insufficient benefit from at least a three-month trial of etanercept and/or
 - tocilizumab such that they do not meet the renewal criteria for AOSD; or
- 2 All of the following:
 - 2.1 Patient diagnosed with AOSD according to the Yamaguchi criteria (J Rheumatol 1992;19:424-430); and

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- 2.2 Patient has tried and not responded to at least 6 months of glucocorticosteroids at a dose of at least 0.5 mg/kg, non-steroidal antiinflammatory drugs (NSAIDs) and methotrexate; and
- 2.3 Patient has persistent symptoms of disabling poorly controlled and active disease.

Continuation – adult-onset Still's disease

Rheumatologist

Re-assessment required after 6 months

The patient has a sustained improvement in inflammatory markers and functional status.

Initiation – severe Behcet's disease

Any relevant practitioner

Re-assessment required after 3 months

All of the following:

- 1 The patient has severe Behcet's disease that is significantly impacting the patient's quality of life (see Notes); and
- 2 Either:
 - 2.1 The patient has severe ocular, neurological, gastrointestinal, rheumatological, mucocutaneous and/or vasculitic symptoms and has not responded adequately to treatment with infliximab (see Notes); or
 - 2.2 The patient has severe ocular, neurological, gastrointestinal, rheumatological, mucocutaneous and/or vasculitic symptoms and has experienced intolerable side effects from treatment with infliximab; and
- 3 The patient is experiencing significant loss of quality of life; and
- 4 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Note: Behcet's disease diagnosed according to the International Study Group for Behcet's disease. Lancet 1990;335(8697):1078-80. Quality of life measured using an appropriate quality of life scale such as that published in Gilworth et al, J Rheumatol. 2004;31:931-7.

Continuation - severe Behcet's disease

Any relevant practitioner

Re-assessment required after 6 months

Both:

- 1 Patient has had a good clinical response to initial treatment with measurably improved quality of life; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Initiation - severe ocular inflammation

Re-assessment required after 4 months Fither:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for infliximab for severe ocular inflammation; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from infliximab; or
 - 1.2.2 The patient has received insufficient benefit from infliximab to meet the renewal criteria for infliximab for severe ocular inflammation; or

2 Both:

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- 2.1 Patient has severe, vision-threatening ocular inflammation requiring rapid control; and
- 2.2 Any of the following:
 - 2.2.1 Treatment with high-dose steroids (intravenous methylprednisolone) followed by high dose oral steroids has proven ineffective at controlling symptoms; or
 - 2.2.2 Patient developed new inflammatory symptoms while receiving high dose steroids; or
 - 2.2.3 Patient is aged under 8 years and treatment with high dose oral steroids and other immunosuppressants has proven ineffective at controlling symptoms.

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Continuation - severe ocular inflammation

Re-assessment required after 12 months

Both:

- 1 Any of the following:
 - 1.1 The patient has had a good clinical response following 3 initial doses; or
 - 1.2 Following each 12-month treatment period, the patient has had a sustained reduction in inflammation (Standardisation of Uveitis Nomenclature (SUN) criteria < ½+ anterior chamber or vitreous cells, absence of active vitreous or retinal lesions, or resolution of uveitic cystoid macular oedema); or
 - 1.3 Following each 12-month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 10mg daily, or steroid drops less than twice daily if under 18 years old; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Note: A trial withdrawal should be considered after every 24 months of stability, unless the patient is deemed to have extremely high risk of irreversible vision loss if adalimumab is withdrawn.

Initiation - chronic ocular inflammation

Re-assessment required after 4 months

Either:

1 Both:

1.1 The patient has had an initial Special Authority approval for infliximab for chronic ocular inflammation; and

- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from infliximab; or
 - 1.2.2 The patient has received insufficient benefit from infliximab to meet the renewal criteria for infliximab for chronic ocular inflammation; or
- 2 Both:
 - 2.1 Patient has severe uveitis uncontrolled with treatment of steroids and other immunosuppressants with a severe risk of vision loss; and
 - 2.2 Any of the following:
 - 2.2.1 Patient is 18 years or older and treatment with at least two other immunomodulatory agents has proven ineffective; or
 - 2.2.2 Patient is under 18 years and treatment with methotrexate has proven ineffective or is not tolerated at a therapeutic dose; or
 - 2.2.3 Patient is under 8 years and treatment with steroids or methotrexate has proven ineffective or is not tolerated at a therapeutic dose; or disease requires control to prevent irreversible vision loss prior to achieving a therapeutic dose of methotrexate.

Continuation - chronic ocular inflammation

Re-assessment required after 12 months Both:

- 1 Any of the following:
 - 1.1 The patient has had a good clinical response following 12 weeks' initial treatment; or
 - 1.2 Following each 12-month treatment period, the patient has had a sustained reduction in inflammation (Standardisation of Uveitis Nomenclature (SUN) criteria < ½+ anterior chamber or vitreous cells, absence of active vitreous or retinal lesions, or resolution of uveitic cystoid macular oedema); or
 - 1.3 Following each 12-month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 10mg daily, or steroid drops less than twice daily if under 18 years old; and
- 2 Adalimumab to be administered at doses no greater than 40 mg every 14 days.

Note: A trial withdrawal should be considered after every 24 months of stability, unless the patient is deemed to have extremely high risk of irreversible vision loss if adalimumab is withdrawn.

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Initiation - hidradenitis suppurativa

Dermatologist

Re-assessment required after 4 months

All of the following:

- 1 Patient has hidradenitis suppurativa Hurley Stage II or Hurley Stage III lesions in distinct anatomic areas; and
- 2 Patient has tried, but had an inadequate response to at least a 90 day trial of systemic antibiotics or patient has demonstrated intolerance to or has contraindications for systemic antibiotics; and
- 3 The patient has 3 or more active lesions (e.g. inflammatory nodules, abscesses, draining fistulae); and
- 4 The patient has a Dermatology Quality of Life Index of 10 or more and the assessment is no more than 1 month old at time of application; and
- 5 Following the initial loading doses, adalimumab is to be administered at doses no greater than 40mg every 7 days.

Continuation - hidradenitis suppurativa

Dermatologist

Re-assessment required after 6 months

All of the following:

- 1 The patient has a reduction in active lesions (e.g. inflammatory nodules, abscesses, draining fistulae) of 25% or more from baseline; and
- 2 The patient has a Dermatology Quality of Life Index improvement of 4 or more from baseline; and
- 3 Adalimumab is to be administered at doses no greater than 40mg every 7 days. Fortnightly dosing has been considered.

AFLIBERCEPT - Restricted see terms below

| Inj 40 mg per ml, 0.1 ml vial | | Eylea |
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→ Restricted (RS1659)

Initiation – Wet Age Related Macular Degeneration

Ophthalmologist

Re-assessment required after 3 months Either:

1 All of the following:

- 1.1 Any of the following:
 - 1.1.1 Wet age-related macular degeneration (wet AMD); or
 - 1.1.2 Polypoidal choroidal vasculopathy; or
 - 1.1.3 Choroidal neovascular membrane from causes other than wet AMD; and
- 1.2 Either:
 - 1.2.1 The patient has developed severe endophthalmitis or severe posterior uveitis following treatment with bevacizumab; or
 - 1.2.2 There is worsening of vision or failure of retina to dry despite three intraocular injections of bevacizumab four weeks apart; and
- 1.3 There is no structural damage to the central fovea of the treated eye; and
- 1.4 Patient has not previously been treated with ranibizumab for longer than 3 months; or
- 2 Either:
 - 2.1 Patient has current approval to use ranibizumab for treatment of wAMD and was found to be intolerant to ranibizumab within 3 months; or
 - 2.2 Patient has previously* (*before June 2018) received treatment with ranibizumab for wAMD and disease was stable while on treatment.

Continuation – Wet Age Related Macular Degeneration

Ophthalmologist *Re-assessment required after 12 months* All of the following:

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- 1 Documented benefit must be demonstrated to continue; and
- 2 Patient's vision is 6/36 or better on the Snellen visual acuity score; and
- 3 There is no structural damage to the central fovea of the treated eye.

Initiation - Diabetic Macular Oedema

Ophthalmologist

Re-assessment required after 4 months

All of the following:

- 1 Patient has centre involving diabetic macular oedema (DMO); and
- 2 Patient's disease is non responsive to 4 doses of intravitreal bevacizumab when administered 4-6 weekly; and
- 3 Patient has reduced visual acuity between 6/9 6/36 with functional awareness of reduction in vision; and
- 4 Patient has DMO within central OCT (ocular coherence tomography) subfield > 350 micrometers; and
- 5 There is no centre-involving sub-retinal fibrosis or foveal atrophy.

Continuation - Diabetic Macular Oedema

Ophthalmologist

Re-assessment required after 12 months All of the following:

- 1 There is stability or two lines of Snellen visual acuity gain; and
- 2 There is structural improvement on OCT scan (with reduction in intra-retinal cysts, central retinal thickness, and sub-retinal fluid); and
- 3 Patient's vision is 6/36 or better on the Snellen visual acuity score; and
- 4 There is no centre-involving sub-retinal fibrosis or foveal atrophy; and
- 5 After each consecutive 12 months treatment with aflibercept, patient has retrialled with at least one injection of bevacizumab and had no response.

BASILIXIMAB - Restricted see terms below

➡ Restricted (RS1203)

Initiation

For use in solid organ transplants.

BEVACIZUMAB - Restricted see terms below

- Inj 25 mg per ml, 4 ml vial
- Inj 25 mg per ml, 16 ml vial

➡ Restricted (RS1691)

Initiation – Recurrent Respiratory Papillomatosis

Otolaryngologist

Re-assessment required after 12 months All of the following:

- 1 Maximum of 6 doses: and
- 2 The patient has recurrent respiratory papillomatosis; and
- 3 The treatment is for intra-lesional administration.

Continuation – Recurrent Respiratory Papillomatosis

Otolaryngologist

Re-assessment required after 12 months All of the following:

- 1 Maximum of 6 doses; and
- 2 The treatment is for intra-lesional administration; and
- 3 There has been a reduction in surgical treatments or disease regrowth as a result of treatment.

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| continued Initiation – ocular conditions Either: 1 Ocular neovascularisation: or | | | |
| 2 Exudative ocular angiopathy. | | | |
| CETUXIMAB - Restricted see terms below ↓ Inj 5 mg per ml, 20 ml vial ↓ Inj 5 mg per ml, 100 ml vial → Restricted (RS1613) Initiation Medical oncologist All of the following: 1 Patient has locally advanced, non-metastatic, squamous cell 2 Patient is contraindicated to, or is intolerant of, cisplatin; and 3 Patient has good performance status; and 4 To be administered in combination with radiation therapy. | 1,820.00 | 1 1 neck; and | Erbitux Erbitux |
| INFLIXIMAB – Restricted see terms below ↓ Inj 100 mg | | 1 | Remicade |
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- 1 The patient has had an initial Special Authority approval for adalimumab and/or etanercept for rheumatoid arthritis; and 2 Either:
 - 2.1 The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept; or
 - 2.2 Following at least a four month trial of adalimumab and/or etanercept, the patient did not meet the renewal criteria for adalimumab and/or etanercept: and
- 3 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance.

Continuation - rheumatoid arthritis

Rheumatologist *Re-assessment required after 6 months* All of the following:

1 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and

2 Either:

- 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
- 2.2 The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; and
- 3 Infliximab to be administered at doses no greater than 3 mg/kg every 8 weeks.

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Initiation - ankylosing spondylitis

Rheumatologist

Re-assessment required after 3 months

Both:

1 The patient has had an initial Special Authority approval for adalimumab and/or etanercept for ankylosing spondylitis; and 2 Either:

- 2.1 The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept; or
- 2.2 Following 12 weeks of adalimumab and/or etanercept treatment, the patient did not meet the renewal criteria for adalimumab and/or etanercept for ankylosing spondylitis.

Continuation - ankylosing spondylitis

Rheumatologist

Re-assessment required after 6 months

All of the following:

- 1 Following 12 weeks of infliximab treatment, BASDAI has improved by 4 or more points from pre-infliximab baseline on a 10 point scale, or by 50%, whichever is less; and
- 2 Physician considers that the patient has benefited from treatment and that continued treatment is appropriate; and
- 3 Infliximab to be administered at doses no greater than 5 mg/kg every 6-8 weeks.

Initiation – psoriatic arthritis

Rheumatologist

Re-assessment required after 4 months

Both:

- 1 The patient has had an initial Special Authority approval for adalimumab and/or etanercept for psoriatic arthritis; and
- 2 Either:
 - 2.1 The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept; or
 - 2.2 Following 3-4 months' initial treatment with adalimumab and/or etanercept, the patient did not meet the renewal criteria for adalimumab and/or etanercept for psoriatic arthritis.

Continuation - psoriatic arthritis

Rheumatologist

Re-assessment required after 6 months Both:

1 Either:

- 1.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
- 1.2 The patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to prior infliximab treatment in the opinion of the treating physician; and

2 Infliximab to be administered at doses no greater than 5 mg/kg every 8 weeks.

Initiation - severe ocular inflammation

Re-assessment required after 3 doses Either:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for adalimumab for severe ocular inflammation; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for severe ocular inflammation; or

2 Both:

2.1 Patient has severe, vision-threatening ocular inflammation requiring rapid control; and

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- 2.2 Any of the following:
 - 2.2.1 Treatment with high-dose steroids (intravenous methylprednisolone) followed by high dose oral steroids has proven ineffective at controlling symptoms; or
 - 2.2.2 Patient developed new inflammatory symptoms while receiving high dose steroids; or
 - 2.2.3 Patient is aged under 8 years and treatment with high dose oral steroids and other immunosuppressants has proven ineffective at controlling symptoms.

Continuation - severe ocular inflammation

Re-assessment required after 12 months

Any of the following:

- 1 The patient has had a good clinical response following 3 initial doses; or
- 2 Following each 12-month treatment period, the patient has had a sustained reduction in inflammation (Standardisation of Uveitis Nomenclature (SUN) criteria < ½+ anterior chamber or vitreous cells, absence of active vitreous or retinal lesions, or resolution of uveitic cystoid macular oedema); or
- 3 Following each 12-month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 10mg daily, or steroid drops less than twice daily if under 18 years old.

Note: A trial withdrawal should be considered after every 24 months of stability, unless the patient is deemed to have extremely high risk of irreversible vision loss if infliximab is withdrawn.

Initiation - chronic ocular inflammation

Re-assessment required after 3 doses

Either:

- 1 Both:
 - 1.1 The patient has had an initial Special Authority approval for adalimumab for chronic ocular inflammation; and
 - 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab to meet the renewal criteria for adalimumab for chronic ocular inflammation; or

2 Both:

- 2.1 Patient has severe uveitis uncontrolled with treatment of steroids and other immunosuppressants with a severe risk of vision loss; and
- 2.2 Any of the following:
 - 2.2.1 Patient is 18 years or older and treatment with at least two other immunomodulatory agents has proven ineffective; or
 - 2.2.2 Patient is under 18 years and treatment with methotrexate has proven ineffective or is not tolerated at therapeutic dose; or
 - 2.2.3 Patient is under 8 years and treatment with steroids or methotrexate has proven ineffective or is not tolerated at a therapeutic dose; or disease requires control to prevent irreversible vision loss prior to achieving a therapeutic dose of methotrexate.

Continuation - chronic ocular inflammation

Re-assessment required after 12 months

Any of the following:

- 1 The patient has had a good clinical response following 3 initial doses; or
- 2 Following each 12-month treatment period, the patient has had a sustained reduction in inflammation (Standardisation of Uveitis Nomenclature (SUN) criteria < ½+ anterior chamber or vitreous cells, absence of active vitreous or retinal lesions, or resolution of uveitic cystoid macular oedema); or
- 3 Following each 12-month treatment period, the patient has a sustained steroid sparing effect, allowing reduction in prednisone to < 10mg daily, or steroid drops less than twice daily if under 18 years old.
- Note: A trial withdrawal should be considered after every 24 months of stability, unless the patient is deemed to have extremely

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high risk of irreversible vision loss if infliximab is withdrawn.

Initiation – Pulmonary sarcoidosis

Both:

- 1 Patient has life-threatening pulmonary sarcoidosis that is refractory to other treatments; and
- 2 Treatment is to be prescribed by, or has been recommended by, a physician with expertise in the treatment of pulmonary sarcoidosis.

Initiation - Crohn's disease (adults)

Gastroenterologist

Re-assessment required after 3 months

All of the following:

- 1 Patient has severe active Crohn's disease; and
- 2 Any of the following:
 - 2.1 Patient has a Crohn's Disease Activity Index (CDAI) score of greater than or equal to 300; or
 - 2.2 Patient has extensive small intestine disease affecting more than 50 cm of the small intestine; or
 - 2.3 Patient has evidence of short gut syndrome or would be at risk of short gut syndrome with further bowel resection; or
 - 2.4 Patient has an ileostomy or colostomy, and has intestinal inflammation; and
- 3 Patient has tried but had an inadequate response to, or has experienced intolerable side effects from, prior systemic therapy with immunomodulators at maximum tolerated doses (unless contraindicated) and corticosteroids; and
- 4 Surgery (or further surgery) is considered to be clinically inappropriate; and
- 5 Patient must be reassessed for continuation after 3 months of therapy.

Continuation - Crohn's disease (adults)

Gastroenterologist

Re-assessment required after 6 months Both:

- 1 Any of the following:
 - 1.1 CDAI score has reduced by 100 points from the CDAI score when the patient was initiated on infliximab; or
 - 1.2 CDAI score is 150 or less; or
 - 1.3 The patient has demonstrated an adequate response to treatment but CDAI score cannot be assessed; and
- 2 Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle.

Initiation - Crohn's disease (children)

Gastroenterologist

Re-assessment required after 3 months All of the following:

- 1 Paediatric patient has severe active Crohn's disease; and
- 2 Either:

2.1 Patient has a Paediatric Crohn's Disease Activity Index (PCDAI) score of greater than or equal to 30; or

- 2.2 Patient has extensive small intestine disease; and
- 3 Patient has tried but had an inadequate response to, or has experienced intolerable side effects from, prior systemic therapy with immunomodulators at maximum tolerated doses (unless contraindicated) and corticosteroids; and
- 4 Surgery (or further surgery) is considered to be clinically inappropriate; and
- 5 Patient must be reassessed for continuation after 3 months of therapy.

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Continuation - Crohn's disease (children)

Gastroenterologist

Re-assessment required after 6 months Both:

- 1 Any of the following:
 - 1.1 PCDAI score has reduced by 10 points from the PCDAI score when the patient was initiated on infliximab; or
 - 1.2 PCDAI score is 15 or less; or
 - 1.3 The patient has demonstrated an adequate response to treatment but PCDAI score cannot be assessed; and
- 2 Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle.

Initiation - fistulising Crohn's disease

Gastroenterologist

Re-assessment required after 4 months Both:

- 1 Patient has confirmed Crohn's disease; and
- 2 Either:
 - 2.1 Patient has one or more complex externally draining enterocutaneous fistula(e); or
 - 2.2 Patient has one or more rectovaginal fistula(e).

Continuation – fistulising Crohn's disease

Gastroenterologist

Re-assessment required after 6 months

Both:

- 1 Either:
 - 1.1 The number of open draining fistulae have decreased from baseline by at least 50%; or
 - 1.2 There has been a marked reduction in drainage of all fistula(e) from baseline (in the case of adult patients, as demonstrated by a reduction in the Fistula Assessment score), together with less induration and patient reported pain; and
- 2 Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle.

Initiation - acute severe fulminant ulcerative colitis

Gastroenterologist

Limited to 6 weeks treatment

Both:

- 1 Patient has acute, severe fulminant ulcerative colitis; and
- 2 Treatment with intravenous or high dose oral corticosteroids has not been successful.

Continuation - severe fulminant ulcerative colitis

Gastroenterologist

Re-assessment required after 6 months Both:

- 1 Where maintenance treatment is considered appropriate, infliximab should be used in combination with immunomodulators and reassessed every 6 months; and
- 2 Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle.

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Initiation - severe ulcerative colitis

Gastroenterologist

Re-assessment required after 3 months

All of the following:

- 1 Patient has histologically confirmed ulcerative colitis; and
- 2 Either:
 - 2.1 Patient is 18 years or older and the Simple Clinical Colitis Activity Index (SCCAI) is greater than or equal to 4; or
 - 2.2 Patient is under 18 years and the Paediatric Ulcerative Colitis Activity Index (PUCAI) score is greater than or equal to 65; and
- 3 Patient has tried but had an inadequate response to, or has experienced intolerable side effects from, prior systemic therapy with immunomodulators at maximum tolerated doses for an adequate duration (unless contraindicated) and corticosteroids; and
- 4 Surgery (or further surgery) is considered to be clinically inappropriate.

Continuation - severe ulcerative colitis

Gastroenterologist

Re-assessment required after 6 months

All of the following:

- 1 Patient is continuing to maintain remission and the benefit of continuing infliximab outweighs the risks; and
- 2 Either:
 - 2.1 Patient is 18 years or older and the SCCAI score has reduced by 2 points or more from the SCCAI score when the patient was initiated on infliximab; or
 - 2.2 Patient is under 18 years and the PUCAI score has reduced by 30 points or more from the PUCAI score when the patient was initiated on infliximab; and
- 3 Infliximab to be administered at doses up to 5 mg/kg every 8 weeks. Up to 10 mg/kg every 8 weeks (or equivalent) can be used for up to 3 doses if required for secondary non-response to treatment for re-induction. Another re-induction may be considered sixteen weeks after completing the last re-induction cycle.

Initiation - plaque psoriasis

Dermatologist

Re-assessment required after 3 doses Either:

1 Both:

- 1.1 The patient has had an initial Special Authority approval for adalimumab or etanercept for severe chronic plaque psoriasis; and
- 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from adalimumab or etanercept; or
 - 1.2.2 The patient has received insufficient benefit from adalimumab or etanercept to meet the renewal criteria for adalimumab or etanercept for severe chronic plaque psoriasis; or
- 2 All of the following:
 - 2.1 Either:
 - 2.1.1 Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) score of greater than 10, where lesions have been present for at least 6 months from the time of initial diagnosis; or
 - 2.1.2 Patient has severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot, where the plaque or plaques have been present for at least 6 months from the time of initial diagnosis; and
 - 2.2 Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, cyclosporin, or acitretin; and

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- 2.3 A PASI assessment has been completed for at least the most recent prior treatment course (but preferably all prior treatment courses), preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course; and
- 2.4 The most recent PASI assessment is no more than 1 month old at the time of initiation.

Note: "Inadequate response" is defined as: for whole body severe chronic plaque psoriasis, a PASI score of greater than 10, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment; for severe chronic plaque psoriasis of the face, hand or foot, at least 2 of the 3 PASI symptom subscores for erythema, thickness and scaling are rated as severe or very severe, and the skin area affected is 30% or more of the face, palm of a hand or sole of a foot, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment.

Continuation – plaque psoriasis

Dermatologist

Re-assessment required after 3 doses Both:

1 Either:

- 1.1 Both:
 - 1.1.1 Patient had "whole body" severe chronic plaque psoriasis at the start of treatment; and
 - 1.1.2 Following each prior infliximab treatment course the patient has a PASI score which is reduced by 75% or more, or is sustained at this level, when compared with the pre-infliximab treatment baseline value; or
- 1.2 Both:
 - 1.2.1 Patient had severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot at the start of treatment; and
 - 1.2.2 Either:
 - 1.2.2.1 Following each prior infliximab treatment course the patient has a reduction in the PASI symptom subscores for all 3 of erythema, thickness and scaling, to slight or better, or sustained at this level, as compared to the treatment course baseline values; or
 - 1.2.2.2 Following each prior infliximab treatment course the patient has a reduction of 75% or more in the skin area affected, or sustained at this level, as compared to the pre-infliximab treatment baseline value; and
- 2 Infliximab to be administered at doses no greater than 5 mg/kg every 8 weeks.

Initiation – neurosarcoidosis

Neurologist

Re-assessment required after 18 months

All of the following:

- 1 Biopsy consistent with diagnosis of neurosarcoidosis; and
- 2 Patient has CNS involvement; and
- 3 Patient has steroid-refractory disease; and
- 4 Either:
 - 4.1 IV cyclophosphamide has been tried; or
 - 4.2 Treatment with IV cyclophosphamide is clinically inappropriate.

Continuation – neurosarcoidosis

Neurologist

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Re-assessment required after 18 months Either:

- 1 A withdrawal period has been tried and the patient has relapsed; or
- 2 All of the following:
 - 2.1 A withdrawal period has been considered but would not be clinically appropriate; and
 - 2.2 There has been a marked reduction in prednisone dose; and

e.g. Brand indicates brand example only. It is not a contracted product.

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- 2.3 Either:
 - 2.3.1 There has been an improvement in MRI appearances; or
 - 2.3.2 Marked improvement in other symptomology.

Initiation - severe Behcet's disease

Re-assessment required after 4 months

All of the followina:

- 1 The patient has severe Behcet's disease which is significantly impacting the patient's guality of life (see Notes); and 2 Either:
- - 2.1 The patient has severe ocular, neurological and/or vasculitic symptoms and has not responded adequately to one or more treatment(s) appropriate for the particular symptom(s) (see Notes): or
 - 2.2 The patient has severe gastrointestinal, rheumatologic and/or mucocutaneous symptoms and has not responded adequately to two or more treatment appropriate for the particular symptom(s) (see Notes); and
- 3 The patient is experiencing significant loss of quality of life.

Notes:

- 1 Behcet's disease diagnosed according to the International Study Group for Behcet's Disease. Lancet 1990;335(8697):1078-80. Quality of life measured using an appropriate quality of life scale such as that published in Gilworth et al J Rheumatol. 2004;31:931-7.
- 2 Treatments appropriate for the particular symptoms are those that are considered standard conventional treatments for these symptoms, for example intravenous/oral steroids and other immunosuppressants for ocular symptoms; azathioprine, steroids, thalidomide, interferon alpha and ciclosporin for mucocutaneous symptoms; and colchicine, steroids and methotrexate for rheumatological symptoms.

Continuation – severe Behcet's disease

Re-assessment required after 6 months

Both:

- 1 Patient has had a good clinical response to initial treatment with measurably improved quality of life; and
- 2 Infliximab to be administered at doses no greater than 5 mg/kg every 8 weeks.

Initiation – pyoderma gangrenosum

Dermatologist

All of the following:

- 1 Patient has pyoderma gangrenosum*; and
- 2 Patient has received three months of conventional therapy including a minimum of three pharmaceuticals (e.g. prednisone, ciclosporin, azathioprine, or methotrexate) and not received an adequate response; and
- 3 A maximum of 8 doses.

Note: Indications marked with * are unapproved indications.

Continuation – pyoderma gangrenosum

Dermatologist

All of the following:

- 1 Patient has shown clinical improvement; and
- 2 Patient continues to require treatment: and
- 3 A maximum of 8 doses.

MEPOLIZUMAB - Restricted see terms below

| Inj 100 mg vial | .1,638.00 | 1 | Nucala |
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| ➡ Restricted (RS1733) | | | |
| Initiation – Severe eosinophilic asthma | | | |
| Respiratory physician or clinical immunologist | | | |
| Re-assessment required after 12 months | | | |
| All of the following: | | | |

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- 1 Patient must be aged 12 years or older; and
- 2 Patient must have a diagnosis of severe eosinophilic asthma documented by a respiratory physician or clinical immunologist; and
- 3 Conditions that mimic asthma eg. vocal cord dysfunction, central airway obstruction, bronchiolitis etc. have been excluded; and
- 4 Patient has a blood eosinophil count of greater than 0.5×10^{9} cells/L in the last 12 months; and
- 5 Patient must be adherent to optimised asthma therapy including inhaled corticosteroids (equivalent to at least 1000 mcg per day of fluticasone propionate) plus long acting beta-2 agonist, or budesonide/formoterol as part of the single maintenance and reliever therapy regimen, unless contraindicated or not tolerated; and

6 Either:

- 6.1 Patient has had at least 4 exacerbations needing systemic corticosteroids in the previous 12 months, where an exacerbation is defined as either documented use of oral corticosteroids for at least 3 days or parenteral corticosteroids; or
- 6.2 Patient has received continuous oral corticosteroids of at least the equivalent of 10 mg per day over the previous 3 months; and
- 7 Patient has an Asthma Control Test (ACT) score of 10 or less. Baseline measurements of the patient's asthma control using the ACT and oral corticosteroid dose must be made at the time of application, and again at around 52 weeks after the first dose to assess response to treatment.

Continuation – Severe eosinophilic asthma

Respiratory physician or clinical immunologist

Re-assessment required after 2 years

Both:

- 1 An increase in the Asthma Control Test (ACT) score of at least 5 from baseline; and
- 2 Either:
 - 2.1 Exacerbations have been reduced from baseline by 50% as a result of treatment with mepolizumab; or
 - 2.2 Reduction in continuous oral corticosteroid use by 50% or by 10 mg/day while maintaining or improving asthma control.

OBINUTUZUMAB - Restricted see terms below

| t | Inj 25 mg per ml, 40 ml vial | 5,910.00 | 1 | Gazyva |
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| • | Restricted (RS1550) | | | |

Initiation

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Haematologist

Limited to 6 months treatment

All of the following:

- 1 The patient has progressive Binet stage A, B or C CD20+ chronic lymphocytic leukaemia requiring treatment; and
- 2 The patient is obinutuzumab treatment naive; and
- 3 The patient is not eligible for full dose FCR due to comorbidities with a score > 6 on the Cumulative Illness Rating Scale (CIRS) or reduced renal function (creatinine clearance < 70mL/min); and
- 4 Patient has adequate neutrophil and platelet counts* unless the cytopenias are a consequence of marrow infiltration by CLL; and
- 5 Patient has good performance status; and
- 6 Obinutuzumab to be administered at a maximum cumulative dose of 8,000 mg and in combination with chlorambucil for a maximum of 6 cycles.

Notes: Chronic lymphocytic leukaemia includes small lymphocytic lymphoma. Comorbidity refers only to illness/impairment other than CLL induced illness/impairment in the patient. 'Good performance status' means ECOG score of 0-1, however, in patients temporarily debilitated by their CLL disease symptoms a higher ECOG (2 or 3) is acceptable where treatment with obinutuzumab is expected to improve symptoms and improve ECOG score to < 2.

* greater than or equal to 1.5×10^9 /L and platelets greater than or equal to 75×10^9 /L

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| OMALIZUMAB – Restricted see terms below | | | |
| Inj 150 mg prefilled syringe | | 1 | Xolair |
| Inj 150 mg vial | | 1 | Xolair |

⇒ Restricted (RS1652)

Initiation - severe asthma

Clinical immunologist or respiratory specialist

Re-assessment required after 6 months

All of the following:

- 1 Patient must be aged 6 years or older ; and
- 2 Patient has a diagnosis of severe asthma; and
- 3 Past or current evidence of atopy, documented by skin prick testing or RAST; and
- 4 Total serum human immunoglobulin E (IgE) between 76 IU/mL and 1300 IU/ml at baseline; and
- 5 Proven adherence with optimal inhaled therapy including high dose inhaled corticosteroid (budesonide 1,600 mcg per day or fluticasone propionate 1,000 mcg per day or equivalent), plus long-acting beta-2 agonist therapy (at least salmeterol 50 mcg bd or eformoterol 12 mcg bd) for at least 12 months, unless contraindicated or not tolerated; and
- 6 Either:
 - 6.1 Patient has received courses of systemic corticosteroids equivalent to at least 28 days treatment in the past 12 months, unless contraindicated or not tolerated; or
 - 6.2 Patient has had at least 4 exacerbations needing systemic corticosteroids in the previous 12 months, where an exacerbation is defined as either documented use of oral corticosteroids for at least 3 days or parenteral steroids; and
- 7 Patient has an Asthma Control Test (ACT) score of 10 or less; and
- 8 Baseline measurements of the patient's asthma control using the ACT and oral corticosteroid dose must be made at the time of application, and again at around 26 weeks after the first dose to assess response to treatment.

Continuation - severe asthma

Respiratory specialist

Re-assessment required after 6 months

Both:

- 1 An increase in the Asthma Control Test (ACT) score of at least 5 from baseline; and
- 2 A reduction in the maintenance oral corticosteroid dose or number of exacerbations of at least 50% from baseline.

Initiation - severe chronic spontaneous urticaria

Clinical immunologist or dermatologist

Re-assessment required after 6 months

All of the following:

- 1 Patient must be aged 12 years or older; and
- 2 Either:
 - 2.1 Both:
 - 2.1.1 Patient is symptomatic with Urticaria Activity Score 7 (UAS7) of 20 or above; and
 - 2.1.2 Patient has a Dermatology life quality index (DLQI) of 10 or greater; and
- 3 Any of the following:
 - 3.1 Patient has been taking high dose antihistamines (e.g. 4 times standard dose) and ciclosporin (> 3 mg/kg day) for at least 6 weeks; or
 - 3.2 Patient has been taking high dose antihistamines (e.g. 4 times standard dose) and at least 3 courses of systemic corticosteroids (> 20 mg prednisone per day for at least 5 days) in the previous 6 months; or
 - 3.3 Patient has developed significant adverse effects whilst on corticosteroids or ciclosporin; and
- 4 Either:
 - 4.1 Treatment to be stopped if inadequate response* following 4 doses; or
 - 4.2 Complete response* to 6 doses of omalizumab.

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Continuation - severe chronic spontaneous urticaria

Clinical immunologist or dermatologist

Re-assessment required after 6 months

Either:

1 Patient has previously had a complete response* to 6 doses of omalizumab; or

2 Both:

- 2.1 Patient has previously had a complete response* to 6 doses of omalizumab; and
- 2.2 Patient has relapsed after cessation of omalizumab therapy.

Note: *Inadequate response defined as less than 50% reduction in baseline UAS7 and DLQI score, or an increase in Urticaria Control Test (UCT) score of less than 4 from baseline. Patient is to be reassessed for response after 4 doses of omalizumab. Complete response is defined as UAS7 less than or equal to 6 and DLQI less than or equal to 5; or UCT of 16. Relapse of chronic urticaria on stopping prednisone/ciclosporin does not justify the funding of omalizumab.

PERTUZUMAB - Restricted see terms below

➡ Restricted (RS1551)

Initiation

Re-assessment required after 12 months All of the following:

- 1 The patient has metastatic breast cancer expressing HER-2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 Either:
 - 2.1 Patient is chemotherapy treatment naive; or
 - 2.2 Patient has not received prior treatment for their metastatic disease and has had a treatment free interval of at least 12 months between prior (neo)adjuvant chemotherapy treatment and diagnosis of metastatic breast cancer; and
- 3 The patient has good performance status (ECOG grade 0-1); and
- 4 Pertuzumab to be administered in combination with trastuzumab; and
- 5 Pertuzumab maximum first dose of 840 mg, followed by maximum of 420 mg every 3 weeks; and
- 6 Pertuzumab to be discontinued at disease progression.

Continuation

Re-assessment required after 12 months

Both:

- 1 The patient has metastatic breast cancer expressing HER-2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 The cancer has not progressed at any time point during the previous 12 months whilst on pertuzumab and trastuzumab.

RANIBIZUMAB - Restricted see terms below

- Inj 10 mg per ml, 0.23 ml vial
- Inj 10 mg per ml, 0.3 ml vial

→ Restricted (RS1637)

Initiation - Wet Age Related Macular Degeneration

Ophthalmologist

Re-assessment required after 3 months Fither:

itner:

- 1 All of the following:
 - 1.1 Any of the following:
 - 1.1.1 Wet age-related macular degeneration (wet AMD); or
 - 1.1.2 Polypoidal choroidal vasculopathy; or

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- 1.1.3 Choroidal neovascular membrane from causes other than wet AMD; and
- 1.2 Either:
 - 1.2.1 The patient has developed severe endophthalmitis or severe posterior uveitis following treatment with bevacizumab; or
 - 1.2.2 There is worsening of vision or failure of retina to dry despite three intraocular injections of bevacizumab four weeks apart; and
- 1.3 There is no structural damage to the central fovea of the treated eye; and
- 1.4 Patient has not previously been treated with aflibercept for longer than 3 months; or
- 2 Patient has current approval to use aflibercept for treatment of wAMD and was found to be intolerant to aflibercept within 3 months.

Continuation - Wet Age Related Macular Degeneration

Ophthalmologist

Re-assessment required after 12 months

All of the following:

- 1 Documented benefit must be demonstrated to continue; and
- 2 Patient's vision is 6/36 or better on the Snellen visual acuity score; and
- 3 There is no structural damage to the central fovea of the treated eye.

RITUXIMAB (MABTHERA) - Restricted see terms below

| t | Inj 10 mg per ml, 10 ml vial1,075.50 | 2 | Mabthera |
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| t | Inj 10 mg per ml, 50 ml vial2,688.30 | 1 | Mabthera |

→ Restricted (RS1734)

Initiation - haemophilia with inhibitors

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – haemophilia with inhibitors

Haematologist

All of the following:

- 1 Patient was previously treated with rituximab for haemophilia with inhibitors; and
- 2 An initial response lasting at least 12 months was demonstrated; and
- 3 Patient now requires repeat treatment.

Initiation - post-transplant

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – post-transplant

All of the following:

- 1 The patient has had a rituximab treatment-free interval of 12 months or more; and
- 2 The patient has B-cell post-transplant lymphoproliferative disorder*; and
- 3 To be used for no more than 6 treatment cycles.

Note: Indications marked with * are unapproved indications.

Initiation – indolent, low-grade lymphomas or hairy cell leukaemia*

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation - indolent, low-grade lymphomas or hairy cell leukaemia*

Re-assessment required after 9 months

All of the following:

- 1 The patient has had a rituximab treatment-free interval of 12 months or more; and
- 2 The patient has indolent, low-grade NHL or hairy cell leukaemia* with relapsed disease following prior chemotherapy; and
- 3 To be used for no more than 6 treatment cycles.

Note: 'Indolent, low-grade lymphomas' includes follicular, mantle, marginal zone and lymphoplasmacytic/Waldenstrom

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macroglobulinaemia. *Unapproved indication. 'Hairy cell leukaemia' also includes hairy cell leukaemia variant. Initiation – aggressive CD20 positive NHL

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – aggressive CD20 positive NHL

All of the following:

- 1 The patient has had a rituximab treatment-free interval of 12 months or more; and
- 2 The patient has relapsed refractory/aggressive CD20 positive NHL; and
- 3 To be used with a multi-agent chemotherapy regimen given with curative intent; and
- 4 To be used for a maximum of 4 treatment cycles.

Note: 'Aggressive CD20 positive NHL' includes large B-cell lymphoma and Burkitt's lymphoma/leukaemia.

Initiation – Chronic lymphocytic leukaemia

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – Chronic lymphocytic leukaemia

Re-assessment required after 12 months

Both:

- 1 Either:
 - 1.1 The patient's disease has relapsed within 36 months of previous treatment and rituximab treatment is to be used in combination with funded venetoclax; or
 - 1.2 All of the following:
 - 1.2.1 The patient's disease has relapsed following no more than one prior line of treatment with rituximab for CLL; and
 - 1.2.2 The patient has had an interval of 36 months or more since commencement of initial rituximab treatment; and
 - 1.2.3 The patient does not have chromosome 17p deletion CLL; and
 - 1.2.4 It is planned that the patient receives full dose fludarabine and cyclophosphamide (orally or dose equivalent intravenous administration) or bendamustin; and
- 2 Rituximab to be administered in combination with fludarabine and cyclophosphamide, bendamustine or venetoclax for a maximum of 6 treatment cycles.

Note: 'Chronic lymphocytic leukaemia (CLL)' includes small lymphocytic lymphoma. A line of chemotherapy treatment is considered to comprise a known standard therapeutic chemotherapy regimen and supportive treatments.

Initiation - rheumatoid arthritis - prior TNF inhibitor use

Rheumatologist

Limited to 4 months treatment

All of the following:

- 1 Both:
 - 1.1 The patient has had an initial community Special Authority approval for at least one of etanercept and/or adalimumab for rheumatoid arthritis; and
 - 1.2 Either:
 - 1.2.1 The patient has experienced intolerable side effects from a reasonable trial of adalimumab and/or etanercept; or
 - 1.2.2 Following at least a four month trial of adalimumab and/or etanercept, the patient did not meet the renewal criteria for adalimumab and/or etanercept for rheumatoid arthritis; and
- 2 Either:

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- 2.1 Rituximab to be used as an adjunct to methotrexate or leflunomide therapy; or
- 2.2 Patient is contraindicated to both methotrexate and leflunomide, requiring rituximab monotherapy to be used; and
- 3 Maximum of two 1,000 mg infusions of rituximab given two weeks apart.

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Initiation - rheumatoid arthritis - TNF inhibitors contraindicated

Rheumatologist

Limited to 4 months treatment

All of the following:

- 1 Treatment with a Tumour Necrosis Factor alpha inhibitor is contraindicated; and
- 2 Patient has had severe and active erosive rheumatoid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP) antibody positive) for six months duration or longer; and
- 3 Patient has tried and not responded to at least three months of oral or parenteral methotrexate at a dose of at least 20 mg weekly or a maximum tolerated dose; and
- 4 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with sulfasalazine and hydroxychloroquine sulphate (at maximum tolerated doses); and
- 5 Any of the following:
 - 5.1 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with the maximum tolerated dose of cyclosporin; or
 - 5.2 Patient has tried and not responded to at least three months of oral or parenteral methotrexate in combination with intramuscular gold; or
 - 5.3 Patient has tried and not responded to at least three months of therapy at the maximum tolerated dose of leflunomide alone or in combination with oral or parenteral methotrexate; and
- 6 Either:
 - 6.1 Patient has persistent symptoms of poorly controlled and active disease in at least 20 swollen, tender joints; or
 - 6.2 Patient has persistent symptoms of poorly controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
- 7 Either:
 - 7.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 7.2 C-reactive protein levels not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months; and
- 8 Either:
 - 8.1 Rituximab to be used as an adjunct to methotrexate or leflunomide therapy; or
 - 8.2 Patient is contraindicated to both methotrexate and leflunomide, requiring rituximab monotherapy to be used; and
- 9 Maximum of two 1,000 mg infusions of rituximab given two weeks apart.

Continuation - rheumatoid arthritis - re-treatment in 'partial responders' to rituximab

Rheumatologist

Re-assessment required after 4 months

All of the following:

- 1 Any of the following:
 - 1.1 At 4 months following the initial course of rituximab infusions the patient had between a 30% and 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 1.2 At 4 months following the second course of rituximab infusions the patient had at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 1.3 At 4 months following the third and subsequent courses of rituximab infusions, the patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; and
- 2 Rituximab re-treatment not to be given within 6 months of the previous course of treatment; and

3 Either:

- 3.1 Rituximab to be used as an adjunct to methotrexate or leflunomide therapy; or
- 3.2 Patient is contraindicated to both methotrexate and leflunomide, requiring rituximab monotherapy to be used; and

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4 Maximum of two 1,000 mg infusions of rituximab given two weeks apart.

Continuation - rheumatoid arthritis - re-treatment in 'responders' to rituximab

Rheumatologist

Re-assessment required after 4 months All of the following:

- 1 Either:
 - 1.1 At 4 months following the initial course of rituximab infusions the patient had at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
 - 1.2 At 4 months following the second and subsequent courses of rituximab infusions, the patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; and

2 Rituximab re-treatment not to be given within 6 months of the previous course of treatment; and

- 3 Either:
 - 3.1 Rituximab to be used as an adjunct to methotrexate or leflunomide therapy; or
 - 3.2 Patient is contraindicated to both methotrexate and leflunomide, requiring rituximab monotherapy to be used; and
- 4 Maximum of two 1,000 mg infusions of rituximab given two weeks apart.

Initiation - severe cold haemagglutinin disease (CHAD)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – severe cold haemagglutinin disease (CHAD)

Haematologist

Re-assessment required after 8 weeks

Either:

- 1 Previous treatment with lower doses of rituximab (100 mg weekly for 4 weeks) have proven ineffective and treatment with higher doses (375 mg/m² weekly for 4 weeks) is now planned; or
- 2 All of the following:
 - 2.1 Patient was previously treated with rituximab for severe cold haemagglutinin disease*; and
 - 2.2 An initial response lasting at least 12 months was demonstrated; and
 - 2.3 Patient now requires repeat treatment.

Note: Indications marked with * are unapproved indications.

Initiation - warm autoimmune haemolytic anaemia (warm AIHA)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation - warm autoimmune haemolytic anaemia (warm AIHA)

Haematologist

Re-assessment required after 8 weeks Either:

- 1 Previous treatment with lower doses of rituximab (100 mg weekly for 4 weeks) have proven ineffective and treatment with higher doses (375 mg/m² weekly for 4 weeks) is now planned; or
- 2 All of the following:
 - 2.1 Patient was previously treated with rituximab for warm autoimmune haemolytic anaemia*; and
 - 2.2 An initial response lasting at least 12 months was demonstrated; and
 - 2.3 Patient now requires repeat treatment.

Note: Indications marked with * are unapproved indications.

Initiation – immune thrombocytopenic purpura (ITP)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

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Continuation - immune thrombocytopenic purpura (ITP)

Haematologist

Re-assessment required after 8 weeks

Either:

- 1 Previous treatment with lower doses of rituximab (100 mg weekly for 4 weeks) have proven ineffective and treatment with higher doses (375 mg/m² weekly for 4 weeks) is now planned; or
- 2 All of the following:
 - 2.1 Patient was previously treated with rituximab for immune thrombocytopenic purpura*; and
 - 2.2 An initial response lasting at least 12 months was demonstrated; and
 - 2.3 Patient now requires repeat treatment.

Note: Indications marked with * are unapproved indications.

Initiation – thrombotic thrombocytopenic purpura (TTP)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation - thrombotic thrombocytopenic purpura (TTP)

Haematologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient was previously treated with rituximab for thrombotic thrombocytopenic purpura*; and
- 2 An initial response lasting at least 12 months was demonstrated; and
- 3 Patient now requires repeat treatment; and
- 4 The total rituximab dose used would not exceed the equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Initiation – pure red cell aplasia (PRCA)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – pure red cell aplasia (PRCA)

Haematologist

Re-assessment required after 6 weeks

Patient was previously treated with rituximab for pure red cell aplasia* associated with a demonstrable B-cell lymphoproliferative disorder and demonstrated an initial response lasting at least 12 months.

Note: Indications marked with * are unapproved indications.

Initiation – ANCA associated vasculitis

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation - ANCA associated vasculitis

Re-assessment required after 8 weeks

All of the following:

- 1 Patient has been diagnosed with ANCA associated vasculitis*; and
- 2 Patient has previously responded to treatment with rituximab but is now experiencing an acute flare of vasculitis; and
- 3 The total rituximab dose would not exceed the equivalent of 375 mg/m² of body-surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Initiation – treatment refractory systemic lupus erythematosus (SLE)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation - treatment refractory systemic lupus erythematosus (SLE)

Rheumatologist or nephrologist

All of the following:

- 1 Patient's SLE* achieved at least a partial response to the previous round of prior rituximab treatment; and
- 2 The disease has subsequently relapsed; and

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3 Maximum of two 1000 mg infusions of rituximab.

Note: Indications marked with * are unapproved indications.

Initiation – Antibody-mediated renal transplant rejection

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Initiation – ABO-incompatible renal transplant

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Initiation – Steroid dependent nephrotic syndrome (SDNS) or frequently relapsing nephrotic syndrome (FRNS) No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – Steroid dependent nephrotic syndrome (SDNS) or frequently relapsing nephrotic syndrome (FRNS) Nephrologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient who was previously treated with rituximab for nephrotic syndrome*; and
- 2 Treatment with rituximab was previously successful and has demonstrated sustained response for > 6 months, but the condition has relapsed and the patient now requires repeat treatment; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.
- Note: Indications marked with a * are unapproved indications.

Initiation - Steroid resistant nephrotic syndrome (SRNS)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – Steroid resistant nephrotic syndrome (SRNS)

Nephrologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient who was previously treated with rituximab for nephrotic syndrome*; and
- 2 Treatment with rituximab was previously successful and has demonstrated sustained response for greater than 6 months, but the condition has relapsed and the patient now requires repeat treatment; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.
- Note: Indications marked with a * are unapproved indications.

Initiation – Neuromyelitis Optica Spectrum Disorder (NMOSD)

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation - Neuromyelitis Optica Spectrum Disorder (NMOSD)

Relevant specialist or medical practitioner on the recommendation of a Relevant specialist

Re-assessment required after 2 years

All of the following:

- 1 One of the following dose regimens is to be used: 2 doses of 1,000 mg rituximab administered fortnightly, or 4 doses of 375 mg/m2 administered weekly for four weeks; and
- 2 The patients has responded to the most recent course of rituximab; and
- 3 The patient has not received rituximab in the previous 6 months.

Initiation – Severe Refractory Myasthenia Gravis

No new patient can start on rituximab (Mabthera brand) under this Initiation criteria from 1 March 2020.

Continuation – Severe Refractory Myasthenia Gravis

Neurologist or medical practitioner on the recommendation of a Neurologist

Re-assessment required after 2 years

All of the following:

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1 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart; and

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| 2 An initial response lasting at least 12 months was demonstra | ited; and | | | |
| 3 Either: | | | | |
| 3.1 The patient has relapsed despite treatment with cortic pacied of at least 12 months, or | costeroids and at least | one other | immunosuppressant for a | |
| period of at least 12 months; or 3.2 Both: | | | | |
| 3.2.1 The patient's myasthenia gravis has relapsed | despite treatment with | at least or | e immunosuppressant for a | |
| period of at least 12 months; and | | | | |
| 3.2.2 Corticosteroids have been trialed for at least 1 side effects. | 2 months and have be | en discont | inued due to unacceptable | |
| RITUXIMAB (RIXIMYO) – Restricted see terms below | | | | |
| Inj 10 mg per ml, 10 ml vial | | 2 | Riximyo | |
| Inj 10 mg per ml, 50 ml vial | | 1 | Riximyo | |
| → Restricted (RS1764) | | | | |
| Initiation – haemophilia with inhibitors Haematologist | | | | |
| Any of the following: | | | | |
| 1 Patient has mild congenital haemophilia complicated by inhib | pitors; or | | | |
| 2 Patient has severe congenital haemophilia complicated by in | hibitors and has failed | immune to | plerance therapy; or | |
| 3 Patient has acquired haemophilia. | | | | |
| Continuation – haemophilia with inhibitors | | | | |
| Haematologist | | | | |
| All of the following: 1 Patient was previously treated with rituximab for haemophilia | with inhibitors, and | | | |
| 2 An initial response lasting at least 12 months was demonstra | | | | |
| 3 Patient now requires repeat treatment. | alou, and | | | |
| Initiation – post-transplant | | | | |
| Both: | | | | |
| 1 The patient has B-cell post-transplant lymphoproliferative dis | order*; and | | | |
| 2 To be used for a maximum of 8 treatment cycles. | | | | |
| Note: Indications marked with * are unapproved indications. | | | | |
| Continuation – post-transplant All of the following: | | | | |
| 1 The patient has had a rituximab treatment-free interval of 12 | months or more: and | | | |
| 2 The patient has B-cell post-transplant lymphoproliferative dis | , | | | |
| 3 To be used for no more than 6 treatment cycles. | | | | |
| Note: Indications marked with * are unapproved indications. | | | | |
| Initiation - indolent, low-grade lymphomas or hairy cell leukaer | nia* | | | |
| Re-assessment required after 9 months | | | | |
| Either: | | | | |
| Both: 1.1 The patient has indolent low grade NHL or hairy cell I | oukaomia* with relance | d discoso | following prior | |
| chemotherapy; and | euraenna wiin relapse | u uisedst | | |
| 1.0. To be used for a maximum of 6 treatment avalage or | | | | |

- 1.2 To be used for a maximum of 6 treatment cycles; or
- 2 Both:
 - 2.1 The patient has indolent, low grade lymphoma or hairy cell leukaemia* requiring first-line systemic chemotherapy; and
 - 2.2 To be used for a maximum of 6 treatment cycles.

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Note: 'Indolent, low-grade lymphomas' includes follicular, mantle, marginal zone and lymphoplasmacytic/Waldenstrom macroglobulinaemia. *Unapproved indication. 'Hairy cell leukaemia' also includes hairy cell leukaemia variant.

Continuation - indolent, low-grade lymphomas or hairy cell leukaemia*

Re-assessment required after 12 months

All of the following:

- 1 The patient has had a rituximab treatment-free interval of 12 months or more; and
- 2 The patient has indolent, low-grade NHL or hairy cell leukaemia* with relapsed disease following prior chemotherapy; and
- 3 To be used for no more than 6 treatment cycles.

Note: 'Indolent, low-grade lymphomas' includes follicular, mantle, marginal zone and lymphoplasmacytic/Waldenstrom macroglobulinaemia. *Unapproved indication. 'Hairy cell leukaemia' also includes hairy cell leukaemia variant. Initiation – aggressive CD20 positive NHL

Either:

- 1 All of the following:
 - 1.1 The patient has treatment naive aggressive CD20 positive NHL; and
 - 1.2 To be used with a multi-agent chemotherapy regimen given with curative intent; and
 - 1.3 To be used for a maximum of 8 treatment cycles; or

2 Both:

- 2.1 The patient has aggressive CD20 positive NHL with relapsed disease following prior chemotherapy; and
- 2.2 To be used for a maximum of 6 treatment cycles.
- Note: 'Aggressive CD20 positive NHL' includes large B-cell lymphoma and Burkitt's lymphoma/leukaemia.

Continuation - aggressive CD20 positive NHL

All of the following:

- 1 The patient has had a rituximab treatment-free interval of 12 months or more; and
- 2 The patient has relapsed refractory/aggressive CD20 positive NHL; and
- 3 To be used with a multi-agent chemotherapy regimen given with curative intent; and
- 4 To be used for a maximum of 4 treatment cycles.

Note: 'Aggressive CD20 positive NHL' includes large B-cell lymphoma and Burkitt's lymphoma/leukaemia.

Initiation – Chronic lymphocytic leukaemia

Re-assessment required after 12 months

All of the following:

- 1 The patient has progressive Binet stage A, B or C chronic lymphocytic leukaemia (CLL) requiring treatment; and
- 2 Any of the following:
 - 2.1 The patient is rituximab treatment naive; or
 - 2.2 Either:
 - 2.2.1 The patient is chemotherapy treatment naive; or

2.2.2 Both:

- 2.2.2.1 The patient's disease has relapsed following no more than three prior lines of chemotherapy treatment; and
- 2.2.2.2 The patient has had a treatment-free interval of 12 months or more if previously treated with fludarabine and cyclophosphamide chemotherapy; or
- 2.3 The patient's disease has relapsed within 36 months of previous treatment and rituximab treatment is to be used in combination with funded venetoclax; and
- 3 The patient has good performance status; and
- 4 Either:

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- 4.1 The patient does not have chromosome 17p deletion CLL; or
- 4.2 Rituximab treatment is to be used in combination with funded venetoclax for relapsed/refractory chronic lymphocytic leukaemia; and

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- 5 Rituximab to be administered in combination with fludarabine and cyclophosphamide, bendamustine or venetoclax for a maximum of 6 treatment cycles; and
- 6 It is planned that the patient receives full dose fludarabine and cyclophosphamide (orally or dose equivalent intravenous administration), bendamustine or venetoclax.

Note: 'Chronic lymphocytic leukaemia (CLL)' includes small lymphocytic lymphoma. A line of chemotherapy treatment is considered to comprise a known standard therapeutic chemotherapy regimen and supportive treatments. 'Good performance status' means ECOG score of 0-1, however, in patients temporarily debilitated by their CLL disease symptoms a higher ECOG (2 or 3) is acceptable where treatment with rituximab is expected to improve symptoms and improve ECOG score to < 2.

Continuation – Chronic lymphocytic leukaemia

Re-assessment required after 12 months Both:

1 Either:

- 1.1 The patient's disease has relapsed within 36 months of previous treatment and rituximab treatment is to be used in combination with funded venetoclax; or
- 1.2 All of the following:
 - 1.2.1 The patient's disease has relapsed following no more than one prior line of treatment with rituximab for CLL; and
 - 1.2.2 The patient has had an interval of 36 months or more since commencement of initial rituximab treatment; and
 - 1.2.3 The patient does not have chromosome 17p deletion CLL; and
 - 1.2.4 It is planned that the patient receives full dose fludarabine and cyclophosphamide (orally or dose equivalent intravenous administration) or bendamustin; and
- 2 Rituximab to be administered in combination with fludarabine and cyclophosphamide, bendamustine or venetoclax for a maximum of 6 treatment cycles.

Note: 'Chronic lymphocytic leukaemia (CLL)' includes small lymphocytic lymphoma. A line of chemotherapy treatment is considered to comprise a known standard therapeutic chemotherapy regimen and supportive treatments.

Initiation - severe cold haemagglutinin disease (CHAD)

Haematologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient has cold haemagglutinin disease*; and
- 2 Patient has severe disease which is characterized by symptomatic anaemia, transfusion dependence or disabling circulatory symptoms; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Continuation – severe cold haemagglutinin disease (CHAD)

Haematologist

Re-assessment required after 8 weeks

Either:

1 Previous treatment with lower doses of rituximab (100 mg weekly for 4 weeks) have proven ineffective and treatment with higher doses (375 mg/m² weekly for 4 weeks) is now planned; or

2 All of the following:

- 2.1 Patient was previously treated with rituximab for severe cold haemagglutinin disease*; and
- 2.2 An initial response lasting at least 12 months was demonstrated; and
- 2.3 Patient now requires repeat treatment.

Note: Indications marked with * are unapproved indications.

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Initiation – warm autoimmune haemolytic anaemia (warm AIHA)

Haematologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient has warm autoimmune haemolytic anaemia*; and
- 2 One of the following treatments has been ineffective: steroids (including if patient requires ongoing steroids at doses equivalent to > 5 mg prednisone daily), cytotoxic agents (e.g. cyclophosphamide monotherapy or in combination), intravenous immunoglobulin; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Continuation - warm autoimmune haemolytic anaemia (warm AIHA)

Haematologist

Re-assessment required after 8 weeks

Either:

- 1 Previous treatment with lower doses of rituximab (100 mg weekly for 4 weeks) have proven ineffective and treatment with higher doses (375 mg/m² weekly for 4 weeks) is now planned; or
- 2 All of the following:
 - 2.1 Patient was previously treated with rituximab for warm autoimmune haemolytic anaemia*; and
 - 2.2 An initial response lasting at least 12 months was demonstrated; and
 - 2.3 Patient now requires repeat treatment.
- Note: Indications marked with * are unapproved indications.

Initiation – immune thrombocytopenic purpura (ITP)

Haematologist

Re-assessment required after 8 weeks

All of the following:

1 Either:

- 1.1 Patient has immune thrombocytopenic purpura* with a platelet count of less than or equal to 20,000 platelets per microlitre; or
- 1.2 Patient has immune thrombocytopenic purpura* with a platelet count of 20,000 to 30,000 platelets per microlitre and significant mucocutaneous bleeding; and
- 2 Any of the following:
 - 2.1 Treatment with steroids and splenectomy have been ineffective; or
 - 2.2 Treatment with steroids has been ineffective and splenectomy is an absolute contraindication; or
 - 2.3 Other treatments including steroids have been ineffective and patient is being prepared for elective surgery (e.g. splenectomy); and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Continuation - immune thrombocytopenic purpura (ITP)

Haematologist

Re-assessment required after 8 weeks

Either:

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- 1 Previous treatment with lower doses of rituximab (100 mg weekly for 4 weeks) have proven ineffective and treatment with higher doses (375 mg/m² weekly for 4 weeks) is now planned; or
- 2 All of the following:
 - 2.1 Patient was previously treated with rituximab for immune thrombocytopenic purpura*; and

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- 2.2 An initial response lasting at least 12 months was demonstrated; and
- 2.3 Patient now requires repeat treatment.
- Note: Indications marked with * are unapproved indications.
- Initiation thrombotic thrombocytopenic purpura (TTP)

Haematologist

Re-assessment required after 8 weeks

Both:

- 1 The total rituximab dose used would not exceed the equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks; and
- 2 Either:
 - 2.1 Patient has thrombotic thrombocytopenic purpura* and has experienced progression of clinical symptoms or persistent thrombocytopenia despite plasma exchange; or
 - 2.2 Patient has acute idiopathic thrombotic thrombocytopenic purpura* with neurological or cardiovascular pathology.

Note: Indications marked with * are unapproved indications.

Continuation – thrombotic thrombocytopenic purpura (TTP)

Haematologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient was previously treated with rituximab for thrombotic thrombocytopenic purpura*; and
- 2 An initial response lasting at least 12 months was demonstrated; and
- 3 Patient now requires repeat treatment; and
- 4 The total rituximab dose used would not exceed the equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Initiation – pure red cell aplasia (PRCA)

Haematologist

Re-assessment required after 6 weeks

Patient has autoimmune pure red cell aplasia* associated with a demonstrable B-cell lymphoproliferative disorder.

Note: Indications marked with * are unapproved indications.

Continuation - pure red cell aplasia (PRCA)

Haematologist

Re-assessment required after 6 weeks

Patient was previously treated with rituximab for pure red cell aplasia* associated with a demonstrable B-cell lymphoproliferative disorder and demonstrated an initial response lasting at least 12 months.

Note: Indications marked with * are unapproved indications.

Initiation – ANCA associated vasculitis

Re-assessment required after 8 weeks

All of the following:

- 1 Patient has been diagnosed with ANCA associated vasculitis*; and
- 2 The total rituximab dose would not exceed the equivalent of 375 mg/m² of body-surface area per week for a total of 4 weeks; and
- 3 Any of the following:
 - 3.1 Induction therapy with daily oral or pulse intravenous cyclophosphamide has failed to achieve significant improvement of disease after at least 3 months; or
 - 3.2 Patient has previously had a cumulative dose of cyclophosphamide > 15 g or a further repeat 3 month induction course of cyclophosphamide would result in a cumulative dose > 15 g; or
 - 3.3 Cyclophosphamide and methotrexate are contraindicated; or

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- 3.4 Patient is a female of child-bearing potential; or
- 3.5 Patient has a previous history of haemorrhagic cystitis, urological malignancy or haematological malignancy.

Note: Indications marked with * are unapproved indications.

Continuation – ANCA associated vasculitis

Re-assessment required after 8 weeks

All of the following:

- 1 Patient has been diagnosed with ANCA associated vasculitis*; and
- 2 Patient has previously responded to treatment with rituximab but is now experiencing an acute flare of vasculitis; and
- 3 The total rituximab dose would not exceed the equivalent of 375 mg/m² of body-surface area per week for a total of 4 weeks.

Note: Indications marked with * are unapproved indications.

Initiation - treatment refractory systemic lupus erythematosus (SLE)

Rheumatologist or nephrologist

All of the following:

- 1 The patient has severe, immediately life- or organ-threatening SLE*; and
- 2 The disease has proved refractory to treatment with steroids at a dose of at least 1 mg/kg; and
- 3 The disease has relapsed following prior treatment for at least 6 months with maximal tolerated doses of azathioprine, mycophenolate mofetil and high dose cyclophosphamide, or cyclophosphamide is contraindicated; and
- 4 Maximum of four 1000 mg infusions of rituximab.

Note: Indications marked with * are unapproved indications.

Continuation - treatment refractory systemic lupus erythematosus (SLE)

Rheumatologist or nephrologist

All of the following:

- 1 Patient's SLE* achieved at least a partial response to the previous round of prior rituximab treatment; and
- 2 The disease has subsequently relapsed; and
- 3 Maximum of two 1000 mg infusions of rituximab.

Note: Indications marked with * are unapproved indications.

Initiation - Antibody-mediated organ transplant rejection

Patient has been diagnosed with antibody-mediated organ transplant rejection*.

Note: Indications marked with * are unapproved indications.

Initiation – ABO-incompatible organ transplant

Patient is to undergo an ABO-incompatible solid organ transplant*.

Note: Indications marked with * are unapproved indications.

Initiation – Steroid dependent nephrotic syndrome (SDNS) or frequently relapsing nephrotic syndrome (FRNS) Nephrologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient is a child with SDNS* or FRNS*; and
- 2 Treatment with steroids for at least a period of 3 months has been ineffective or associated with evidence of steroid toxicity; and
- 3 Treatment with ciclosporin for at least a period of 3 months has been ineffective and/or discontinued due to unacceptable side effects; and
- 4 Treatment with mycophenolate for at least a period of 3 months with no reduction in disease relapses; and
- 5 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.

Note: Indications marked with a * are unapproved indications.

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Continuation – Steroid dependent nephrotic syndrome (SDNS) or frequently relapsing nephrotic syndrome (FRNS) Nephrologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient who was previously treated with rituximab for nephrotic syndrome*; and
- 2 Treatment with rituximab was previously successful and has demonstrated sustained response for > 6 months, but the condition has relapsed and the patient now requires repeat treatment; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.

Note: Indications marked with a * are unapproved indications.

Initiation – Steroid resistant nephrotic syndrome (SRNS)

Nephrologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient is a child with SRNS* where treatment with steroids and ciclosporin for at least 3 months have been ineffective; and
- 2 Treatment with tacrolimus for at least 3 months has been ineffective; and
- 3 Genetic causes of nephrotic syndrome have been excluded; and
- 4 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.

Note: Indications marked with a * are unapproved indications.

Continuation – Steroid resistant nephrotic syndrome (SRNS)

Nephrologist

Re-assessment required after 8 weeks

All of the following:

- 1 Patient who was previously treated with rituximab for nephrotic syndrome*; and
- 2 Treatment with rituximab was previously successful and has demonstrated sustained response for greater than 6 months, but the condition has relapsed and the patient now requires repeat treatment; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.
- Note: Indications marked with a * are unapproved indications.

Initiation - Neuromyelitis Optica Spectrum Disorder (NMOSD)

Re-assessment required after 6 months

Both:

- 1 One of the following dose regimens is to be used: 2 doses of 1,000 mg rituximab administered fortnightly, or 4 doses of 375 mg/m2 administered weekly for four weeks; and
- 2 Either:
 - 2.1 The patient has experienced a severe episode or attack of NMOSD (rapidly progressing symptoms and clinical investigations supportive of a severe attack of NMOSD); or
 - 2.2 All of the following:
 - 2.2.1 The patient has experienced a breakthrough attack of NMOSD; and
 - 2.2.2 The patient is receiving treatment with mycophenolate; and
 - 2.2.3 The patients is receiving treatment with corticosteroids.

Continuation - Neuromyelitis Optica Spectrum Disorder (NMOSD)

Re-assessment required after 2 years

All of the following:

1 One of the following dose regimens is to be used: 2 doses of 1,000 mg rituximab administered fortnightly, or 4 doses of

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- 375 mg/m2 administered weekly for four weeks; and
- 2 The patients has responded to the most recent course of rituximab; and
- 3 The patient has not received rituximab in the previous 6 months.

Initiation - Severe Refractory Myasthenia Gravis

Neurologist

Re-assessment required after 2 years

Both:

- 1 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart; and
- 2 Either:
 - 2.1 Treatment with corticosteroids and at least one other immunosuppressant for at least a period of 12 months has been ineffective; or
 - 2.2 Both:
 - 2.2.1 Treatment with at least one other immunosuppressant for a period of at least 12 months; and
 - 2.2.2 Corticosteroids have been trialed for at least 12 months and have been discontinued due to unacceptable side effects.

Continuation - Severe Refractory Myasthenia Gravis

Neurologist

Re-assessment required after 2 years

All of the following:

- 1 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart; and
- 2 An initial response lasting at least 12 months was demonstrated; and
- 3 Either:
 - 3.1 The patient has relapsed despite treatment with corticosteroids and at least one other immunosuppressant for a period of at least 12 months; or
 - 3.2 Both:
 - 3.2.1 The patient's myasthenia gravis has relapsed despite treatment with at least one immunosuppressant for a period of at least 12 months; and
 - 3.2.2 Corticosteroids have been trialed for at least 12 months and have been discontinued due to unacceptable side effects.

Initiation - Severe antisynthetase syndrome

Re-assessment required after 12 months

All of the following:

- 1 Patient has confirmed antisynthetase syndrome; and
- 2 Patient has severe, immediately life or organ threatening disease, including interstitial lung disease; and
- 3 Either:
 - 3.1 Treatment with at least 3 immunosuppressants (oral steroids, cyclophosphamide, methotrexate, mycophenolate, ciclosporin, azathioprine) has not be effective at controlling active disease; or
 - 3.2 Rapid treatment is required due to life threatening complications; and

4 Maximum of four 1,000 mg infusions of rituximab.

Continuation - Severe antisynthetase syndrome

Re-assessment required after 12 months

All of the following:

- 1 Patient's disease has responded to the previous rituximab treatment with demonstrated improvement in inflammatory markers, muscle strength and pulmonary function; and
- 2 The patient has not received rituximab in the previous 6 months; and

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3 Maximum of two cycles of 2 × 1,000 mg infusions of rituximab given two weeks apart.

Initiation – graft versus host disease

All of the following:

- 1 Patient has refractory graft versus host disease following transplant; and
- 2 Treatment with at least 3 immunosuppressants (oral steroids, ciclosporin, tacrolimus, mycophenolate, sirolimus) has not be effective at controlling active disease; and
- 3 The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks.

Initiation - severe chronic inflammatory demyelinating polyneuropathy

Neurologist

Re-assessment required after 6 months

All of the following:

1 Patient has severe chronic inflammatory demyelinating polyneuropathy (CIPD); and

2 Either:

- 2.1 Both:
 - 2.1.1 Treatment with steroids and intravenous immunoglobulin and/or plasma exchange has not been effective at controlling active disease; and
 - 2.1.2 At least one other immunosuppressant (cyclophosphamide, ciclosporin, tacrolimus, mycophenolate) has not been effective at controlling active disease; or
- 2.2 Rapid treatment is required due to life threatening complications; and
- 3 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart.

Continuation - severe chronic inflammatory demyelinating polyneuropathy

Neurologist or medical practitioner on the recommendation of a Neurologist

Re-assessment required after 6 months

All of the following:

- 1 Patient's disease has responded to the previous rituximab treatment with demonstrated improvement in neurological function compared to baseline; and
- 2 The patient has not received rituximab in the previous 6 months; and
- 3 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart.

Initiation – anti-NMDA receptor autoimmune encephalitis

Neurologist

Re-assessment required after 6 months All of the following:

- 1 Patient has severe anti-NMDA receptor autoimmune encephalitis; and
- 2 Either:
 - 2.1 Both:
 - 2.1.1 Treatment with steroids and intravenous immunoglobulin and/or plasma exchange has not been effective at controlling active disease; and
 - 2.1.2 At least one other immunosuppressant (cyclophosphamide, ciclosporin, tacrolimus, mycophenolate) has not been effective at controlling active disease; or
 - 2.2 Rapid treatment is required due to life threatening complications; and
- 3 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart.

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Continuation - anti-NMDA receptor autoimmune encephalitis

Neurologist

Re-assessment required after 6 months

All of the following:

- 1 Patient's disease has responded to the previous rituximab treatment with demonstrated improvement in neurological function; and
- 2 The patient has not received rituximab in the previous 6 months; and
- 3 The patient has experienced a relapse and now requires further treatment; and
- 4 One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart.

Initiation - CD20+ low grade or follicular B-cell NHL

Re-assessment required after 9 months

Either:

- 1 Both:
 - 1.1 The patient has CD20+ low grade or follicular B-cell NHL with relapsed disease following prior chemotherapy; and
 - 1.2 To be used for a maximum of 6 treatment cycles; or
- 2 Both:
 - 2.1 The patient has CD20+ low grade or follicular B-cell NHL requiring first-line systemic chemotherapy; and
 - 2.2 To be used for a maximum of 6 treatment cycles.

Continuation - CD20+ low grade or follicular B-cell NHL

Re-assessment required after 24 months

Both:

ſ

- 1 Rituximab is to be used for maintenance in CD20+ low grade or follicular B-cell NHL following induction with first-line systemic chemotherapy; and
- 2 Patient is intended to receive rituximab maintenance therapy for 2 years at a dose of 375 mg/m2 every 8 weeks (maximum of 12 cycles).

SECUKINUMAB - Restricted see terms below

→ Restricted (RS1653)

Initiation - severe chronic plaque psoriasis, second-line biologic

Dermatologist

Re-assessment required after 4 months

All of the following:

- 1 The patient has had an initial Special Authority approval for adalimumab or etanercept, or has trialled infliximab in a DHB hospital in accordance with the General Rules of the Pharmaceutical Schedule, for severe chronic plaque psoriasis; and 2 Either.
- 2 Either:
 - 2.1 The patient has experienced intolerable side effects from adalimumab, etanercept or infliximab; or
 - 2.2 The patient has received insufficient benefit from adalimumab, etanercept or infliximab; and
- 3 A Psoriasis Area and Severity Index (PASI) assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the most recent prior treatment course, preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course; and
- 4 The most recent PASI or DQLI assessment is no more than 1 month old at the time of application.

Continuation - severe chronic plaque psoriasis, second-line biologic

Dermatologist *Re-assessment required after 6 months* Both:

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| \$ 1 | Per Manufacturer | |

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- 1 Either:
 - 1.1 Patient's PASI score has reduced by 75% or more (PASI 75) as compared to baseline PASI prior to commencing secukinumab; or
 - 1.2 Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing secukinumab; and
- 2 Secukinumab to be administered at a maximum dose of 300 mg monthly.

Initiation - severe chronic plaque psoriasis, first-line biologic

Dermatologist

Re-assessment required after 4 months

All of the following:

1 Either:

- 1.1 Patient has "whole body" severe chronic plaque psoriasis with a Psoriasis Area and Severity Index (PASI) score of greater than 10, where lesions have been present for at least 6 months from the time of initial diagnosis; or
- 1.2 Patient has severe chronic plaque psoriasis of the face, or palm of a hand or sole of a foot, where the plaque or plaques have been present for at least 6 months from the time of initial diagnosis; and
- 2 Patient has tried, but had an inadequate response (see Note) to, or has experienced intolerable side effects from, at least three of the following (at maximum tolerated doses unless contraindicated): phototherapy, methotrexate, ciclosporin, or acitretin; and
- 3 A PASI assessment or Dermatology Quality of Life Index (DLQI) assessment has been completed for at least the most recent prior treatment course, preferably while still on treatment but no longer than 1 month following cessation of each prior treatment course; and
- 4 The most recent PASI or DQLI assessment is no more than 1 month old at the time of application.

Note: A treatment course is defined as a minimum of 12 weeks of treatment. "Inadequate response" is defined as: for whole body severe chronic plaque psoriasis, a PASI score of greater than 10, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment; for severe chronic plaque psoriasis of the face, hand or foot, at least 2 of the 3 PASI symptom sub scores for erythema, thickness and scaling are rated as severe or very severe, and the skin area affected is 30% or more of the face, palm of a hand or sole of a foot, as assessed preferably while still on treatment but no longer than 1 month following cessation of the most recent prior treatment.

Continuation - severe chronic plaque psoriasis, first-line biologic

Dermatologist

Re-assessment required after 6 months Both:

1 Fither

- 1.1 Patient's PASI score has reduced by 75% or more (PASI 75) as compared to baseline PASI prior to commencing secukinumab; or
- 1.2 Patient has a Dermatology Quality of Life Index (DLQI) improvement of 5 or more, as compared to baseline DLQI prior to commencing secukinumab; and
- 2 Secukinumab to be administered at a maximum dose of 300 mg monthly.

SILTUXIMAB - Restricted see terms below

| t | Inj 100 mg vial | .770.57 | 1 | Sylvant |
|---|-----------------|---------|---|---------|
| t | Inj 400 mg vial | ,082.33 | 1 | Sylvant |

→ Restricted (RS1525)

Initiation

Haematologist or rheumatologist *Re-assessment required after 6 months* All of the following:

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continued...

- 1 Patient has severe HHV-8 negative idiopathic multicentric Castleman's Disease; and
- 2 Treatment with an adequate trial of corticosteroids has proven ineffective; and
- 3 Siltuximab is to be administered at doses no greater than 11 mg/kg every 3 weeks.

Continuation

Haematologist or rheumatologist

Re-assessment required after 12 months

The treatment remains appropriate and the patient has sustained improvement in inflammatory markers and functional status.

TOCILIZUMAB - Restricted see terms below

| t | Inj 20 mg per ml, 4 ml vial220.00 | 1 | Actemra |
|---|--------------------------------------|---|---------|
| t | Inj 20 mg per ml, 10 ml vial550.00 | 1 | Actemra |
| t | Inj 20 mg per ml, 20 ml vial1,100.00 | 1 | Actemra |

→ Restricted (RS1710)

Initiation - cytokine release syndrome

Therapy limited to 3 doses

Either:

- 1 All of the following:
 - 1.1 The patient is enrolled in the Children's Oncology Group AALL1731 trial; and
 - 1.2 The patient has developed grade 3 or 4 cytokine release syndrome associated with the administration of blinatumomab for the treatment of acute lymphoblastic leukaemia; and
 - Tocilizumab is to be administered at doses no greater than 8 mg/kg IV for a maximum of 3 doses (if less than 30kg, maximum of 12 mg/kg); or
- 2 All of the following:
 - 2.1 The patient is enrolled in the Malaghan Institute of Medical Research Phase I ENABLE trial; and
 - 2.2 The patient has developed CRS or CAR T-Cell Related Encephalopathy Syndrome (CRES) associated with the administration of CAR T-cell therapy for the treatment of relapsed or refractory B-cell non-Hodgkin lymphoma; and
 - 2.3 Tocilizumab is to be administered according to the consensus guidelines for CRS and CRES for CAR T-cell therapy (Neelapu et al. Nat Rev Clin Oncol 2018;15:47-62) at doses no greater than 8 mg/kg IV for a maximum of 3 doses.

Initiation - previous use

Any relevant practitioner

Limited to 6 months treatment

Both:

- 1 Patient was being treated with tocilizumab prior to 1 February 2019; and
- 2 Any of the following:
 - 2.1 rheumatoid arthritis; or
 - 2.2 systemic juvenile idiopathic arthritis; or
 - 2.3 adult-onset Still's disease; or
 - 2.4 polyarticular juvenile idiopathic arthritis; or
 - 2.5 idiopathic multicentric Castleman's disease.

Initiation - Rheumatoid Arthritis (patients previously treated with adalimumab or etanercept)

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Limited to 6 months treatment

All of the following:

- 1 The patient has had an initial Special Authority approval for adalimumab and/or etanercept for rheumatoid arthritis; and 2 Either:
 - 2.1 The patient has experienced intolerable side effects from adalimumab and/or etanercept; or
 - 2.2 The patient has received insufficient benefit from at least a three-month trial of adalimumab and/or etanercept such that they do not meet the renewal criteria for rheumatoid arthritis; and

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- 3 Either:
 - 3.1 The patient is seronegative for both anti-cyclic citrullinated peptide (CCP) antibodies and rheumatoid factor; or 3.2 Both:
 - 3.2.1 The patient has been started on rituximab for rheumatoid arthritis in a DHB hospital in accordance with the Section H rules; and
 - 3.2.2 Either:
 - 3.2.2.1 The patient has experienced intolerable side effects from rituximab; or
 - 3.2.2.2 At four months following the initial course of rituximab the patient has received insufficient benefit such that they do not meet the renewal criteria for rheumatoid arthritis.

Initiation - Rheumatoid Arthritis

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has had severe and active erosive rheumatoid arthritis (either confirmed by radiology imaging, or the patient is cyclic citrullinated peptide (CCP) antibody positive) for six months duration or longer; and
- 2 Tocilizumab is to be used as monotherapy; and
- 3 Either:
 - 3.1 Treatment with methotrexate is contraindicated; or
 - 3.2 Patient has tried and did not tolerate oral and/or parenteral methotrexate; and
- 4 Either:
 - 4.1 Patient has tried and not responded to at least three months therapy at the maximum tolerated dose of ciclosporin alone or in combination with another agent; or
 - 4.2 Patient has tried and not responded to at least three months therapy at the maximum tolerated dose of leflunomide alone or in combination with another agent; and
- 5 Either:
 - 5.1 Patient has persistent symptoms of poorly controlled and active disease in at least 20 active, swollen, tender joints; or
 - 5.2 Patient has persistent symptoms of poorly controlled and active disease in at least four active joints from the following: wrist, elbow, knee, ankle, and either shoulder or hip; and
- 6 Either:
 - 6.1 Patient has a C-reactive protein level greater than 15 mg/L measured no more than one month prior to the date of this application; or
 - 6.2 C-reactive protein levels not measured as patient is currently receiving prednisone therapy at a dose of greater than 5 mg per day and has done so for more than three months.

Initiation - systemic juvenile idiopathic arthritis

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Re-assessment required after 6 months

Both:

- 1 Patient diagnosed with systemic juvenile idiopathic arthritis; and
- 2 Patient has tried and not responded to a reasonable trial of all of the following, either alone or in combination: oral or parenteral methotrexate; non-steroidal anti-inflammatory drugs (NSAIDs); and systemic corticosteroids.

Initiation - adult-onset Still's disease

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Re-assessment required after 6 months

Either: 1 Both:

1.1 Fither:

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- 1.1.1 The patient has had an initial Special Authority approval for adalimumab and/or etanercept for adult-onset Still's disease (AOSD); or
- 1.1.2 The patient has been started on tocilizumab for AOSD in a DHB hospital in accordance with the General Rules of the Pharmaceutical Schedule; and

1.2 Either:

- 1.2.1 The patient has experienced intolerable side effects from adalimumab and/or etanercept; or
- 1.2.2 The patient has received insufficient benefit from at least a three-month trial of adalimumab and/or etanercept such that they do not meet the renewal criteria for AOSD; or
- 2 All of the following:
 - 2.1 Patient diagnosed with AOSD according to the Yamaguchi criteria (J Rheumatol 1992;19:424-430); and
 - 2.2 Patient has tried and not responded to at least 6 months of glucocorticosteroids at a dose of at least 0.5 mg/kg, non-steroidal antiinflammatory drugs (NSAIDs) and methotrexate; and
 - 2.3 Patient has persistent symptoms of disabling poorly controlled and active disease.

Initiation - polyarticular juvenile idiopathic arthritis

Rheumatologist or Practitioner on the recommendation of a rheumatologist *Re-assessment required after 4 months* Either:

- 1 Both:
 - 1.1 The patient has had an initial Special Authority approval for both etanercept and adalimumab for juvenile idiopathic arthritis (JIA); and
 - 1.2 The patient has experienced intolerable side effects, or has received insufficient benefit from, both etanercept and adalimumab; or
- 2 All of the following:
 - 2.1 Treatment with a tumour necrosis factor alpha inhibitor is contraindicated; and
 - 2.2 Patient has had severe active polyarticular course JIA for 6 months duration or longer; and
 - 2.3 Patient has tried and not responded to at least three months of oral or parenteral methotrexate (at a dose of 10-20 mg/m² weekly or at the maximum tolerated dose) in combination with either oral corticosteroids (prednisone 0.25 mg/kg or at the maximum tolerated dose) or a full trial of serial intra-articular corticosteroid injections; and
 - 2.4 To be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
 - 2.5 Both:
 - 2.5.1 Either:
 - 2.5.1.1 Patient has persistent symptoms of poorly-controlled and active disease in at least 20 swollen, tender joints; or
 - 2.5.1.2 Patient has persistent symptoms of poorly-controlled and active disease in at least four joints from the following: wrist, elbow, knee, ankle, shoulder, cervical spine, hip; and
 - 2.5.2 Physician's global assessment indicating severe disease.

Initiation - idiopathic multicentric Castleman's disease

Haematologist, rheumatologist or Practitioner on the recommendation of a haematologist or rheumatologist

Re-assessment required after 6 months

All of the following:

- 1 Patient has severe HHV-8 negative idiopathic multicentric Castleman's disease; and
- 2 Treatment with an adequate trial of corticosteroids has proven ineffective; and
- 3 Tocilizumab to be administered at doses no greater than 8 mg/kg IV every 3-4 weeks.

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Continuation – Rheumatoid Arthritis

Rheumatologist or Practitioner on the recommendation of a rheumatologist *Re-assessment required after 6 months*

Either:

- 1 Following 6 months' initial treatment, the patient has at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician; or
- 2 On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician.

Continuation - systemic juvenile idiopathic arthritis

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Re-assessment required after 6 months

Either:

- 1 Following up to 6 months' initial treatment, the patient has achieved at least an American College of Rheumatology paediatric 30% improvement criteria (ACR Pedi 30) response from baseline; or
- 2 On subsequent reapplications, the patient demonstrates at least a continuing ACR Pedi 30 response from baseline.

Continuation - adult-onset Still's disease

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Re-assessment required after 6 months

the patient has a sustained improvement in inflammatory markers and functional status.

Continuation - polyarticular juvenile idiopathic arthritis

Rheumatologist or Practitioner on the recommendation of a rheumatologist

Re-assessment required after 6 months

Both:

- 1 Treatment is to be used as an adjunct to methotrexate therapy or monotherapy where use of methotrexate is limited by toxicity or intolerance; and
- 2 Either:
 - 2.1 Following 3 to 4 months' initial treatment, the patient has at least a 50% decrease in active joint count and an improvement in physician's global assessment from baseline; or
 - 2.2 On subsequent reapplications, the patient demonstrates at least a continuing 30% improvement in active joint count and continued improvement in physician's global assessment from baseline.

Continuation - idiopathic multicentric Castleman's disease

Haematologist, rheumatologist or Practitioner on the recommendation of a haematologist or rheumatologist Re-assessment required after 12 months

the treatment remains appropriate and the patient has a sustained improvement in inflammatory markers and functional status.

| TRASTUZUMAB – Restricted see terms below | | |
|--|---|-----------|
| Inj 150 mg vial | 1 | Herceptin |
| Inj 440 mg vial | 1 | Herceptin |

→ Restricted (RS1554)

Initiation - Early breast cancer

Limited to 12 months treatment

All of the following:

- 1 The patient has early breast cancer expressing HER 2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 Maximum cumulative dose of 106 mg/kg (12 months' treatment); and
- 3 Any of the following:
 - 3.1 9 weeks' concurrent treatment with adjuvant chemotherapy is planned; or
 - 3.2 12 months' concurrent treatment with adjuvant chemotherapy is planned; or
 - 3.3 12 months' sequential treatment following adjuvant chemotherapy is planned; or

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- 3.4 12 months' treatment with neoadjuvant and adjuvant chemotherapy is planned; or
- 3.5 Other treatment regimen, in association with adjuvant chemotherapy, is planned.

Initiation - metastatic breast cancer (trastuzumab-naive patients)

Limited to 12 months treatment

All of the following:

- 1 The patient has metastatic breast cancer expressing HER-2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 Either:
 - 2.1 The patient has not previously received lapatinib treatment for HER-2 positive metastatic breast cancer; or
 - 2.2 Both:
 - 2.2.1 The patient started lapatinib treatment for metastatic breast cancer but discontinued lapatinib within 3 months of starting treatment due to intolerance; and
 - 2.2.2 The cancer did not progress whilst on lapatinib; and
- 3 Either:
 - 3.1 Trastuzumab will not be given in combination with pertuzumab; or
 - 3.2 All of the following:
 - 3.2.1 Trastuzumab to be administered in combination with pertuzumab; and
 - 3.2.2 Patient has not received prior treatment for their metastatic disease and has had a treatment-free interval of at least 12 months between prior (neo)adjuvant chemotherapy treatment and diagnosis of metastatic breast cancer; and
 - 3.2.3 The patient has good performance status (ECOG grade 0-1); and
- 4 Trastuzumab not to be given in combination with lapatinib; and
- 5 Trastuzumab to be discontinued at disease progression.

Initiation - metastatic breast cancer (patients previously treated with trastuzumab)

Limited to 12 months treatment

All of the following:

- 1 The patient has metastatic breast cancer expressing HER-2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 Either:
 - 2.1 The patient has not previously received lapatinib treatment for HER-2 positive metastatic breast cancer; or
 - 2.2 Both:
 - 2.2.1 The patient started lapatinib treatment for metastatic breast cancer but discontinued lapatinib within 3 months of starting treatment due to intolerance; and
 - 2.2.2 The cancer did not progress whilst on lapatinib; and
- 3 Either:
 - 3.1 Trastuzumab will not be given in combination with pertuzumab; or
 - 3.2 All of the following:
 - 3.2.1 Trastuzumab to be administered in combination with pertuzumab; and
 - 3.2.2 Patient has not received prior treatment for their metastatic disease and has had a treatment-free interval of at least 12 months between prior (neo)adjuvant chemotherapy treatment and diagnosis of metastatic breast cancer; and
 - 3.2.3 The patient has good performance status (ECOG grade 0-1); and
- 4 Trastuzumab not to be given in combination with lapatinib; and
- 5 Trastuzumab to be discontinued at disease progression.

Continuation - metastatic breast cancer

Re-assessment required after 12 months All of the following:

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continued...

- 1 The patient has metastatic breast cancer expressing HER-2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 The cancer has not progressed at any time point during the previous 12 months whilst on trastuzumab; and
- 3 Trastuzumab not to be given in combination with lapatinib; and
- 4 Trastuzumab to be discontinued at disease progression.

TRASTUZUMAB EMTANSINE - Restricted see terms below

| t | Inj 100 mg vial2 | ,320.00 | 1 | Kadcyla |
|---|------------------|---------|---|---------|
| | Inj 160 mg vial | ,712.00 | 1 | Kadcyla |
| | | | | |

⇒ Restricted (RS1715)

Initiation

Re-assessment required after 6 months

All of the following:

- 1 Patient has metastatic breast cancer expressing HER-2 IHC 3+ or ISH+ (including FISH or other current technology); and
- 2 Patient has previously received trastuzumab and chemotherapy, separately or in combination; and
- 3 Either:
 - 3.1 The patient has received prior therapy for metastatic disease*; or
 - 3.2 The patient developed disease recurrence during, or within six months of completing adjuvant therapy*; and
- 4 Patient has a good performance status (ECOG 0-1); and
- 5 Either:
 - 5.1 Patient does not have symptomatic brain metastases; or
 - 5.2 Patient has brain metastases and has received prior local CNS therapy; and
- 6 Treatment to be discontinued at disease progression.

Continuation

Re-assessment required after 6 months

Both:

- 1 The cancer has not progressed at any time point during the previous approval period whilst on trastuzumab emtansine; and
- 2 Treatment to be discontinued at disease progression.

Note: *Note: Prior or adjuvant therapy includes anthracycline, other chemotherapy, biological drugs, or endocrine therapy.

Programmed Cell Death-1 (PD-1) Inhibitors

| NIVOLUMAB – Restricted see terms below | | | |
|--|----------|---|--------|
| Inj 10 mg per ml, 4 ml vial | 1,051.98 | 1 | Opdivo |
| Inj 10 mg per ml, 10 ml vial | 2,629.96 | 1 | Opdivo |
| ➡ Restricted (RS1742) | | | |

Initiation

Medical oncologist *Re-assessment required after 4 months* All of the following:

- 1 Patient has metastatic or unresectable melanoma (excluding uveal) stage III or IV; and
- 2 Patient has measurable disease as defined by RECIST version 1.1; and
- 3 The patient has ECOG performance score of 0-2; and
- 4 Either:
 - 4.1 Patient has not received funded pembrolizumab; or

4.2 Both:

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- 4.2.1 Patient has received an initial Special Authority approval for pembrolizumab and has discontinued pembrolizumab within 12 weeks of starting treatment due to intolerance; and
- 4.2.2 The cancer did not progress while the patient was on pembrolizumab; and
- 5 Baseline measurement of overall tumour burden is documented (see Note); and
- 6 Documentation confirming that the patient has been informed and acknowledges that funded treatment with nivolumab will not be continued if their disease progresses.

Continuation

Medical oncologist

Re-assessment required after 4 months Either:

- 1 All of the following:
 - 1.1 Any of the following:
 - 1.1.1 Patient's disease has had a complete response to treatment according to RECIST criteria (see Note); or
 - 1.1.2 Patient's disease has had a partial response to treatment according to RECIST criteria (see Note); or
 - 1.1.3 Patient has stable disease according to RECIST criteria (see Note); and
 - 1.2 Patient's disease has not progressed clinically and disease response to treatment has been clearly documented in patient notes; and
 - 1.3 No evidence of progressive disease according to RECIST criteria (see Note); and
 - 1.4 The treatment remains clinically appropriate and the patient is benefitting from the treatment; or

2 All of the following:

- 2.1 Patient has previously discontinued treatment with nivolumab for reasons other than severe toxicity or disease progression; and
- 2.2 Patient has signs of disease progression; and
- 2.3 Disease has not progressed during previous treatment with nivolumab.

Notes: Baseline assessment and disease responses to be assessed according to the Response Evaluation Criteria in Solid Tumours (RECIST) version 1.1 (Eisenhauer EA, et al. Eur J Cancer 2009;45:228-47). Assessments of overall tumour burden and measurable disease to be undertaken on a minimum of one lesion and maximum of 5 target lesions (maximum two lesions per organ). Target lesions should be selected on the basis of their size (lesions with the longest diameter), be representative of all involved organs, and suitable for reproducible repeated measurements. Measurable disease includes by CT or MRI imaging or caliper measurement by clinical exam. Target lesion measurements should be assessed using the same method of assessment and the same technique used to characterise each identified and reported lesion at baseline and every 12 weeks. Response definitions as follows:

- Complete Response: Disappearance of all target lesions. Any pathological lymph nodes (whether target or non-target) must have reduction in short axis to < 10 mm.
- Partial Response: At least a 30% decrease in the sum of diameters of target lesions, taking as reference the baseline sum diameters.
- Progressive Disease: At least a 20% increase in the sum of diameters of target lesions, taking as reference the smallest sum on study (this includes the baseline sum if that is the smallest on study). In addition to the relative increase of 20%, the sum must also demonstrate an absolute increase of at least 5 mm. (Note: the appearance of one or more new lesions is also considered progression).
- Stable Disease: Neither sufficient shrinkage to qualify for partial response nor sufficient increase to qualify for progressive disease.

PEMBROLIZUMAB - Restricted see terms below

202

| Inj 25 mg per ml, 4 ml vial | 1 | Keytruda |
|---------------------------------------|-------|----------|
| ➡ Restricted (RS1741) | | - |
| Initiation | | |
| Medical oncologist | | |
| Re-assessment required after 4 months | | |
| All of the following: | | |

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- 1 Patient has metastatic or unresectable melanoma (excluding uveal) stage III or IV; and
- 2 Patient has measurable disease as defined by RECIST version 1.1; and
- 3 The patient has ECOG performance score of 0-2; and
- 4 Either:
 - 4.1 Patient has not received funded nivolumab; or
 - 4.2 Both:
 - 4.2.1 Patient has received an initial Special Authority approval for nivolumab and has discontinued nivolumab within 12 weeks of starting treatment due to intolerance; and
 - 4.2.2 The cancer did not progress while the patient was on nivolumab; and
- 5 Baseline measurement of overall tumour burden is documented (see Note); and
- 6 Documentation confirming that the patient has been informed and acknowledges that funded treatment with pembrolizumab will not be continued if their disease progresses.

Continuation

Medical oncologist

Re-assessment required after 4 months Either:

1 All of the following:

- 1.1 Any of the following:
 - 1.1.1 Patient's disease has had a complete response to treatment according to RECIST criteria (see Note); or
 - 1.1.2 Patient's disease has had a partial response to treatment according to RECIST criteria (see Note); or
 - 1.1.3 Patient has stable disease according to RECIST criteria (see Note); and
- 1.2 Patient's disease has not progressed clinically and disease response to treatment has been clearly documented in patient notes; and
- 1.3 No evidence of progressive disease according to RECIST criteria (see Note); and
- 1.4 The treatment remains clinically appropriate and the patient is benefitting from the treatment; or
- 2 All of the following:
 - 2.1 Patient has previously discontinued treatment with pembrolizumab for reasons other than severe toxicity or disease progression; and
 - 2.2 Patient has signs of disease progression; and
 - 2.3 Disease has not progressed during previous treatment with pembrolizumab.

Notes: Baseline assessment and disease responses to be assessed according to the Response Evaluation Criteria in Solid Tumours (RECIST) version 1.1 (Eisenhauer EA, et al. Eur J Cancer 2009;45:228-47). Assessments of overall tumour burden and measurable disease to be undertaken on a minimum of one lesion and maximum of 5 target lesions (maximum two lesions per organ). Target lesions should be selected on the basis of their size (lesions with the longest diameter), be representative of all involved organs, and suitable for reproducible repeated measurements. Measurable disease includes by CT or MRI imaging or caliper measurement by clinical exam. Target lesion measurements should be assessed using the same method of assessment and the same technique used to characterise each identified and reported lesion at baseline and every 12 weeks. Response definitions as follows:

- Complete Response: Disappearance of all target lesions. Any pathological lymph nodes (whether target or non-target) must have reduction in short axis to < 10 mm.
- Partial Response: At least a 30% decrease in the sum of diameters of target lesions, taking as reference the baseline sum diameters.
- Progressive Disease: At least a 20% increase in the sum of diameters of target lesions, taking as reference the smallest sum on study (this includes the baseline sum if that is the smallest on study). In addition to the relative increase of 20%, the sum must also demonstrate an absolute increase of at least 5 mm. (Note: the appearance of one or more new lesions is also considered progression).
- Stable Disease: Neither sufficient shrinkage to qualify for partial response nor sufficient increase to qualify for progressive

| (e | Price x man. excl. GST \$ | Per | Brand or Generic Manufacturer |
|---|---------------------------------|--------------|-------------------------------------|
| continued disease. | | | |
| Other Immunosuppressants | | | |
| ANTITHYMOCYTE GLOBULIN (EQUINE) | 0.054.05 | - | 470444 |
| Inj 50 mg per ml, 5 ml ampoule | 2,351.25 | 5 | ATGAM |
| ANTITHYMOCYTE GLOBULIN (RABBIT) Inj 25 mg vial | | | |
| AZATHIOPRINE | | | |
| Tab 25 mg – 1% DV Jan-20 to 2022 | 7.35 | 60 | Azamun |
| Tab 50 mg - 1% DV Jan-20 to 2022 | 7.60 | 100 | Azamun |
| Inj 50 mg vial – 1% DV Nov-19 to 2022 | | 1 | Imuran |
| BACILLUS CALMETTE-GUERIN (BCG) – Restricted see terms below | | | |
| Inj 2-8 × 10 [°] 8 CFU vial | | 1 | OncoTICE |
| → Restricted (RS1206) | | | |
| Initiation | | | |
| For use in bladder cancer. | | | |
| EVEROLIMUS – Restricted see terms below | | | |
| | 4,555.76 | 30 | Afinitor |
| Tab 10 mg | 6,512.29 | 30 | Afinitor |
| → Restricted (RS1745) | | | |
| Initiation | | | |
| Neurologist or oncologist | | | |
| Re-assessment required after 3 months | | | |
| Both: | | | |
| 1 Patient has tuberous sclerosis; and | | | |
| 2 Patient has progressively enlarging sub-ependymal giant cell astro | cytomas (SEGAs | s) that requ | uire treatment. |
| Continuation – pandemic circumstances | | | |
| Re-assessment required after 6 months | | | |
| All of the following: | | | |
| 1 The patient is clinically benefiting from treatment and continued tre | atment remains a | appropriate | e; and |
| 2 Everolimus to be discontinued at progression of SEGAs; and | | | |
| 3 The regular renewal requirements cannot be met due to COVID-19 | | | |
| Note: MRI should be performed at minimum once every 12 months, more | | • | • |
| of symptoms such as headaches, visual complaints, nausea or vomiting, o | or increase in seiz | zure activit | ty. |
| Continuation | | | |
| Neurologist or oncologist | | | |
| Re-assessment required after 12 months | | | |
| All of the following: | ithin the last 0 | anthai are | L |
| 1 Documented evidence of SEGA reduction or stabilisation by MRI w | | | 1 |
| The treatment remains appropriate and the patient is benefiting fro Everolimus to be discontinued at progression of SEGAs. | in irealment, and | | |
| | froquent coordin | | he performed with new energy |
| Note: MRI should be performed at minimum once every 12 months, more of symptoms such as headaches, visual complaints, nausea or vomiting, or | | • | • |
| MYCOPHENOLATE MOFETIL | | | |
| Tab 500 mg | | 50 | CellCept |

| Tab 500 mg | 0 | CellCept |
|-----------------|------|----------|
| Cap 250 mg | 00 | CellCept |
| | i ml | CellCept |
| Inj 500 mg vial | 1 | CellCept |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|-------|-------------------------------------|
| PICIBANIL | | | |
| Inj 100 mg vial | | | |
| SIROLIMUS – Restricted see terms below | | | |
| ↓ Tab 1 mg | | 100 | Rapamune |
| ↓ Tab 2 mg | | 100 | Rapamune |
| Oral liq 1 mg per ml | | 60 ml | Rapamune |
| ➡ Restricted (RS1208) | | | |
| Initiation | | | |

For rescue therapy for an organ transplant recipient.

Notes: Rescue therapy defined as unresponsive to calcineurin inhibitor treatment as defined by refractory rejection; or intolerant to calcineurin inhibitor treatment due to any of the following:

- GFR < 30 ml/min; or
- · Rapidly progressive transplant vasculopathy; or
- Rapidly progressive obstructive bronchiolitis; or
- HUS or TTP; or
- · Leukoencepthalopathy; or
- Significant malignant disease

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|--|-----------------|-------------------------------------|
| Antiallergy Preparations | | | |
| Allergic Emergencies | | | |
| CATIBANT - Restricted see terms below Inj 10 mg per ml, 3 ml prefilled syringe | | 1 odominal a | Firazyr |
| angloedema (HAE) for patients with confirmed diagnosis of (2 The patient has undergone product training and has agreed Continuation Re-assessment required after 12 months The treatment remains appropriate and the patient is benefiting from | C1-esterase inhibitor def upon an action plan for s | iciency; ar | nd |
| Allergy Desensitisation | | | |
| BEE VENOM – Restricted see terms below Maintenance kit - 6 vials 120 mcg freeze dried venom, with dilu Inj 550 mcg vial with diluent → Restricted (RS1117) nitiation Both: 1 RAST or skin test positive; and 2 Patient has had severe generalised reaction to the sensitisin | | | |
| APER WASP VENOM - Restricted see terms below Treatment kit - 6 vials 120 mcg freeze dried venom, with diluent Inj 550 mcg vial with diluent → Restricted (RS1118) nitiation Both: 1 RAST or skin test positive; and 2 Patient has had severe generalised reaction to the sensitisin | | | |
| YELLOW JACKET WASP VENOM - Restricted see terms below Treatment kit - 6 vials 120 mcg freeze dried venom, with diluent → Restricted (RS1119) nitiation Both: 1 RAST or skin test positive; and 2 Patient has had severe generalised reaction to the sensitisin | | | |
| Allergy Prophylactics | | | |

| Nasal spray 50 mcg per dose - 1% DV Oct-20 to 2023 | 200 dose | SteroClear |
|---|----------|------------|
| Nasal spray 100 mcg per dose - 1% DV Oct-20 to 2023 | 200 dose | SteroClear |

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e.g. Brand indicates brand example only. It is not a contracted product.

| | Price | - | Brand or Generic | |
|---|-------------------------|---------------|---------------------------------|--|
| | (ex man. excl. GS \$ | I) Per | Generic Manufacturer | |
| LUTICASONE PROPIONATE | | | | |
| Nasal spray 50 mcg per dose - 1% DV Nov-18 to 2021 | 1.98 | 120 dose | Flixonase Hayfever & Allergy | |
| PRATROPIUM BROMIDE Aqueous nasal spray 0.03% | 4.61 | 15 ml | Univent | |
| ODIUM CROMOGLICATE Nasal spray 4% | | | | |
| Antihistamines | | | | |
| ETIRIZINE HYDROCHLORIDE | | | | |
| Tab 10 mg – 1% DV Nov-19 to 2022 Oral liq 1 mg per ml | | 100 200 ml | Zista Histaclear | |
| CHLORPHENIRAMINE MALEATE Oral liq 0.4 mg per ml | | | | |
| Inj 10 mg per ml, 1 ml ampoule YPROHEPTADINE HYDROCHLORIDE | | | | |
| Tab 4 mg EXOFENADINE HYDROCHLORIDE | | | | |
| Tab 60 mg Tab 120 mg | | | | |
| Tab 180 mg | | | | |
| ORATADINE Tab 10 mg - 1% DV Feb-20 to 2022 | | 100 | Lorafix | |
| Oral liq 1 mg per ml | | 120 ml | Lorfast | |
| ROMETHAZINE HYDROCHLORIDE Tab 10 mg - 1% DV Sep-18 to 2021 | | 50 | Allersoothe | |
| Tab 25 mg - 1% DV Sep-18 to 2021 | | 50 | Allersoothe | |
| Oral liq 1 mg per ml - 1% DV Sep-18 to 2021 | | 100 ml | Allersoothe | |
| Inj 25 mg per ml, 2 ml ampoule | | 5 | Hospira | |
| Anticholinergic Agents | | | | |
| PRATROPIUM BROMIDE | | | | |
| Aerosol inhaler 20 mcg per dose Nebuliser soln 250 mcg per ml, 1 ml ampoule | 3,35 | 20 | Univent | |
| Nebuliser soln 250 mcg per ml, 2 ml ampoule – 1% DV Jan-20 | | 20 | Univent | |
| Anticholinergic Agents with Beta-Adrenoceptor A | gonists | | | |
| ALBUTAMOL WITH IPRATROPIUM BROMIDE | | | | |
| Aerosol inhaler 100 mcg with ipratropium bromide 20 mcg per de Nebuliser soln 2.5 mg with ipratropium bromide 0.5 mg per 2.5 r | nl | | Dualia | |
| ampoule – 1% DV Oct-18 to 2021 | | 20 | Duolin | |
| | | | | |
| LYCOPYRRONIUM Note: inhaled glycopyrronium treatment must not be used if the or umeclidinium. | patient is also receiv | ing treatmen | t with subsidised tiotropiu | |
| Powder for inhalation 50 mcg per dose | 61.00 | 30 dose | Seebri Breezhaler | |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | Price (ex man. excl. GS \$ | ST) Per | Brand or Generic Manufacturer |
|---|----------------------------------|--------------------------|--|
| TIOTROPIUM BROMIDE | | | |
| Note: tiotropium treatment must not be used if the patient is also or umeclidinium. | receiving treatment | t with subsidi | sed inhaled glycopyrronium |
| Soln for inhalation 2.5 mcg per dose | 50.37 | 60 dose | Spiriva Respimat |
| Powder for inhalation 18 mcg per dose | 50.37 | 30 dose | Spiriva |
| UMECLIDINIUM Note: Umeclidinium must not be used if the patient is also receiv tiotropium bromide. Powder for inhalation 62.5 mcg per dose | Ū | ubsidised inh 30 dose | naled glycopyrronium or Incruse Ellipta |

Long-Acting Muscarinic Antagonists with Long-Acting Beta-Adrenoceptor Agonists

→ Restricted (RS1518)

Initiation

Re-assessment required after 2 years

Both:

- 1 Patient has been stabilised on a long acting muscarinic antagonist; and
- 2 The prescriber considers that the patient would receive additional benefit from switching to a combination product.

Continuation

Re-assessment required after 2 years

Both:

- 1 Patient is compliant with the medication; and
- 2 Patient has experienced improved COPD symptom control (prescriber determined).

Note: Combination long acting muscarinic antagonist and long acting beta-2 agonist must not be used if the patient is also receiving treatment with a combination inhaled corticosteroid and long acting beta-2 agonist.

| GLYCOPYRRONIUM WITH INDACATEROL – Restricted see terms above | | |
|---|---------|--------------------|
| Powder for Inhalation 50 mcg with indacaterol 110 mcg | 30 dose | Ultibro Breezhaler |
| TIOTROPIUM BROMIDE WITH OLODATEROL - Restricted see terms above | | |
| t Soln for inhalation 2.5 mcg with olodaterol 2.5 mcg | 60 dose | Spiolto Respimat |
| UMECLIDINIUM WITH VILANTEROL – Restricted see terms above | | |
| t Powder for inhalation 62.5 mcg with vilanterol 25 mcg77.00 | 30 dose | Anoro Ellipta |

Antifibrotics

| NINTEDANIB – Restricted see terms below | | | |
|---|----------|----|------|
| Cap 100 mg | 2,554.00 | 60 | Ofev |
| ↓ Cap 150 mg | | 60 | Ofev |

➡ Restricted (RS1756)

Initiation – idiopathic pulmonary fibrosis

Respiratory specialist

Re-assessment required after 12 months

All of the following:

- 1 Patient has been diagnosed with idiopathic pulmonary fibrosis; and
- 2 Forced vital capacity is between 50% and 90% predicted; and
- 3 Nintedanib is to be discontinued at disease progression (See Note); and
- 4 Nintedanib is not to be used in combination with subsidised pirfenidone; and
- 5 Any of the following:

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

continued...

- 5.1 The patient has not previously received treatment with pirfenidone; or
- 5.2 Patient has previously received pirfenidone, but discontinued pirfenidone within 12 weeks due to intolerance; or
- 5.3 Patient has previously received pirfenidone, but the patient's disease has not progressed (disease progression defined as 10% or more decline in predicted FVC within any 12 month period since starting treatment with pirfenidone).

Continuation - idiopathic pulmonary fibrosis

Re-assessment required after 12 months

All of the following:

- 1 Treatment remains clinically appropriate and patient is benefitting from and tolerating treatment; and
- 2 Nintedanib is not to be used in combination with subsidised pirfenidone; and
- 3 Nintedanib is to be discontinued at disease progression (See Note).

Note: disease progression is defined as a decline in percent predicted FVC of 10% or more within any 12 month period.

PIRFENIDONE - Restricted see terms below

| t | Tab 801 mg | 90 | Esbriet |
|---|--------------------|-----|---------|
| t | Cap 267 mg3,645.00 | 270 | Esbriet |

⇒ Restricted (RS1757)

Initiation - idiopathic pulmonary fibrosis

Respiratory specialist

Re-assessment required after 12 months

All of the following:

- 1 Patient has been diagnosed with idiopathic pulmonary fibrosis; and
- 2 Forced vital capacity is between 50% and 90% predicted; and
- 3 Pirfenidone is to be discontinued at disease progression (See Notes); and
- 4 Pirfenidone is not to be used in combination with subsidised nintedanib; and
- 5 Any of the following:
 - 5.1 The patient has not previously received treatment with nintedanib; or
 - 5.2 Patient has previously received nintedanib, but discontinued nintedanib within 12 weeks due to intolerance; or
 - 5.3 Patient has previously received nintedanib, but the patient's disease has not progressed (disease progression defined as 10% or more decline in predicted FVC within any 12 month period since starting treatment with nintedanib).

Continuation - idiopathic pulmonary fibrosis

Re-assessment required after 12 months

All of the following:

- 1 Treatment remains clinically appropriate and patient is benefitting from and tolerating treatment; and
- 2 Pirfenidone is not to be used in combination with subsidised nintedanib; and
- 3 Pirfenidone is to be discontinued at disease progression (See Note).

Note: disease progression is defined as a decline in percent predicted FVC of 10% or more within any 12 month period.

Beta-Adrenoceptor Agonists

SALBUTAMOL

| Oral liq 400 mcg per ml – 1% DV Nov-18 to 2021 | 150 ml | Ventolin |
|---|----------|----------|
| Inj 500 mcg per ml, 1 ml ampoule | | |
| Inj 1 mg per ml, 5 ml ampoule | | |
| Aerosol inhaler, 100 mcg per dose | 200 dose | SalAir |
| 6.00 | | Ventolin |
| Nebuliser soln 1 mg per ml, 2.5 ml ampoule - 1% DV Oct-18 to 2021 | 20 | Asthalin |
| Nebuliser soln 2 mg per ml, 2.5 ml ampoule - 1% DV Oct-18 to 20214.03 | 20 | Asthalin |
| | | |

| | Price (ex man. excl. G \$ | ST) Per | Brand or Generic Manufacturer |
|--|---------------------------------|------------|-------------------------------------|
| TERBUTALINE SULPHATE Powder for inhalation 250 mcg per dose Inj 0.5 mg per ml, 1 ml ampoule | | | |
| Powder for inhalation, 200 mcg per dose (equivalent to 250 mcg metered dose), breath activated | | 120 dose | Bricanyl Turbuhaler |
| Cough Suppressants | | | |
| PHOLCODINE Oral liq 1 mg per ml – 1% DV Jun-20 to 2022 | | 200 ml | AFT Pholcodine Linctus BP |
| Decongestants | | | |
| OXYMETAZOLINE HYDROCHLORIDE Aqueous nasal spray 0.25 mg per ml Aqueous nasal spray 0.5 mg per ml | | | |
| PSEUDOEPHEDRINE HYDROCHLORIDE Tab 60 mg | | | |
| SODIUM CHLORIDE Aqueous nasal spray isotonic | | | |
| SODIUM CHLORIDE WITH SODIUM BICARBONATE Soln for nasal irrigation | | | |
| XYLOMETAZOLINE HYDROCHLORIDE Aqueous nasal spray 0.05% Aqueous nasal spray 0.1% Nasal drops 0.05% Nasal drops 0.1% | | | |
| Inhaled Corticosteroids | | | |
| BECLOMETHASONE DIPROPIONATE | | | |

| Aerosol inhaler 50 mcg per dose | 8.54 | 200 dose | Beclazone 50 | |
|----------------------------------|-------|----------|---------------|--|
| | 9.30 | | Qvar | |
| Aerosol inhaler 100 mcg per dose | 12.50 | 200 dose | Beclazone 100 | |
| | 15.50 | | Qvar | |
| Aerosol inhaler 250 mcg per dose | 22.67 | 200 dose | Beclazone 250 | |
| | | | | |

BUDESONIDE

Nebuliser soln 250 mcg per ml, 2 ml ampoule Nebuliser soln 500 mcg per ml, 2 ml ampoule Powder for inhalation 100 mcg per dose Powder for inhalation 200 mcg per dose Powder for inhalation 400 mcg per dose

| | Price | | Brand or |
|---|-------------------|------------|-------------------------|
| | (ex man. excl. GS | ST) Per | Generic Manufacturer |
| | ð | Per | Manufacturer |
| LUTICASONE | | | |
| Aerosol inhaler 50 mcg per dose – 1% DV Sep-20 to 2023 | | 120 dose | Flixotide |
| | 4.68 | | Floair |
| Powder for inhalation 50 mcg per dose | | 60 dose | Flixotide Accuhaler |
| Powder for inhalation 100 mcg per dose | | 60 dose | Flixotide Accuhaler |
| Aerosol inhaler 125 mcg per dose - 1% DV Sep-20 to 2023 | | 120 dose | Flixotide |
| | 7.22 | | Floair |
| Aerosol inhaler 250 mcg per dose – 1% DV Sep-20 to 2023 | | 120 dose | Flixotide |
| | 10.18 | | Floair |
| Powder for inhalation 250 mcg per dose | 24.51 | 60 dose | Flixotide Accuhaler |
| Floair Aerosol inhaler 50 mcg per dose to be delisted 1 September 2 | 2020) | | |
| Floair Aerosol inhaler 125 mcg per dose to be delisted 1 September | 2020) | | |
| Floair Aerosol inhaler 250 mcg per dose to be delisted 1 September | 2020) | | |
| Leukotriene Receptor Antagonists | | | |
| IONTELUKAST | | | |
| Tab 4 mg – 1% DV Jan-20 to 2022 | | 28 | Montelukast Mylar |
| Tab 5 mg – 1% DV Jan-20 to 2022 | | 28 | Montelukast Mylar |
| Tab 10 mg - 1% DV Jan-20 to 2022 | | 28 | Montelukast Mylar |
| ······································ | | | ,, |
| | | | |
| Long-Acting Beta-Adrenoceptor Agonists | | | |
| Long-Acting Beta-Adrenoceptor Agonists | | | |

EFORMOTEROL FUMARATE DIHYDRATE

Powder for inhalation 4.5 mcg per dose, breath activated (equivalent to

eformoterol fumarate 6 mcg metered dose)

INDACATEROL

| Powder for inhalation 150 mcg per dose Powder for inhalation 300 mcg per dose | | | Onbrez Breezhaler Onbrez Breezhaler |
|--|-------|----------|--|
| SALMETEROL | | | |
| Aerosol inhaler 25 mcg per dose | 9.90 | 120 dose | Meterol |
| | 25.00 | | Serevent |
| Powder for inhalation 50 mcg per dose | 25.00 | 60 dose | Serevent Accuhaler |

Inhaled Corticosteroids with Long-Acting Beta-Adrenoceptor Agonists

BUDESONIDE WITH EFORMOTEROL

Powder for inhalation 100 mcg with eformoterol fumarate 6 mcg Powder for inhalation 200 mcg with eformoterol fumarate 6 mcg Powder for inhalation 400 mcg with eformoterol fumarate 12 mcg Aerosol inhaler 100 mcg with eformoterol fumarate 6 mcg Aerosol inhaler 200 mcg with eformoterol fumarate 6 mcg FLUTICASONE FUROATE WITH VILANTEROL Powder for inhalation 100 mcg with vilanterol 25 mcg44.08 30 dose Breo Ellipta

| | Price | | Brand or |
|--|-------------------|---------------------|--------------------|
| | (ex man. excl. GS | (ex man. excl. GST) | |
| | \$ | Per | Manufacturer |
| FLUTICASONE WITH SALMETEROL | | | |
| Aerosol inhaler 50 mcg with salmeterol 25 mcg - 1% DV Sep-2 | 0 to 202314.58 | 120 dose | RexAir |
| | 25.79 | | Seretide |
| Powder for inhalation 100 mcg with salmeterol 50 mcg | | 60 dose | Seretide Accuhaler |
| Aerosol inhaler 125 mcg with salmeterol 25 mcg - 1% DV Sep- | 20 | | |
| to 2023 | | 120 dose | RexAir |
| | 32.60 | | Seretide |
| Powder for inhalation 250 mcg with salmeterol 50 mcg | | 60 dose | Seretide Accuhaler |
| (RexAir Aerosol inhaler 50 mcg with salmeterol 25 mcg to be deliste (RexAir Aerosol inhaler 125 mcg with salmeterol 25 mcg to be delist | | , | |

Mast Cell Stabilisers

NEDOCROMIL

Aerosol inhaler 2 mg per dose

(Any Aerosol inhaler 2 mg per dose to be delisted 1 February 2021)

SODIUM CROMOGLICATE

Aerosol inhaler 5 mg per dose

(Any Aerosol inhaler 5 mg per dose to be delisted 1 May 2021)

Methylxanthines

AMINOPHYLLINE

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| AMINOPHYLLINE Inj 25 mg per ml, 10 ml ampoule124.37 | 5 | DBL Aminophylline |
|---|--------|-------------------|
| CAFFEINE CITRATE Oral liq 20 mg per ml (caffeine 10 mg per ml) – 1% DV Nov-19 to 202215.10 | 25 ml | Biomed |
| Inj 20 mg per ml (caffeine 10 mg per ml), 2.5 ml ampoule – 1% DV Nov-19 to 2022 | 5 | Biomed |
| THEOPHYLLINE Tab long-acting 250 mg – 1% DV Jan-20 to 2022 | 100 | Nuelin-SR |
| Oral liq 80 mg per 15 ml – 1% DV Jan-20 to 2022 | 500 ml | Nuelin |

Mucolytics and Expectorants

| DORNASE ALFA – Restricted see terms below | | | |
|---|------------|-------|-----------|
| I Nebuliser soln 2.5 mg per 2.5 ml ampoule | 250.00 | 6 | Pulmozyme |
| ➡ Restricted (RS1352) | | | |
| Initiation – cystic fibrosis | | | |
| The patient has cystic fibrosis and has been approved by the Cystic Fibro | sis Panel. | | |
| Initiation – significant mucus production | | | |
| Limited to 4 weeks treatment | | | |
| Both: | | | |
| 1 Patient is an in-patient; and | | | |
| 2 The mucus production cannot be cleared by first line chest techniq | ues. | | |
| Initiation – pleural emphyema | | | |
| Limited to 3 days treatment | | | |
| Both: | | | |
| 1 Patient is an in-patient; and | | | |
| 2 Patient diagnoses with pleural emphyema. | | | |
| SODIUM CHLORIDE | | | |
| Nebuliser soln 7%. 90 ml bottle – 1% DV Nov-19 to 2022 | 24 50 | 90 ml | Biomed |
| | | 00111 | Diomou |
| | | | |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

| (e | Price x man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|----------------------------------|-----|-------------------------------------|
| Pulmonary Surfactants | | | |
| BERACTANT Soln 200 mg per 8 ml vial | | | |
| PORACTANT ALFA Soln 120 mg per 1.5 ml vial | 425.00 | 1 | Curosurf |
| Soln 240 mg per 3 ml vial | | 1 | Curosurf |
| Respiratory Stimulants | | | |
| DOXAPRAM Inj 20 mg per ml, 5 ml vial | | | |
| | | | |

Sclerosing Agents

TALC

Powder Soln (slurry) 100 mg per ml, 50 ml

| | Price | | Brand or |
|---|--------------------------|---------------|-------------------------|
| | (ex man. excl. GST \$ |) Per | Generic Manufacturer |
| Anti-Infective Preparations | | | |
| Antibacterials | | | |
| CHLORAMPHENICOL Eye oint 1% – 1% DV May-20 to 2022 Ear drops 0.5% | 1.55 | 5 g | Devatis |
| Eye drops 0.5% – 1% DV Nov-19 to 2022 Eye drops 0.5%, single dose | 1.54 | 10 ml | Chlorafast |
| CIPROFLOXACIN Eye drops 0.3% | 9.99 | 5 ml | Ciprofloxacin Teva |
| FRAMYCETIN SULPHATE Ear/eye drops 0.5% | | | |
| GENTAMICIN SULPHATE Eye drops 0.3% PROPAMIDINE ISETHIONATE | 11.40 | 5 ml | Genoptic |
| Eye drops 0.1% SODIUM FUSIDATE [FUSIDIC ACID] | | | |
| Eye drops 1% SULPHACETAMIDE SODIUM Eye drops 10% | 5.29 | 5 g | Fucithalmic |
| TOBRAMYCIN Eye oint 0.3% Eye drops 0.3% | | 3.5 g 5 ml | Tobrex Tobrex |
| Antifungals | | | |
| NATAMYCIN Eye drops 5% | | | |
| Antivirals | | | |
| ACICLOVIR Eye oint 3% | 14.92 | 4.5 g | ViruPOS |
| Combination Preparations | | | |
| CIPROFLOXACIN WITH HYDROCORTISONE Ear drops ciprofloxacin 0.2% with 1% hydrocortisone | 16.30 | 10 ml | Ciproxin HC Otic |
| DEXAMETHASONE WITH FRAMYCETIN AND GRAMICIDIN Ear/eye drops 500 mcg with framycetin sulphate 5 mg and gramicio 50 mcg per ml | lin | | |
| DEXAMETHASONE WITH NEOMYCIN SULPHATE AND POLYMYXIN Eye oint 0.1% with neomycin sulphate 0.35% and polymyxin b sulp | hate | | |
| 6,000 u per g Eye drops 0.1% with neomycin sulphate 0.35% and polymyxin b sulphate 6,000 u per ml | | 3.5 g 5 ml | Maxitrol Maxitrol |
| DEXAMETHASONE WITH TOBRAMYCIN Eye drops 0.1% with tobramycin 0.3% | | 5 ml | Tobradex |

t Item restricted (see → above); t Item restricted (see → below)

e.g. Brand indicates brand example only. It is not a contracted product.

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SENSORY ORGANS

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|------------------------------------|-------------|-------------------------------------|
| FLUMETASONE PIVALATE WITH CLIOQUINOL Ear drops 0.02% with clioquinol 1% | | | |
| TRIAMCINOLONE ACETONIDE WITH GRAMICIDIN, NEOMYCIN AN | ID NYSTATIN | | |
| Ear drops 1 mg with nystatin 100,000 u, neomycin sulphate 2.5 m gramicidin 250 mcg per g | • | 7.5 ml | Kenacomb |
| Anti-Inflammatory Preparations | | | |
| Corticosteroids | | | |
| DEXAMETHASONE | | | |
| Eye oint 0.1% | | 3.5 g | Maxidex |
| Eye drops 0.1% | | 5 ml | Maxidex |
| Ccular implant 700 mcg | 1,444.50 | 1 | Ozurdex |
| → Restricted (RS1606) | | | |
| Initiation – Diabetic macular oedema | | | |
| Ophthalmologist Re-assessment required after 12 months | | | |
| All of the following: | | | |
| 1 Patients have diabetic macular oedema with pseudophakic len | s: and | | |
| 2 Patient has reduced visual acuity of between 6/9 - 6/48 with fu | | f reduction | in vision; and |
| 3 Either: | | | |
| 3.1 Patient's disease has progressed despite 3 injections w | | | |
| 3.2 Patient is unsuitable or contraindicated to treatment with | n anti-VEGF agents; a | and | |
| 4 Dexamethasone implants are to be administered not more freq maximum of 3 implants per eye per year. | uently than once eve | ry 4 month | s into each eye, and up to a |
| Continuation – Diabetic macular oedema | | | |
| Ophthalmologist | | | |
| Re-assessment required after 12 months Both: | | | |
| 1 Patient's vision is stable or has improved (prescriber determine | nd). and | | |
| Dexamethasone implants are to be administered not more freq maximum of 3 implants per eye per year. | | ry 4 month | s into each eye, and up to a |
| Initiation - Women of child bearing age with diabetic macular oed | lema | | |
| Ophthalmologist | | | |
| Re-assessment required after 12 months | | | |
| All of the following: | | | |
| Patients have diabetic macular oedema; and Patient has reduced visual acuity of between 6/9 – 6/48 with fu | nctional awaranosa a | fraduction | in vision: and |
| 3 Patient is of child bearing potential and has not yet completed a | | | ni visiun, anu |
| 4 Dexamethasone implants are to be administered not more freq | | rv 4 month | s into each eve, and up to a |
| maximum of 3 implants per eye per year. | | , | |
| Continuation - Women of child bearing age with diabetic macular | oedema | | |
| Ophthalmologist | | | |
| Pa accompany required after 12 menths | | | |

Re-assessment required after 12 months

All of the following:

- 1 Patient's vision is stable or has improved (prescriber determined); and
- 2 Patient is of child bearing potential and has not yet completed a family; and
- 3 Dexamethasone implants are to be administered not more frequently than once every 4 months into each eye, and up to a maximum of 3 implants per eye per year.

SENSORY ORGANS

| | Price | | Brand or |
|--|------------------------|------------|-------------------------|
| | (ex man. excl. G \$ | ST) Per | Generic Manufacturer |
| | Ŷ | 101 | Manalaotalor |
| FLUOROMETHOLONE | | | |
| Eye drops 0.1% | | 5 ml | FML |
| PREDNISOLONE ACETATE | | | |
| Eye drops 0.12% Eye drops 1% | 7.00 | 5 ml | Pred Forte |
| | 5.93 | 10 ml | Prednisolone- AFT |
| PREDNISOLONE SODIUM PHOSPHATE | 00.50 | 00 | Minima Decideirates |
| Eye drops 0.5%, single dose (preservative free) | | 20 dose | Minims Prednisolone |
| Non-Steroidal Anti-Inflammatory Drugs | | | |
| DICLOFENAC SODIUM | | | |
| Eye drops 0.1% | | 5 ml | Voltaren Ophtha |
| KETOROLAC TROMETAMOL Eye drops 0.5% | | | |
| | | | |
| Decongestants and Antiallergics | | | |
| Antiallergic Preparations | | | |
| LEVOCABASTINE | | | |
| Eye drops 0.05% | | | |
| LODOXAMIDE Eye drops 0.1% | 0.71 | 10 ml | Lomido |
| | 0.71 | 10 ml | Lomide |
| Eye drops 0.1% – 1% DV Oct-20 to 2022 | 2.20 | 5 ml | Olopatadine Teva |
| | 10.00 | | Patanol |
| (Patanol Eye drops 0.1% to be delisted 1 October 2020) | | | |
| SODIUM CROMOGLICATE Eye drops 2% – 1% DV Jan-20 to 2022 | 1 79 | 5 ml | Rexacrom |
| | | 5 111 | nexacrom |
| Decongestants | | | |
| | | 45 | Newberry Frida |
| Eye drops 0.1% | 4.15 | 15 ml | Naphcon Forte |
| Diagnostic and Surgical Preparations | | | |
| Diagnostic Dyes | | | |
| FLUORESCEIN SODIUM | | | |
| Eye drops 2%, single dose Inj 10%, 5 ml vial | 105.00 | 10 | Elucrossite |
| Ophthalmic strips 1 mg | 123.00 | 12 | Fluorescite |
| FLUORESCEIN SODIUM WITH LIGNOCAINE HYDROCHLORIDI | E | | |
| Eye drops 0.25% with lignocaine hydrochloride 4%, single dos | | | |
| LISSAMINE GREEN | | | |
| Ophthalmic strips 1.5 mg | | | |
| | | | |
| ROSE BENGAL SODIUM Ophthalmic strips 1% | | | |

t Item restricted (see \Rightarrow above); t Item restricted (see \Rightarrow below)

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e.g. Brand indicates brand example only. It is not a contracted product.

SENSORY ORGANS

| | | Price . excl. GST) \$ | Per | Brand or Generic Manufacturer |
|---|-----------------------|-----------------------------|-------------|--|
| Irrigation Solutions | | | | |
| MIXED SALT SOLUTION FOR EYE IRRIGATION Eye irrigation solution calcium chloride 0.048% with magnesium ch 0.03%, potassium chloride 0.075%, sodium acetate 0.39%, so chloride 0.64% and sodium citrate 0.17%, 15 ml dropper bottle Eye irrigation solution calcium chloride 0.048% with magnesium ch | odium e nloride | 5.00 | 15 ml | Balanced Salt Solution |
| 0.03%, potassium chloride 0.075%, sodium acetate 0.39%, so chloride 0.64% and sodium citrate 0.17%, 250 ml | dium | | | e.g. Balanced Salt Solution |
| Eye irrigation solution calcium chloride 0.048% with magnesium ch 0.03%, potassium chloride 0.075%, sodium acetate 0.39%, so chloride 0.64% and sodium citrate 0.17%, 500 ml bottle | dium | 10.50 | 500 ml | Balanced Salt Solution |
| Ocular Anaesthetics | | | | |
| OXYBUPROCAINE HYDROCHLORIDE Eye drops 0.4%, single dose PROXYMETACAINE HYDROCHLORIDE Eye drops 0.5% TETRACAINE [AMETHOCAINE] HYDROCHLORIDE Eye drops 0.5%, single dose Eye drops 1%, single dose | | | | |
| Viscoelastic Substances | | | | |
| HYPROMELLOSE Inj 2%, 1 ml syringe Inj 2%, 2 ml syringe | | | | |
| SODIUM HYALURONATE [HYALURONIC ACID] Inj 14 mg per ml, 0.85 ml syringe – 1% DV Oct-19 to 2022 Inj 14 mg per ml, 0.55 ml syringe – 1% DV Oct-19 to 2022 Inj 23 mg per ml, 0.6 ml syringe – 1% DV Oct-19 to 2022 Inj 10 mg per ml, 0.85 ml syringe – 1% DV Oct-19 to 2022 SODIUM HYALURONATE [HYALURONIC ACID] WITH CHONDROITI Inj 30 mg per ml with chondroitin sulphate 40 mg per ml, 0.35 ml s | IN SULP | 50.00 60.00 28.50 | 1 1 1 | Healon GV Healon GV Healon 5 Healon |
| and inj 10 mg sodium hyaluronate [hyaluronic acid] per ml, 0.4 syringe Inj 30 mg per ml with chondroitin sulphate 40 mg per ml, 0.5 ml sy and inj 10 mg sodium hyaluronate [hyaluronic acid] per ml, 0.5 | ringe | 64.00 | 1 | Duovisc |
| syringe Inj 30 mg per ml with chondroitin sulphate 40 mg per ml, 0.75 ml s | | | 1 1 | Duovisc Viscoat |
| Other | | | | |

Other

DISODIUM EDETATE

Inj 150 mg per ml, 20 ml ampoule

Inj 150 mg per ml, 20 ml vial

Inj 150 mg per ml, 100 ml vial

| | F | Price | | | Brand or |
|---|----------|-------------|------|----------------|----------------------------------|
| | (ex man. | excl. \$ | GST) | Per | Generic Manufacturer |
| RIBOFLAVIN 5-PHOSPHATE | | | | | |
| Soln trans epithelial riboflavin Inj 0.1% | | | | | |
| Inj 0.1% plus 20% dextran T500 | | | | | |
| Glaucoma Preparations | | | | | |
| Beta Blockers | | | | | |
| BETAXOLOL | | | | | |
| E TAXOLOL Eye drops 0.25% | | .11.80 |) | 5 ml | Betoptic S |
| Eye drops 0.5% | | 7.50 |) | 5 ml | Betoptic |
| | | | | 5 1 | A |
| Eye drops 0.25% Eye drops 0.5% | | | | 5 ml 5 ml | Arrow-Timolol Arrow-Timolol |
| Eye drops 0.5%, gel forming | | | | 2.5 ml | Timoptol XE |
| Carbonic Anhydrase Inhibitors | | | | | |
| ACETAZOLAMIDE | | | | | |
| Tab 250 mg | | .17.03 | 3 | 100 | Diamox |
| | | | | | |
| BRINZOLAMIDE Eye drops 1% | | | | | |
| DORZOLAMIDE | | | | | |
| Eye drops 2% | | | | | |
| DORZOLAMIDE WITH TIMOLOL Eye drops 2% with timolol 0.5% – 1% DV Jan-19 to 2021 | | 2.87 | 7 | 5 ml | Dortimopt |
| Miotics | | | | | |
| ACETYLCHOLINE CHLORIDE Inj 20 mg vial with diluent | | | | | |
| CARBACHOL | | | | | |
| Inj 150 mcg vial | | | | | |
| | | 4.00 | | 451 | la seta Osmina |
| Eye drops 1% Eye drops 2% | | | | 15 ml 15 ml | Isopto Carpine Isopto Carpine |
| Eye drops 2%, single dose | | | | | |
| Eye drops 4% | | 7.99 |) | 15 ml | Isopto Carpine |
| Prostaglandin Analogues | | | | | |
| BIMATOPROST | | 0.07 | 、 | 0! | Dimeter and Markin's |
| Eye drops 0.03% – 1% DV Feb-19 to 2021 | | 3.30 | J | 3 ml | Bimatoprost Multichem |
| ATANOPROST Eye drops 0.005% – 1% DV Apr-19 to 2021 | | 1.57 | 7 | 2.5 ml | Teva |
| | | | | | |
| TRAVOPROST | | | | | |

e.g. Brand indicates brand example only. It is not a contracted product.

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SENSORY ORGANS

| | Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|------------------------------------|--------------|-------------------------------------|
| Sympathomimetics | | | |
| APRACLONIDINE Eye drops 0.5% BRIMONIDINE TARTRATE Eye drops 0.2% BRIMONIDINE TARTRATE WITH TIMOLOL Eye drops 0.2% with timolol 0.5% | | 5 ml 5 ml | lopidine Arrow-Brimonidine |
| Mydriatics and Cycloplegics | | | |
| | | | |
| Anticholinergic Agents ATROPINE SULPHATE Eye drops 0.5% Eye drops 1%, single dose | | | |
| Eye drops 1% – 1% DV Oct-20 to 2023 CYCLOPENTOLATE HYDROCHLORIDE Eye drops 0.5%, single dose | 17.36 | 15 ml | Atropt |
| Eye drops 1% Eye drops 1%, single dose | 8.76 | 15 ml | Cyclogyl |
| TROPICAMIDE Eye drops 0.5% | 7.15 | 15 ml | Mydriacyl |
| Eye drops 0.5%, single dose Eye drops 1% Eye drops 1%, single dose | 8.66 | 15 ml | Mydriacyl |
| Sympathomimetics | | | |
| PHENYLEPHRINE HYDROCHLORIDE Eye drops 2.5%, single dose Eye drops 10%, single dose | | | |
| Ocular Lubricants | | | |
| CARBOMER Ophthalmic gel 0.3%, single dose Ophthalmic gel 0.2% | 8.25 | 30 | Poly Gel |
| CARMELLOSE SODIUM WITH PECTIN AND GELATINE Eye drops 0.5% Eye drops 0.5%, single dose Eye drops 1% Eye drops 1%, single dose | | | |
| HYPROMELLOSE Eye drops 0.5% | 3 92 | 15 ml | Methopt |
| HYPROMELLOSE WITH DEXTRAN Eye drops 0.3% with dextran 0.1% Eye drops 0.3% with dextran 0.1%, single dose | | 15 ml | Poly-Tears |
| MACROGOL 400 AND PROPYLENE GLYCOL Eye drops 0.4% with propylene glycol 0.3% preservative free, | single dose4.30 | 24 | Systane Unit Dose |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|---|----------|----------------------|------|-------|-------------------------------------|
| PARAFFIN LIQUID WITH SOFT WHITE PARAFFIN Eye oint 42.5% with soft white paraffin 57.3% | | | | | |
| PARAFFIN LIQUID WITH WOOL FAT Eye oint 3% with wool fat 3% | | 3.6 | 3 | 3.5 g | Poly-Visc |
| POLYVINYL ALCOHOL WITH POVIDONE Eye drops 1.4% with povidone 0.6%, single dose | | | | | |
| RETINOL PALMITATE Oint 138 mcg per g | | 3.8 | 0 | 5 g | VitA-POS |
| SODIUM HYALURONATE [HYALURONIC ACID] Eye drops 1 mg per ml | | .22.0 | 0 | 10 ml | Hylo-Fresh |
| Other Otological Preparations | | | | | |

ACETIC ACID WITH PROPYLENE GLYCOL

Ear drops 2.3% with propylene glycol 2.8%

DOCUSATE SODIUM

Ear drops 0.5%

| VARI | ous |
|------|-----|
|------|-----|

| (ex | P k man. | | GST) | _ | Brand or Generic |
|---|-------------|------|------|-----|-------------------------------|
| | | \$ | | Per | Manufacturer |
| Agents Used in the Treatment of Poisonings | | | | | |
| Antidotes | | | | | |
| ACETYLCYSTEINE Tab eff 200 mg Inj 200 mg per ml, 10 ml ampoule – 1% DV Sep-18 to 2021 | | 58.7 | 6 | 10 | DBL Acetylcysteine |
| AMYL NITRITE Liq 98% in 3 ml capsule | | | | | ,., |
| DIGOXIN IMMUNE FAB Inj 38 mg vial Inj 40 mg vial | | | | | |
| ETHANOL Liq 96% | | | | | |
| ETHANOL WITH GLUCOSE Inj 10% with glucose 5%, 500 ml bottle | | | | | |
| ETHANOL, DEHYDRATED Inj 100%, 5 ml ampoule Inj 96% | | | | | |
| FLUMAZENIL Inj 0.1 mg per ml, 5 ml ampoule – 1% DV Dec-18 to 2021 | 1 | 32.6 | 3 | 10 | Hameln |
| HYDROXOCOBALAMIN Inj 5 g vial Inj 2.5 g vial | | | | | |
| NALOXONE HYDROCHLORIDE | | | | | |
| Inj 400 mcg per ml, 1 ml ampoule - 1% DV Aug-18 to 2021 | | 22.6 | 0 | 5 | DBL Naloxone Hydrochloride |
| PRALIDOXIME IODIDE Inj 25 mg per ml, 20 ml ampoule | | | | | |
| SODIUM NITRITE Inj 30 mg per ml, 10 ml ampoule | | | | | |
| SODIUM THIOSULFATE Inj 250 mg per ml, 10 ml vial Inj 250 mg per ml. 50 ml vial Inj 500 mg per ml, 10 ml vial Inj 500 mg per ml, 20 ml ampoule | | | | | |
| SOYA OIL Inj 20%, 500 ml bag Inj 20%, 500 ml bottle | | | | | |
| Antitoxins | | | | | |
| | | | | | |

BOTULISM ANTITOXIN Inj 250 ml vial DIPHTHERIA ANTITOXIN Inj 10,000 iu vial

| Price | | Brand or |
|-------------------|-----|--------------|
| (ex man. excl. GS | | Generic |
| \$ | Per | Manufacturer |

Antivenoms

RED BACK SPIDER ANTIVENOM Inj 500 u vial

SNAKE ANTIVENOM

Inj 50 ml vial

Removal and Elimination

| Oral liq 200 mg per ml | 43.50 | 250 ml | Carbasorb-X |
|--|----------|--------|-------------|
| DEFERASIROX – Restricted see terms below | | | |
| Tab 125 mg dispersible | 276.00 | 28 | Exjade |
| I Tab 250 mg dispersible | 552.00 | 28 | Exjade |
| I Tab 500 mg dispersible | 1,105.00 | 28 | Exjade |
| - Destricted (DC1111) | - | | |

Restricted (RS1444)

Initiation

Haematologist *Re-assessment required after 2 years* All of the following:

1 The patient has been diagnosed with chronic iron overload due to congenital inherited anaemia; and

2 Deferasirox is to be given at a daily dose not exceeding 40 mg/kg/day; and

- 3 Any of the following:
 - 3.1 Treatment with maximum tolerated doses of deferiprone monotherapy or deferiprone and desferrioxamine combination therapy have proven ineffective as measured by serum ferritin levels, liver or cardiac MRI T2*; or
 - 3.2 Treatment with deferiprone has resulted in severe persistent vomiting or diarrhoea; or
 - 3.3 Treatment with deferiprone has resulted in arthritis; or
 - 3.4 Treatment with deferiprone is contraindicated due to a history of agranulocytosis (defined as an absolute neutrophil count (ANC) of < 0.5 cells per μL) or recurrent episodes (greater than 2 episodes) of moderate neutropenia (ANC 0.5 1.0 cells per μL).</p>

Continuation

Haematologist

Re-assessment required after 2 years Either:

- 1 For the first renewal following 2 years of therapy, the treatment has been tolerated and has resulted in clinical improvement in all three parameters namely serum ferritin. cardiac MRI T2* and liver MRI T2* levels; or
 - 2 For subsequent renewals, the treatment has been tolerated and has resulted in clinical stability or continued improvement in all three parameters namely serum ferritin, cardiac MRI T2* and liver MRI T2* levels.

DEFERIPRONE - Restricted see terms below

| t | Tab 500 mg | 7 | 100 | Ferriprox |
|---|------------------------|---|--------|-----------|
| | Oral liq 100 mg per ml | | 250 ml | Ferriprox |

➡ Restricted (RS1445)

Initiation

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Patient has been diagnosed with chronic iron overload due to congenital inherited anaemia or acquired red cell aplasia. DESFERBIOXAMINE MESILATE

| Inj 500 mg vial - 1% DV Mar-19 to 2021 | 84.53 | 10 | DBL Desferrioxamine |
|--|-------|----|---------------------|
| | | | Mesylate for Inj |
| | | | BP |

DICOBALT EDETATE

Inj 15 mg per ml, 20 ml ampoule

| | | | VANIOUS |
|---|----------------------------------|-----------|-------------------------------------|
| | Price (ex man. excl. GS \$ | T) Per | Brand or Generic Manufacturer |
| DIMERCAPROL | • | | |
| Inj 50 mg per ml, 2 ml ampoule | | | |
| DIMERCAPTOSUCCINIC ACID | | | |
| Cap 100 mg | | | e.g. PCNZ, Optimus |
| Cap Too Ing | | | Healthcare, |
| | | | Chemet |
| Cap 200 mg | | | e.g. PCNZ, Optimus |
| | | | Healthcare, |
| SODIUM CALCIUM EDETATE | | | Chemet |
| Inj 200 mg per ml, 2.5 ml ampoule | | | |
| Inj 200 mg per ml, 5 ml ampoule | | | |
| | | | |
| Antiseptics and Disinfectants | | | |
| CHLORHEXIDINE | | | |
| Soln 4% | | | |
| Soln 4%, | 1.86 | 50 ml | healthE |
| Soln 5% | | 500 ml | healthE |
| (healthE Soln 4%, to be delisted 1 November 2020) | | | |
| CHLORHEXIDINE WITH CETRIMIDE | | | |
| Crm 0.1% with cetrimide 0.5% | | | |
| Foaming soln 0.5% with cetrimide 0.5% | | | |
| CHLORHEXIDINE WITH ETHANOL | | | |
| Soln 0.5% with ethanol 70% | | | |
| Soln 2% with ethanol 70% | | | |
| Soln 0.5% with ethanol 70%, non-staining (pink) 100 ml | | 1 | healthE |
| Soln 2% with ethanol 70%, non-staining (pink) 100 ml | | 1 | healthE |
| Soln 0.5% with ethanol 70%, non-staining (pink) 25 ml | | 1 | healthE |
| Soln 0.5% with ethanol 70%, staining (red) 100 ml | | 1 1 | healthE healthE |
| Soln 2% with ethanol 70%, staining (red) 100 ml Soln 0.5% with ethanol 70%, non-staining (pink) 500 ml | | 1 | healthE |
| Soln 0.5% with ethanol 70%, staining (pink) 500 ml | | 1 | healthE |
| Soln 2% with ethanol 70%, staining (red) 500 ml | | 1 | healthE |
| (healthE Soln 0.5% with ethanol 70%, non-staining (pink) 100 ml to be a | | | Hourte |
| (healthE Soln 2% with ethanol 70%, non-staining (pink) 100 ml to be del | | , | |
| (healthE Soln 0.5% with ethanol 70%, staining (red) 100 ml to be deliste | | | |
| (healthE Soln 2% with ethanol 70%, staining (red) 100 ml to be delisted | | , | |
| (healthE Soln 0.5% with ethanol 70%, non-staining (pink) 500 ml to be o | | | |
| (healthE Soln 0.5% with ethanol 70%, staining (red) 500 ml to be deliste | d 1 November 20 | 20) | |
| (healthE Soln 2% with ethanol 70%, staining (red) 500 ml to be delisted | 1 November 2020 |)) | |
| IODINE WITH ETHANOL | | | |
| Soln 1% with ethanol 70% | | | |
| Soln 1% with ethanol 70%, 100 ml | | 1 | healthE |
| (healthE Soln 1% with ethanol 70%, 100 ml to be delisted 1 November 2 | 2020) | | |
| ISOPROPYL ALCOHOL | | | |
| Soln 70%, 500 ml | 5.65 | 1 | healthE |
| | | | |

VARIOUS

| | l (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------------|----------------------|------|--------|-------------------------------------|
| POVIDONE-IODINE | | | | | |
| Vaginal tab 200 mg | | | | | |
| → Restricted (RS1354) | | | | | |
| nitiation | | | | | |
| Rectal administration pre-prostate biopsy. | | | | | |
| Oint 10% - 1% DV Oct-20 to 2023 | | 7.4 | 0 | 65 g | Betadine |
| Soln 10% - 1% DV Nov-19 to 2021 | | 2.5 | 5 | 100 ml | Riodine |
| Soln 5% | | | | | |
| Soln 7.5% | | | | | |
| Soln 10%, - 1% DV Dec-19 to 2022 | | | | 15 ml | Riodine |
| | | 5.4 | 0 | 500 ml | Riodine |
| Pad 10% | | | | | |
| Swab set 10% | | | | | |
| POVIDONE-IODINE WITH ETHANOL | | | | | |
| Soln 10% with ethanol 30% | | | | | |
| Soln 10% with ethanol 70% | | | | | |
| SODIUM HYPOCHLORITE | | | | | |
| Soln | | | | | |
| | | | | | |
| Contrast Media | | | | | |
| Indirated X roy Contract Madia | | | | | |

Iodinated X-ray Contrast Media

| DIATRIZOATE MEGLUMINE WITH SODIUM AMIDOTRIZOATE | | |
|--|--------|----------------------|
| Oral liq 660 mg per ml with sodium amidotrizoate 100 mg per ml, 100 ml | | |
| bottle | 100 ml | Gastrografin |
| Inj 260 mg with sodium amidotrizoate 40 mg per ml, 250 ml bottle | 1 | Urografin |
| DIATRIZOATE SODIUM | | |
| Oral liq 370 mg per ml, 10 ml sachet156.12 | 50 | loscan |
| IODISED OIL | | |
| Inj 38% w/w (480 mg per ml), 10 ml ampoule | 1 | Lipiodol Ultra Fluid |
| IODIXANOL | | |
| Inj 270 mg per ml (iodine equivalent), 50 ml bottle | 10 | Visipaque |
| Inj 270 mg per ml (iodine equivalent), 30 ml bottle | 10 | Visipaque |
| Inj 320 mg per ml (iodine equivalent), 50 ml bottle | 10 | Visipaque |
| Inj 320 mg per ml (iodine equivalent), 100 ml bottle | 10 | Visipaque |
| Inj 320 mg per ml (iodine equivalent), 200 ml bottle | 10 | Visipaque |
| IOHEXOL | | |
| Inj 240 mg per ml (iodine equivalent), 50 ml bottle | 10 | Omnipaque |
| Inj 300 mg per ml (iodine equivalent), 30 ml bottle | 10 | Omnipaque |
| Inj 300 mg per ml (iodine equivalent), 20 ml bottle | 10 | Omnipaque |
| Inj 300 mg per ml (iodine equivalent), 30 ml bottle | 10 | Omnipaque |
| Inj 350 mg per ml (iodine equivalent), 700 ml bottle | 10 | Omnipaque |
| Inj 350 mg per ml (iodine equivalent), 50 ml bottle | 10 | Omnipaque |
| Inj 350 mg per ml (iodine equivalent), 75 ml bottle | 10 | Omnipaque |
| Inj 350 mg per ml (iodine equivalent), 100 ml bottle | 10 | Omnipaque |
| Inj 350 mg per ml (iodine equivalent), 200 ml bottle | 10 | Omnipaque |
| | 10 | epaquo |

| | Price | | Brand or |
|---|-------------------------|--|--|
| | (ex man. excl. GS \$ | T) Per | Generic Manufacturer |
| Non-iodinated X-ray Contrast Media | | | |
| BARIUM SULPHATE | | | |
| Powder for oral liq 20 mg per g (2% w/w), 22.1 g sachet | 507.50 | 50 | E-Z-Cat Dry |
| Oral liq 400 mg per ml (40% w/v, 30% w/w), bottle | | 148 g | Varibar - Thin Liquid |
| Oral liq 600 mg per g (60% w/w), tube | | 454 g | E-Z-Paste |
| Oral liq 400 mg per ml (40% w/v), bottle | | 250 ml | Varibar - Honey |
| | 38.40 | 240 ml | Varibar - Nectar |
| | 145.04 | 230 ml | Varibar - Pudding |
| Enema 1,250 mg per ml (125% w/v), 500 ml bag | | 12 | Liquibar |
| Oral liq 22 mg per g (2.2% w/w), 250 ml bottle | | 24 | CT Plus+ |
| Oral liq 22 mg per g (2.2% w/w), 450 ml bottle | | 24 | CT Plus+ |
| Oral liq 1 mg per ml (0.1% w/v, 0.1% w/w), 450 ml bottle | | 24 | VoLumen |
| Oral liq 20.9 mg per ml (2.1% w/v, 2% w/w), 250 ml bottle | | 24 | Readi-CAT 2 |
| Powder for oral soln 97.65% w/w, 300 g bottle | | 24 | X-Opaque-HD |
| Oral liq 400 mg per ml (40% w/v, 30% w/w), 20 ml bottle | | 3 | Tagitol V |
| Oral liq 1,250 mg per ml (125% w/v), 2,000 ml bottle | 91.77 | 1 | Liquibar |
| 3ARIUM SULPHATE WITH SODIUM BICARBONATE | | | |
| Grans eff 382.2 mg per g with sodium bicarbonate 551.3 mg per g, | 4 g | | |
| sachet | | 50 | E-Z-Gas II |
| CITRIC ACID WITH SODIUM BICARBONATE | | | |
| Powder 382.2 mg per g with sodium bicarbonate 551.3 mg per g, 4 | g | | |
| sachet | 0 | | e.g. E-Z-GAS II |
| Paramagnetic Contrast Media | | | |
| GADOBENIC ACID | | | |
| Inj 334 mg per ml, 10 ml vial | | 10 | Multihance |
| Inj 334 mg per ml, 20 ml vial | 636.28 | 10 | Multihance |
| GADOBUTROL | | | |
| lnj 1 mmol per ml, 15 ml vial | | | |
| | | | |
| Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 5 ml prefilled | | | |
| Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 5 ml prefilled syringe | | 5 | Gadovist 1.0 |
| | | 5 | Gadovist 1.0 |
| syringe | | 5 5 | Gadovist 1.0 Gadovist 1.0 |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled | | | |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled | | | |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe | | 5 | Gadovist 1.0 |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe | | 5 | Gadovist 1.0 |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe GADODIAMIDE | | 5 10 | Gadovist 1.0 Gadovist 1.0 |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe GADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe | | 5 10 10 | Gadovist 1.0 Gadovist 1.0 Omniscan |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe GADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial | | 5 10 10 10 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe GADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml prefilled syringe | | 5 10 10 10 10 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe SADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 15 ml prefilled syringe SADODTERIC ACID | | 5 10 10 10 10 10 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan Omniscan |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe SADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 15 ml prefilled syringe SADOTERIC ACID Inj 279.32 mg per ml (0.5 mmol per ml), 10 ml prefilled syringe | | 5 10 10 10 10 10 10 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan Omniscan Dotarem |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe SADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml prefilled syringe SADOTERIC ACID Inj 279.32 mg per ml (0.5 mmol per ml), 10 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml bottle | | 5 10 10 10 10 10 10 1 1 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan Dotarem Dotarem |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 15 ml prefilled syringe GADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 15 ml prefilled syringe GADOTERIC ACID Inj 279.32 mg per ml (0.5 mmol per ml), 10 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml bottle Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml prefilled syringe | | 5 10 10 10 10 10 10 1 1 1 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan Dotarem Dotarem Dotarem |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe SADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 15 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 10 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml bottle Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 20 ml prefilled syringe | | 5 10 10 10 10 10 10 1 1 1 1 1 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan Dotarem Dotarem Dotarem Dotarem |
| syringe Inj 604.72 mg per ml (equivalent to 1 mmol per ml), 7.5 ml prefilled syringe SADODIAMIDE Inj 287 mg per ml, 10 ml prefilled syringe Inj 287 mg per ml, 10 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 5 ml vial Inj 287 mg per ml, 15 ml prefilled syringe SADOTERIC ACID Inj 279.32 mg per ml (0.5 mmol per ml), 10 ml prefilled syringe Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml bottle Inj 279.32 mg per ml (0.5 mmol per ml), 15 ml prefilled syringe | | 5 10 10 10 10 10 10 1 1 1 | Gadovist 1.0 Gadovist 1.0 Omniscan Omniscan Omniscan Dotarem Dotarem Dotarem |

VARIOUS

| | | Price excl. GS \$ | T) Per | Brand or Generic Manufacturer |
|--|----|-------------------------|-----------|-------------------------------------|
| GADOXETATE DISODIUM | | | | |
| Inj 181.43 mg per ml (equivalent to 0.25 mmol per ml), 10 ml p syringe | | 300.00 | 1 | Primovist |
| IEGLUMINE GADOPENTETATE | | | | |
| Inj 469 mg per ml, 10 ml prefilled syringe Inj 469 mg per ml, 10 ml vial | | | 5 10 | Magnevist Magnevist |
| IEGLUMINE IOTROXATE | | | | - |
| Inj 105 mg per ml, 100 ml bottle | 1 | 150.00 | 100 ml | Biliscopin |
| Ultrasound Contrast Media | | | | |
| PERFLUTREN | | | | |
| Inj 1.1 mg per ml, 1.5 ml vial | 1 | 180.00 | 1 | Definity |
| | 7 | 720.00 | 4 | Definity |
| Diagnostic Agents | | | | |
| ARGININE | | | | |
| Inj 50 mg per ml, 500 ml bottle Inj 100 mg per ml, 300 ml bottle | | | | |
| IISTAMINE ACID PHOSPHATE | | | | |
| Nebuliser soln 0.6%, 10 ml vial | | | | |
| Nebuliser soln 2.5%, 10 ml vial | | | | |
| Nebuliser soln 5%, 10 ml vial | | | | |
| /ANNITOL | | | | |
| Powder for inhalation | | | | e.g. Aridol |
| IETHACHOLINE CHLORIDE Powder 100 mg | | | | |
| | | | | |
| Inj 100 u ampoule | | | | |
| SINCALIDE | | | | |
| Inj 5 mcg per vial | | | | |
| Diagnostic Dyes | | | | |
| SONNEY'S BLUE DYE | | | | |
| Soln | | | | |
| | | | | |
| Inj 4 mg per ml, 5 ml ampoule | | | | |
| Inj 8 mg per ml, 5 ml ampoule | | | | |
| NDOCYANINE GREEN | | | | |
| Inj 25 mg vial | | | | |
| IETHYLTHIONINIUM CHLORIDE [METHYLENE BLUE] Inj 5 mg per ml, 10 ml ampoule | c. | 240 35 | 5 | Proveblue |
| | | | 5 | TIOVEDIUE |
| | | | | |
| ATENT BLUE V Inj 2.5%, 2 ml ampoule | , | 140.00 | 5 | Obex Medical |

e.g. Brand indicates brand example only. It is not a contracted product.

VARIOUS

Price Brand or (ex man. excl. GST) Generic \$ Per Manufacturer

Irrigation Solutions

CHLORHEXIDINE WITH CETRIMIDE

Irrigation soln 0.015% with cetrimide 0.15%, 500 ml bottle

→ Restricted (RS1683)

Initiation

Re-assessment required after 3 months All of the following:

- 1 Patient has burns that are greater than 30% of total body surface area (BSA); and
- 2 For use in the perioperative preparation and cleansing of large burn areas requiring debridement/skin grafting; and
- 3 The use of 30 ml ampoules is impractical due to the size of the area to be covered.

Continuation

Re-assessment required after 3 months

The treatment remains appropriate for the patient and the patient is benefiting from the treatment.

| Irrigation soln 0.015% with cetrimide 0.15%, 30 ml ampoule - 1% DV Aug-18 to 2021 | 30 | Pfizer |
|--|----|------------------------------|
| GLYCINE | | |
| Irrigation soln 1.5%, 3,000 ml bag - 1% DV Sep-18 to 2021 | 4 | B Braun |
| SODIUM CHLORIDE | | |
| Irrigation soln 0.9%, 3,000 ml bag – 1% DV Sep-18 to 2021 | 4 | B Braun |
| Irrigation soln 0.9%, 30 ml ampoule - 1% DV Sep-18 to 2021 | 20 | Interpharma |
| Irrigation soln 0.9%, 1,000 ml bottle – 1% DV Jun-18 to 2021 | 10 | Baxter Sodium |
| | | Chloride 0.9% |
| Irrigation soln 0.9%, 250 ml bottle – 1% DV Aug-18 to 2021 | 12 | Fresenius Kabi |
| WATER | | |
| Irrigation soln, 3,000 ml bag – 1% DV Sep-18 to 2021 | 4 | B Braun |
| Irrigation soln, 1,000 ml bottle - 1% DV Jun-18 to 2021 | 10 | Baxter Water for |
| Irrigation soln, 250 ml bottle - 1% DV Aug-18 to 2021 | 12 | Irrigation Fresenius Kabi |

Surgical Preparations

BISMUTH SUBNITRATE AND IODOFORM PARAFFIN

Paste

DIMETHYL SULFOXIDE Soln 50% Soln 99%

PHENOL

Inj 6%, 10 ml ampoule

PHENOL WITH IOXAGLIC ACID

Inj 12%, 10 ml ampoule

TROMETAMOL

Inj 36 mg per ml, 500 ml bottle

| | (ex man. | Price . excl. \$ | GST) | Per | Bran Gene Manu | |
|---|---------------------|------------------------|------|-----|----------------------|---|
| Cardioplegia Solutions | | | | | | |
| ELECTROLYTES Inj 15 mmol/l sodium chloride, 9 mmol/l potassium chloride, 1 m potassium hydrogen 2-ketoglutarate, 4 mmol/l magnesium 18 mmol/l histidine hydrochloride, 180 mmol/l histidine, 2 m tryptophan, 30 mmol/l mannitol, 0.015 mmol/l calcium chlor | chloride, Imol/l | | | | | |
| 1,000 ml bag Inj aspartic acid 10.43 mg per ml, citric acid 0.22476 mg per ml acid 11.53 mg per ml, sodium phosphate 0.1725 mg per m potassium chloride 2.15211 mg per ml, sodium citrate 1.80 per ml, sodium hydroxide 6.31 mg per ml and trometamol | glutamic | | | | e.g. | Custodiol-HTK |
| 11.2369 mg per ml, 364 ml bag | | | | | e.g. | Cardioplegia Enriched Paed. Soln. |
| Inj aspartic acid 8.481 mg per ml, citric acid 0.8188 mg per ml, acid 9.375 mg per ml, sodium phosphate 0.6285 mg per m potassium chloride 2.5 mg per ml, sodium citrate 6.585 mg sodium hydroxide 5.133 mg per ml and trometamol 9.097 r ml, 527 ml bag | , per ml, | | | | e.g. | Cardioplegia |
| Inj citric acid 0.07973 mg per ml, sodium phosphate 0.06119 m potassium chloride 2.181 mg per ml, sodium chloride 1.786 sodium citrate 0.6412 mg per ml and trometamol 5.9 mg per | s mg ml, | | | | 0 | Enriched Solution |
| 523 ml bag | | | | | e.g. | Cardioplegia Base Solution |
| Inj 110 mmol/l sodium, 16 mmol/l potassium, 1.2 mmol/l calciun 16 mmol/l magnesium and 160 mmol/l chloride, 1,000 ml b | | | | | e.g. | Cardioplegia Solution AHB7832 |
| Inj 143 mmol/l sodium, 16 mmol/l potassium, 16 mmol/l magnes 1.2 mmol/l calcium, 1,000 ml bag | sium and | | | | e.g. | Cardioplegia Electrolyte Solution |
| MONOSODIUM GLUTAMATE WITH SODIUM ASPARTATE Inj 42.68 mg with sodium aspartate 39.48 mg per ml, 250 ml bo MONOSODIUM L-ASPARTATE Inj 14 mmol per 10 ml, 10 ml | ttle | | | | | · |

Cold Storage Solutions

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SODIUM WITH POTASSIUM Inj 29 mmol/l with potassium 125 mmol/l, 1,000 ml bag

EXTEMPORANEOUSLY COMPOUNDED PREPARATIONS

| (ex m | Price an. excl | GST) | | Brand or Generic |
|--|-------------------|------|--------|---------------------|
| | \$ | | Per | Manufacturer |
| Extemporaneously Compounded Preparations | | | | |
| ACETIC ACID | | | | |
| Liq | | | | |
| ALUM Powder BP | | | | |
| ARACHIS OIL [PEANUT OIL] Liq | | | | |
| ASCORBIC ACID Powder | | | | |
| BENZOIN | | | | |
| Tincture compound BP BISMUTH SUBGALLATE Powder | | | | |
| BORIC ACID Powder | | | | |
| CARBOXYMETHYLCELLULOSE Soln 1.5% | | | | |
| CETRIMIDE Soln 40% | | | | |
| CHLORHEXIDINE GLUCONATE Soln 20 % | | | | |
| CHLOROFORM Liq BP | | | | |
| CITRIC ACID Powder BP | | | | |
| CLOVE OIL Liq | | | | |
| COAL TAR Soln BP - 1% DV Nov-19 to 2022 | 36.2 | 25 | 200 ml | Midwest |
| CODEINE PHOSPHATE Powder | | | | |
| COLLODION FLEXIBLE | | | | |
| COMPOUND HYDROXYBENZOATE Soln – 1% DV Aug-19 to 2022 | 30.0 | 00 | 100 ml | Midwest |
| CYSTEAMINE HYDROCHLORIDE Powder | | | - | |
| DISODIUM HYDROGEN PHOSPHATE WITH SODIUM DIHYDROGEN PHO Inj 37.46 mg with sodium dihydrogen phosphate 47.7 mg in 1.5 ml ampoule | SPHAT | E | | |
| DITHRANOL Powder | | | | |
| GLUCOSE [DEXTROSE] Powder | | | | |

EXTEMPORANEOUSLY COMPOUNDED PREPARATIONS

| | Price | | Brand or |
|--|-------------------|--------|---------------------|
| | (ex man. excl. GS | | Generic |
| | \$ | Per | Manufacturer |
| GLYCERIN WITH SODIUM SACCHARIN | | | |
| Suspension - 1% DV Jul-19 to 2022 | | 473 ml | Ora-Sweet SF |
| | | | |
| GLYCERIN WITH SUCROSE | 00.05 | 470 | . |
| Suspension - 1% DV Jul-19 to 2022 | | 473 ml | Ora-Sweet |
| GLYCEROL | | | |
| Liq - 1% DV Oct-20 to 2023 | 3.23 | 500 ml | healthE Glycerol BP |
| | | | Liquid |
| HYDROCORTISONE | | | |
| Powder | 40.05 | 25 a | ABM |
| | | 25 g | ADIVI |
| LACTOSE | | | |
| Powder | | | |
| MAGNESIUM HYDROXIDE | | | |
| Paste | | | |
| Suspension | | | |
| • | | | |
| MENTHOL | | | |
| Crystals | | | |
| METHADONE HYDROCHLORIDE | | | |
| Powder | | | |
| | | | |
| METHYL HYDROXYBENZOATE | 0.00 | 05 - | Midure et |
| Powder - 1% DV Jul-19 to 2022 | 8.98 | 25 g | Midwest |
| METHYLCELLULOSE | | | |
| Powder - 1% DV Jul-19 to 2022 | | 100 g | Midwest |
| Suspension - 1% DV Jul-19 to 2022 | | 473 ml | Ora-Plus |
| METHYLCELLULOSE WITH GLYCERIN AND SODIUM SACCHARII | N | | |
| Suspension – 1% DV Jul-19 to 2022. | | 473 ml | Ora-Blend SF |
| | | 470111 | |
| METHYLCELLULOSE WITH GLYCERIN AND SUCROSE | | | |
| Suspension - 1% DV Jul-19 to 2022 | | 473 ml | Ora-Blend |
| OLIVE OIL | | | |
| Liq | | | |
| PARAFFIN | | | |
| | | | |
| Liq | | | |
| PHENOBARBITONE SODIUM | | | |
| Powder | | | |
| PHENOL | | | |
| Liq | | | |
| • | | | |
| PILOCARPINE NITRATE | | | |
| Powder | | | |
| POLYHEXAMETHYLENE BIGUANIDE | | | |
| Liq | | | |
| POVIDONE K30 | | | |
| | | | |
| Powder | | | |
| SALICYLIC ACID | | | |
| Powder | | | |
| SILVER NITRATE | | | |
| Crystals | | | |
| - | | | |
| SODIUM BICARBONATE | 10.05 | 500 | NP |
| Powder BP – 1% DV Jan-20 to 2022 | 10.05 | 500 g | Midwest |
| | | | |

t Item restricted (see → above); t Item restricted (see → below)

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e.g. Brand indicates brand example only. It is not a contracted product.

EXTEMPORANEOUSLY COMPOUNDED PREPARATIONS

| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|--|-----------------------------------|----------|-------------------------------------|
| SODIUM CITRATE Powder | | | |
| SODIUM METABISULFITE Powder | | | |
| STARCH Powder | | | |
| SULPHUR Precipitated Sublimed | | | |
| SYRUP Liq (pharmaceutical grade) – 1% DV Jan-20 to 2022 | | 500 ml | Midwest |
| THEOBROMA OIL Oint | | | |
| TRI-SODIUM CITRATE Crystals | | | |
| TRICHLORACETIC ACID Grans | | | |
| UREA Powder BP | | | |
| WOOL FAT Oint, anhydrous | | | |
| XANTHAN Gum 1% | | | |
| ZINC OXIDE Powder | | | |

Price Bi (ex man. excl. GST) G \$ Per M

Brand or Generic Manufacturer

Food Modules

Carbohydrate

➡ Restricted (RS1467)

Initiation – Use as an additive

Any of the following:

- 1 Cystic fibrosis; or
- 2 Chronic kidney disease; or
- 3 Cancer in children; or
- 4 Cancers affecting alimentary tract where there are malabsorption problems in patients over the age of 20 years; or
- 5 Faltering growth in an infant/child; or
- 6 Bronchopulmonary dysplasia; or
- 7 Premature and post premature infant; or
- 8 Inborn errors of metabolism.

Initiation – Use as a module

For use as a component in a modular formula made from at least one nutrient module and at least one further product listed in Section D of the Pharmaceutical Schedule or breast milk.

Note: Patients are required to meet any Special Authority criteria associated with all of the products used in the modular formula.

CARBOHYDRATE SUPPLEMENT - Restricted see terms above

- 1 Powder 95 g carbohydrate per 100 g, 368 g can
- Powder 96 g carbohydrate per 100 g, 400 g can

e.g. Polycal

Fat

➡ Restricted (RS1468)

Initiation - Use as an additive

Any of the following:

- 1 Patient has inborn errors of metabolism; or
- 2 Faltering growth in an infant/child; or
- 3 Bronchopulmonary dysplasia; or
- 4 Fat malabsorption; or
- 5 Lymphangiectasia; or
- 6 Short bowel syndrome; or
- 7 Infants with necrotising enterocolitis; or
- 8 Biliary atresia; or
- 9 For use in a ketogenic diet; or
- 10 Chyle leak; or
- 11 Ascites; or

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12 Patient has increased energy requirements, and for whom dietary measures have not been successful.

Initiation – Use as a module

For use as a component in a modular formula made from at least one nutrient module and at least one further product listed in Section D of the Pharmaceutical Schedule or breast milk.

Note: Patients are required to meet any Special Authority criteria associated with all of the products used in the modular formula.

LONG-CHAIN TRIGLYCERIDE SUPPLEMENT - Restricted see terms above

- 1 Liquid 50 g fat per 100 ml, 200 ml bottle
- Liquid 50 g fat per 100 ml, 500 ml bottle

e.g. Calogen e.g. Calogen

| | SI | PECIAL FOODS |
|--|--------------------|---|
| Price (ex man. excl. \$ | GST) Per | Brand or Generic Manufacturer |
| MEDIUM-CHAIN TRIGLYCERIDE SUPPLEMENT – Restricted see terms on the pre Liquid 50 g fat per 100 ml, 250 ml bottle Liquid 95 g fat per 100 ml, 500 ml bottle WALNUT OIL – Restricted see terms on the previous page Liq | vious page | e.g. Liquigen e.g. MCT Oil |
| Protein | | |
| → Restricted (RS1469) Initiation – Use as an additive Either: Protein losing enteropathy; or High protein needs. Initiation – Use as a module For use as a component in a modular formula made from at least one nutrient module Section D of the Pharmaceutical Schedule or breast milk Note: Patients are required to meet any Special Authority criteria associated with all of PROTEIN SUPPLEMENT – Restricted see terms above Powder 5 g protein, 0.67 g carbohydrate and 0.6 g fat per 6.6 g, 275 g can Powder 6 g protein per 7 g, can | of the products us | |
| Other Supplements | | |
| BREAST MILK FORTIFIER Powder 0.2 g protein, 0.7 g carbohydrate and 0.02 g fat per 1 g sachet Powder 0.5 g protein, 1.2 g carbohydrate and 0.08 g fat per 2 g sachet Powder 0.6 g protein and 1.4 g carbohydrate per 2.2 g sachet CARBOHYDRATE AND FAT SUPPLEMENT - Restricted see terms below Powder 72.7 g carbohydrate and 22.3 g fat per 100 g, 400 g can Restricted (RS1212) Initiation Both: Infant or child aged four years or under; and Any of the following: Cancer in children; or Faltering growth; or Fornchopulmonary dysplasia; or Premature and post premature infants. | | e.g. FM 85 e.g. S26 Human Milk Fortifier e.g. Nutricia Breast Milk Fortifer e.g. Super Soluble Duocal |

| Price | | Brand or |
|---------------------|-----|--------------|
| (ex man. excl. GST) | | Generic |
| \$ | Per | Manufacturer |

Food/Fluid Thickeners

NOTE:

While pre-thickened drinks and supplements have not been included in Section H, DHB hospitals may continue to use such products for patients with dysphagia, provided that:

- use was established prior to 1 July 2013; and
- the product has not been specifically considered and excluded by PHARMAC; and
- use of the product conforms to any applicable indication restrictions for similar products that are listed in Section H (for example, use of thickened high protein products should be in line with the restriction for high protein oral feed in Section H).

PHARMAC intends to make a further decision in relation to pre-thickened drinks and supplements in the future, and will notify of any change to this situation.

CAROB BEAN GUM WITH MAIZE STARCH AND MALTODEXTRIN

| | Powder | e.g. | Feed Thickener Karicare Aptamil |
|---|---|------|------------------------------------|
| | SUAR GUM Powder IAIZE STARCH | e.g. | Guarcol |
| n | Powder | e.g. | Resource Thicken Up; Nutilis |
| | IALTODEXTRIN WITH XANTHAN GUM Powder | e.g. | Instant Thick |
| Ν | IALTODEXTRIN WITH XANTHAN GUM AND ASCORBIC ACID Powder | e.g. | Easy Thick |

Metabolic Products

➡ Restricted (RS1232)

Initiation

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Any of the following:

- 1 For the dietary management of homocystinuria, maple syrup urine disease, phenylketonuria (PKU), glutaric aciduria, isovaleric acidaemia, propionic acidaemia, methylmalonic acidaemia, tyrosinaemia or urea cycle disorders; or
- 2 Patient has adrenoleukodystrophy; or
- 3 For use as a supplement to the Ketogenic diet in patients diagnosed with epilepsy.

Glutaric Aciduria Type 1 Products

AMINO ACID FORMULA (WITHOUT LYSINE AND LOW TRYPTOPHAN) - Restricted see terms above

- Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g fibre per 100 g, 400 g can
- Powder 25 g protein and 51 g carbohydrate per 100 g, 500 g can

- e.g. GA1 Anamix Infant
- e.g. XLYS Low TRY Maxamaid

| | | | SPECIAL FOODS |
|---|-----------------------------------|------------|---|
| | Price (ex man. excl. GST \$ |) Per | Brand or Generic Manufacturer |
| Homocystinuria Products | | | |
| AMINO ACID FORMULA (WITHOUT METHIONINE) - Restricted set Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g fib 100 g, 400 g can Powder 25 g protein and 51 g carbohydrate per 100 g, 500 g can Powder 39 g protein and 34 g carbohydrate per 100 g, 500 g can Liquid 8 g protein, 7 g carbohydrate, 3.8 g fat and 0.25 g fibre per 100 ml, 125 ml bottle | re per | ous page | e.g. HCU Anamix Infant e.g. XMET Maxamaid e.g. XMET Maxamum e.g. HCU Anamix Junior LQ |
| Isovaleric Acidaemia Products | | | |
| AMINO ACID FORMULA (WITHOUT LEUCINE) - Restricted see tel Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g fib 100 g, 400 g can Powder 25 g protein and 51 g carbohydrate per 100 g, 500 g can Powder 39 g protein and 34 g carbohydrate per 100 g, 500 g can | | oage | e.g. IVA Anamix Infant e.g. XLEU Maxamaid e.g. XLEU Maxamum |
| Maple Syrup Urine Disease Products | | | |
| AMINO ACID FORMULA (WITHOUT ISOLEUCINE, LEUCINE AND V Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g fib 100 g, 400 g cap | , | d see term | |
| 100 g, 400 g can Powder 39 g protein and 34 g carbohydrate per 100 g, 500 g can Liquid 8 g protein, 7 g carbohydrate, 3.8 g fat and 0.25 g fibre per 100 ml, 125 ml bottle | | | e.g. MSUD Anamix Infant e.g. MSUD Maxamum e.g. MSUD Anamix Junior LQ |

| | F (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|---------------|----------------------|------|--------|---------------------------------------|
| Phenylketonuria Products | | | | | |
| MINO ACID FORMULA (WITHOUT PHENYLALANINE) - Restricted | see tern | ns <mark>on</mark> | page | 234 | |
| Tab 8.33 mg | | | | | e.g. Phlexy-10 |
| Powder 20 g protein, 2.5 g carbohydrate and 0.22 g fibre per 27.8 g | l | | | | |
| sachet | | | | | e.g. PKU Lophlex |
| | | | | | Powder |
| | _ | | | | (unflavoured) |
| Powder 36 g protein, 32 g carbohydrate and 12.5 g fat per 100 g, 3 | 6 g | | | | 5.4.1 |
| sachet | | | | | e.g. PKU Anamix Junio |
| Bounder 10.1 a protein 40.5 a compensation 00 a fat and 5.0 a fibra | | | | | (van/choc/unfl) |
| Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g fibre 100 g, 400 g can | per | | | | e.g. PKU Anamix Infar |
| Powder 39 g protein and 34 g carbohydrate per 100 g, 500 g can | | | | | e.g. XP Maxamum |
| Powder 8.33 g protein and 8.8 g carbohydrate per 100 g, 600 g carb | | | | | e.g. Phlexy-10 |
| Liquid 10 g protein, 4.4 g carbohydrate and 0.25 g fibre per 100 ml, | | | | | o.g. Thiony To |
| 62.5 ml bottle | | | | | e.g. PKU Lophlex LQ |
| Liquid 20 g protein, 8.8 g carbohydrate and 0.34 g fibre per 100 ml, | | | | | |
| 125 ml bottle | | | | | e.g. PKU Lophlex LQ |
| Liquid 8 g protein, 7 g carbohydrate, 3.8 g fat and 0.25 g fibre per | | | | | , |
| 100 ml, bottle | | . 13.1 | 0 | 125 ml | PKU Anamix Junior LQ |
| | | | | | (Berry) |
| | | | | | PKU Anamix Junior LQ |
| | | | | | (Orange) |
| | | | | | PKU Anamix Junior LQ (Unflavoured) |
| Liquid 16 g protein, 7 g carbohydrate and 0.27 g fibre per 100 ml, 1 | 25 ml | | | | (Offilavoureu) |
| bottle | 20111 | | | | e.g. PKU Lophlex LQ |
| Liquid 16 g protein, 7 g carbohydrate and 0.27 g fibre per 100 ml, | | | | | e.g. The Lophick Lor |
| 62.5 ml bottle | | | | | e.g. PKU Lophlex LQ |
| Liquid 16 g protein, 7 g carbohydrate and 0.4 g fibre per 100 ml, 12 | 5 ml | | | | g |
| bottle | | | | | e.g. PKU Lophlex LQ |
| Liquid 16 g protein, 7 g carbohydrate and 0.4 g fibre per 100 ml, 62 | .5 ml | | | | - , |
| bottle | | | | | e.g. PKU Lophlex LQ |
| Liquid 6.7 g protein, 5.1 g carbohydrate and 2 g fat per 100 ml, 250 | ml | | | | |
| carton | | | | | e.g. Easiphen |
| Semi-solid 18.3 g protein, 18.5 g carbohydrate and 0.92 g fibre per | | | | | |
| 100 g, 109 g pot | | | | | e.g. PKU Lophlex |
| | | | | | Sensations |
| | | | | | 20 (berries) |

Propionic Acidaemia and Methylmalonic Acidaemia Products

| ٨N | IINO ACID FORMULA (WITHOUT ISOLEUCINE, METHIONINE, THREONINE AND VALINE) - | Restricted see terms on |
|----|--|-------------------------------|
| | ye 234 | |
| τ | Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g fibre per | |
| | 100 g, 400 g can | e.g. MMA/PA Anamix |
| t | Powder 25 g protein and 51 g carbohydrate per 100 g, 500 g can | Infant e.g. XMTVI Maxamaid |
| | Powder 39 g protein and 34 g carbohydrate per 100 g, 500 g can | e.g. XMTVI Maxamum |

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SPECIAL FOODS

| | (ex man | Price excl. \$ | GST) | Per | Bran Gene Mani | |
|--|----------|----------------------|--------|-----------|----------------------|--------------------------------|
| Protein Free Supplements | | | | | | |
| PROTEIN FREE SUPPLEMENT – Restricted see terms on page 2 Powder nil added protein and 67 g carbohydrate per 100 g, 400 | | | | | e.g. | Energivit |
| Tyrosinaemia Products | | | | | | |
| AMINO ACID FORMULA (WITHOUT PHENYLALANINE AND TYRO Powder 36 g protein, 32 g carbohydrate and 12.5 g fat per 100 g | , | estric | ted se | e terms o | n page | e 234 |
| sachet | | | | | e.g. | TYR Anamix Junio |
| Powder 13.1 g protein, 49.5 g carbohydrate, 23 g fat and 5.3 g f | ibre per | | | | | |
| 100 g, 400 g can Powder 25 g protein and 51 g carbohydrate per 100 g, 400 g ca | n | | | | • | TYR Anamix Infar XPHEN, TYR |
| | | | | | e.y. | Maxamaid |
| Liquid 8 g protein, 7 g carbohydrate, 3.8 g fat and 0.25 g fibre p | er | | | | | |
| 100 ml, 125 ml bottle | | | | | e.g. | TYR Anamix Junio LQ |
| Urea Cycle Disorders Products | | | | | | |
| AMINO ACID SUPPLEMENT – Restricted see terms on page 234 | | | | | | |
| Powder 25 g protein and 65 g carbohydrate per 100 g, 200 g ca | n | | | | | Dialamine |
| Powder 79 g protein per 100 g, 200 g can | | | | | e.g. | Essential Amino Acid Mix |
| X-Linked Adrenoleukodystrophy Products | | | | | | |
| GLYCEROL TRIERUCATE – Restricted see terms on page 234 Liquid, 1,000 ml bottle | | | | | | |
| GLYCEROL TRIOLEATE – Restricted see terms on page 234 | | | | | | |

1 Liquid, 500 ml bottle

Specialised Formulas

Diabetic Products

→ Restricted (RS1215)

Initiation

Any of the following:

- 1 For patients with type I or type II diabetes suffering weight loss and malnutrition that requires nutritional support; or
- 2 For patients with pancreatic insufficiency; or
- 3 For patients who have, or are expected to, eat little or nothing for 5 days; or
- 4 For patients who have a poor absorptive capacity and/or high nutrient losses and/or increased nutritional needs from causes such as catabolism; or
- 5 For use pre- and post-surgery; or
- 6 For patients being tube-fed; or
- 7 For tube-feeding as a transition from intravenous nutrition.

SPECIAL FOODS

| | (ex man. | Price . excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|---|----------|------------------------|------|----------|-------------------------------------|
| LOW-GI ENTERAL FEED 1 KCAL/ML - Restricted see terms on the p | revious | page | | | |
| t Liquid 5 g protein, 9.6 g carbohydrate and 5.4 g fat per 100 ml, 1,00 bottle | | 7.5 | 0 | 1,000 ml | Glucerna Select RTH (Vanilla) |
| Liquid 4.3 g protein, 11.3 g carbohydrate and 4.2 g fat per 100 ml, 1,000 ml bag | | | | | e.g. Nutrison Advanced Diason |
| LOW-GI ORAL FEED 1 KCAL/ML - Restricted see terms on the previo | ous page | е | | | |
| Liquid 4.5 g protein, 9.8 g carbohydrate, 4.4 g fat and 1.9 g fibre pe 100 ml, can | | 2.1 | 0 | 237 ml | Sustagen Diabetic (Vanilla) |
| t Liquid 5 g protein, 9.6 g carbohydrate and 5.4 g fat per 100 ml, 250 bottle. | | 18 | 8 | 250 ml | Glucerna Select (Vanilla) |
| Liquid 6 g protein, 9.5 g carbohydrate, 4.7 g fat and 2.6 g fibre per | | | 0 | 200 111 | |
| 100 ml, can | | 2.1 | 0 | 237 ml | Resource Diabetic (Vanilla) |
| Liquid 4.9 g protein, 11.7 g carbohydrate, 3.8 g fat and 2 g fibre per 100 ml, 200 ml bottle | | | | | e.g. Diasip |
| Elemental and Semi-Elemental Products | | | | | |
| → Restricted (RS1216) Initiation Any of the following: | | | | | |

- 1 Malabsorption; or
- 2 Short bowel syndrome; or
- 3 Enterocutaneous fistulas; or
- 4 Eosinophilic enteritis (including oesophagitis); or
- 5 Inflammatory bowel disease; or
- 6 Acute pancreatitis where standard feeds are not tolerated; or
- 7 Patients with multiple food allergies requiring enteral feeding.

| AMINO ACID ORAL FEED – Restricted see terms above t Powder 11 g protein, 62 g carbohydrate and 1 g fat per sachet4.50 80 g AMINO ACID ORAL FEED 0.8 KCAL/ML – Restricted see terms above | Vivonex TEN | |
|---|---------------------------------|----------|
| Liquid 2.5 g protein, 11 g carbohydrate and 3.5 g fat per 100 ml, 250 ml carton | e.g. Elemental 02 | 8 Extra |
| PEPTIDE-BASED ENTERAL FEED 1 KCAL/ML – Restricted see terms above t Liquid 4 g protein, 17.6 g carbohydrate and 1.7 g fat per 100 ml, | | |
| 1,000 ml bag | e.g. Nutrison Adv. Peptisorb | anced |
| t Liquid 4 g protein, 17.7 g carbohydrate and 1.7 g fat per 100 ml, | | |
| 1,000 ml bag | e.g. Nutrison Adva Peptisorb | anced |
| (e.g. Nutrison Advanced Peptisorb Liquid 4 g protein, 17.6 g carbohydrate and 1.7 g fat per 100 ml, 1 February 2021) | ,000 ml bag to be de | listed 1 |
| PEPTIDE-BASED ENTERAL FEED 1.5 KCAL/ML - Restricted see terms above t Liquid 6.75 g protein, 18.4 g carbohydrate and 5.5 g fat per 100 ml, bottle18.06 1,000 ml | Vital | |

e.g. Brand indicates brand example only. It is not a contracted product.

SPECIAL FOODS

| | Price (ex man. excl. GST | | Brand or Generic |
|--|-----------------------------|--------------------|---|
| | \$ | Per | Manufacturer |
| PEPTIDE-BASED ORAL FEED - Restricted see terms on the previ Powder 13.7 g protein, 62.9 g carbohydrate and 17.5 g fat per 10 400 g can Powder 13.8 g protein, 59 g carbohydrate and 18 g fat per 100 g |)0 g, | | e.g. Peptamen Junior |
| can | , | | e.g. MCT Pepdite; MCT Pepdite 1+ |
| PEPTIDE-BASED ORAL FEED 1 KCAL/ML – Restricted see terms Liquid 5 g protein, 16 g carbohydrate and 1.69 g fat per 100 ml, o | | 237 ml | Peptamen OS 1.0 (Vanilla) |
| Fat Modified Products | | | |
| FAT-MODIFIED FEED - Restricted see terms below Powder 12.9 g protein, 69.1 g carbohydrate and 12.9 g fat per 10 400 g can → Restricted (RS1470) Initiation Any of the following: Patient has metabolic disorders of fat metabolism; or | 10 g, | | e.g. Monogen |
| Patient has a chyle leak; or Modified as a modular feed, made from at least one nutrient n the Pharmaceutical Schedule, for adults. Note: Patients are required to meet any Special Authority criteria associated and the second seco | | | |
| Hepatic Products | | | |
| → Restricted (RS1217) Initiation For children (up to 18 years) who require a liver transplant. HEPATIC ORAL FEED - Restricted see terms above t Powder 11 g protein, 64 g carbohydrate and 20 g fat per 100 g, c | an78.97 | 400 g | Heparon Junior |
| High Calorie Products | | | |
| → Restricted (RS1317) Initiation Any of the following: Patient is fluid volume or rate restricted; or Patient requires low electrolyte; or Both: | pottle5.50 e per | 500 ml 1,000 ml | Nutrison Concentrated TwoCal HN RTH (Vanilla) |

Products with Hospital Supply Status (HSS) are in **bold** Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| | | rice excl. GST) \$ | Per | Brand or Generic Manufacturer |
|--|-----------------------------------|--------------------------|----------------------------------|--|
| ORAL FEED 2 KCAL/ML – Restricted see terms on the previous page Liquid 8.4 g protein, 22.4 g carbohydrate, 8.9 g fat and 0.8 g fibre 100 ml, bottle. | per | 1.90 | 200 ml | Two Cal HN |
| High Protein Products | | | | |
| HIGH PROTEIN ENTERAL FEED 1.25 KCAL/ML – Restricted see to ↓ Liquid 6.3 g protein, 14.2 g carbohydrate and 4.9 g fat per 100 ml 1,000 ml bottle → Restricted (RS1327) | | | | e.g. Nutrison Protein Plus |
| Initiation Both: The patient has a high protein requirement; and Any of the following: Patient has liver disease; or Patient is obese (BMI > 30) and is undergoing surgery; Patient is fluid restricted; or Patient's needs cannot be more appropriately met using HIGH PROTEIN ENTERAL FEED 1.28 KCAL/ML - Restricted see to Liquid 6.3 g protein, 14.1 g carbohydrate, 4.9 g fat and 1.5 g fibre 100 ml, 1,000 ml bag Restricted (RS1327) initiation Both: The patient has a high protein requirement; and Any of the following: Patient has liver disease; or Patient has liver disease; or Patient has liver disease; or Patient is obese (BMI > 30) and is undergoing surgery; Patient is fluid restricted; or Patient's needs cannot be more appropriately met using | g high calor erms below per | , | | e.g. Nutrison Protein Plus Multi Fibre |
| Infant Formulas | | | | |
| AMINO ACID FORMULA - Restricted see terms on the next page Powder 1.95 g protein, 8.1 g carbohydrate and 3.5 g fat per 100 r 400 g can Powder 13 g protein, 49 g carbohydrate and 23 g fat per 100 g, 4 can Powder 13.3 g protein, 56 g carbohydrate and 22 g fat per 100 g, can Powder 13.5 g protein, 52 g carbohydrate and 24.5 g fat per 100 | 00 g 400 g | 53.00 | 400 g | e.g. Neocate e.g. Neocate SYNEO unflavoured e.g. Neocate Junior Unflavoured Neocate Gold |
| Powder 14.8 g protein, 51.4 g carbohydrate and 23 g fat per 100 Powder 15 g protein, 56 g carbohydrate and 20 g fat per 100 g, cz Powder 2.2 g protein, 7.8 g carbohydrate and 3.4 g fat per 100 m Powder 2.2 g protein, 7.8 g carbohydrate and 3.4 g fat per 100 m | g, can an I, can | 53.00 43.60 53.00 | 400 g 400 g 400 g 400 g | (Unflavoured) Neocate Junior Vanilla Alfamino Junior Elecare LCP (Unflavoured) Elecare (Unflavoured) Elecare (Vanilla) |

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| | | Price | | | Brand or |
|---|----------|-------------|----------|-----------|-------------------------------|
| (| ex man. | excl. \$ | GST) | Per | Generic Manufacturer |
| ► Restricted (RS1765) | | • | | | |
| itiation | | | | | |
| ny of the following: | | | | | |
| Extensively hydrolysed formula has been reasonably trialled for 2 intolerance or allergy or malabsorption; or | -4 weeł | ks and | d is ina | opropriat | te due to documented sever |
| 2 History of anaphylaxis to cows' milk protein formula or dairy produ | icts; or | | | | |
| 3 Eosinophilic oesophagitis; or | | | | | |
| 4 Ultra-short gut; or | | | | | |
| 5 Severe Immune deficiency. | | | | | |
| ontinuation I of the following: | | | | | |
| 5 | | | tain a | | tanaivaly by draly and infant |
| An assessment as to whether the infant can be transitioned to a c formula has been undertaken; and | ows m | lik pro | nein, so | by, or ex | tensively hydrolysed imant |
| 2 The outcome of the assessment is that the infant continues to req | uire an | amin | o acid i | nfant for | mula: and |
| 3 Amino acid formula is required for a nutritional deficit. | | | | | |
| XTENSIVELY HYDROLYSED FORMULA - Restricted see terms belo | w | | | | |
| Powder 1.6 g protein, 7.5 g carbohydrate and 3.1 g fat per 100 ml, 9 | | | | | |
| can | - | .30.4 | 2 | 900 q | Allerpro 1 |
| Powder 1.6 g protein, 7.8 g carbohydrate and 3.2 g fat per 100 ml, 9 | | | | 5 | - F |
| can | | . 30.4 | 2 | 900 g | Allerpro 2 |
| Powder 14 g protein, 53.4 g carbohydrate and 27.3 g fat per 100 g, | | | | | |
| 450 g can | | | | | e.g. Aptamil Gold+ Pep |
| Restricted (RS1502) | | | | | Junior |
| itiation | | | | | |
| ny of the following: | | | | | |
| 1 Both: | | | | | |

- 1 Both:
 - 1.1 Cows' milk formula is inappropriate due to severe intolerance or allergy to its protein content; and
 - 1.2 Either:
 - 1.2.1 Soy milk formula has been reasonably trialled without resolution of symptoms; or
 - 1.2.2 Soy milk formula is considered clinically inappropriate or contraindicated; or
- 2 Severe malabsorption; or
- 3 Short bowel syndrome; or
- 4 Intractable diarrhoea; or
- 5 Biliary atresia; or
- 6 Cholestatic liver diseases causing malsorption; or
- 7 Cystic fibrosis; or
- 8 Proven fat malabsorption; or
- 9 Severe intestinal motility disorders causing significant malabsorption; or
- 10 Intestinal failure; or
- 11 For step down from Amino Acid Formula.

Note: A reasonable trial is defined as a 2-4 week trial, or signs of an immediate IgE mediated allergic reaction.

Continuation

Both:

- 1 An assessment as to whether the infant can be transitioned to a cows' milk protein or soy infant formula has been undertaken; and
- 2 The outcome of the assessment is that the infant continues to require an extensively hydrolysed infant formula.

FRUCTOSE-BASED FORMULA

Powder 14.6 g protein, 49.7 g carbohydrate and 30.8 g fat per 100 g, 400 g can

e.g. Galactomin 19

| Price (ex man. excl. GS \$ | Г) Per | Brand or Generic Manufacturer |
|---|-----------|--|
| LACTOSE-FREE FORMULA | | |
| Powder 1.3 g protein, 7.3 g carbohydrate and 3.5 g fat per 100 ml, 900 g can | | e.g. Karicare Aptamil Gold De-Lact |
| Powder 1.5 g protein, 7.2 g carbohydrate and 3.6 g fat per 100 ml, 900 g can | | e.g. S26 Lactose Free |
| LOW-CALCIUM FORMULA | | |
| Powder 14.6 g protein, 53.7 g carbohydrate and 26.1 g fat per 100 g, 400 g can | | e.g. Locasol |
| PAEDIATRIC ORAL/ENTERAL FEED 1 KCAL/ML - Restricted see terms below | | |
| ↓ Liquid 2.6 g protein, 10.3 g carbohydrate, 5.4 g fat and 0.6 g fibre per 100 ml, bottle | 125 ml | Infatrini |
| Initiation – Fluid restricted or volume intolerance with faltering growth Both: | | |
| 1 Either: | | |
| 1.1 The patient is fluid restricted or volume intolerant; or1.2 The patient has increased nutritional requirements due to faltering growth; a | and | |
| 2 Patient is under 18 months old and weighs less than 8kg. | | |
| Note: 'Volume intolerant' patients are those who are unable to tolerate an adequate volur growth rate. These patients should have first trialled appropriate clinical alternative treatrr and adjusting the frequency of feeding. | | |
| PRETERM FORMULA – Restricted see terms below Liquid 2.2 g protein, 8.4 g carbohydrate and 4.4 g fat per 100 ml, bottle0.75 Liquid 2.3 g protein, 8.6 g carbohydrate and 4.2 g fat per 100 ml, 90 ml | 100 ml | S26 LBW Gold RTF |
| Liquid 2.5 g protein, 8.4 g carbohydrate and 3.9 g fat per 100 ml, 70 ml | | e.g. Pre Nan Gold RTF |
| bottle | | e.g. Karicare Aptamil Gold+Preterm |
| → Restricted (RS1224) Initiation | | |
| For infants born before 33 weeks' gestation or weighing less than 1.5 kg at birth. THICKENED FORMULA | | |
| Powder 1.8 g protein, 8.1 g carbohydrate and 3.3 g fat per 100 ml, 900 g can | | e.g. Karicare Aptamil Thickened AR |
| Ketogenic Diet Products | | |
| HIGH FAT FORMULA – Restricted see terms below Powder 14.4 g protein, 2.9 g carbohydrate and 69.2 g fat per 100 g, can35.50 | 300 g | Ketocal |
| | 000 g | 4:1 (Unflavoured) Ketocal 4:1 (Vanilla) |
| • Powder 15.3 g protein, 7.2 g carbohydrate and 67.7 g fat per 100 g, can 35.50 | 300 g | Ketocal 3:1 (Unflavoured) |
| → Restricted (RS1225) | | |

Initiation

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For patients with intractable epilepsy, pyruvate dehydrogenase deficiency or glucose transported type-1 deficiency and other conditions requiring a ketogenic diet.

| Price (ex man. excl. GST) | | Brand or Generic |
|----------------------------------|-----|---------------------|
| \$ | Per | Manufacturer |

Paediatric Products

| → Restricted (RS1473) Initiation Both: | | |
|--|---------------|--|
| 1 Child is aged one to ten years; and | | |
| 2 Any of the following: 2.1 The child is being fed via a tube or a tube is to be inserted for the purposes of 2.2 Any condition causing malabsorption; or 2.3 Faltering growth in an infant/child; or 2.4 Increased nutritional requirements; or 2.5 The child is being transitioned from TPN or tube feeding to oral feeding; or 2.6 The child has eaten, or is expected to eat, little or nothing for 3 days. | of feeding; c |)r |
| PAEDIATRIC ENTERAL FEED 0.76 KCAL/ML - Restricted see terms above | | |
| Liquid 2.5 g protein, 12.5 g carbohydrate, 3.3 g fat and 0.7 g fibre per 100 ml, bag4.00 | 500 ml | Nutrini Low Energy Multifibre RTH |
| PAEDIATRIC ENTERAL FEED 1 KCAL/ML – Restricted see terms above Liquid 2.8 g protein, 11.2 g carbohydrate and 5 g fat per 100 ml, bag2.68 Liquid 2.8 g protein, 12.3 g carbohydrate and 4.4 g fat per 100 ml, | 500 ml | Pediasure RTH |
| 500 ml bag | | e.g. Nutrini RTH |
| PAEDIATRIC ENTERAL FEED 1.5 KCAL/ML - Restricted see terms above | | |
| Liquid 4.1 g protein, 18.5 g carbohydrate, 6.7 g fat and 0.8 g fibre per 100 ml, bag6.00 | 500 ml | Nutrini Energy Multi Fibre |
| Liquid 4.1 g protein, 18.5 g carbohydrate and 6.7 g fat per 100 ml, 500 ml bag | | e.g. Nutrini Energy RTH |
| PAEDIATRIC ORAL FEED 1 KCAL/ML – Restricted see terms above Liquid 4.2 g protein, 16.7 g carbohydrate and 7.5 g fat per 100 ml, bottle1.07 | 200 ml | Pediasure (Chocolate) Pediasure (Strawberry) Pediasure (Vanilla) |
| Liquid 4.2 g protein, 16.7 g carbohydrate and 7.5 g fat per 100 ml, can | 250 ml | Pediasure (Vanilla) |
| 200 ml bottle t Liquid 4.0 g protein, 18.8 g carbohydrate, 6.8 g fat and 1.5 g fibre per | | e.g. Fortini |
| 100 ml, 200 ml bottle | | e.g. Fortini Multifibre |
| Renal Products | | |
| LOW ELECTROLYTE ENTERAL FEED 1.8 KCAL/ML – Restricted see terms below | | |
| ↓ Liquid 8.1 g protein, 14.74 g carbohydrate, 9.77 g fat and 1.26 g fibre per 100 ml, bottle | 500 ml | Nepro HP RTH |
| For patients with acute or chronic kidney disease. | | |
| LOW ELECTROLYTE ORAL FEED − Restricted see terms on the next page Powder 7.5 g protein, 59 g carbohydrate and 26.3 g fat per 100 g, 400 g can | | e.g. Kindergen |
| | | |

| | | Price | | | Brand or | |
|--|-----------------------------|---------------|--------|----------------------|---|--|
| | (ex man. | excl. \$ | GST) | Per | Generic Manufacturer | |
| → Restricted (RS1227) nitiation For children (up to 18 years) with acute or chronic kidney disease. OW ELECTROLYTE ORAL FEED 1.8 KCAL/ML ↓ Liquid 8 g protein, 14.74 g carbohydrate, 9.77 g fat and 1.26 g fib | • | | | | | |
| 100 ml, carton → Restricted (RS1228) | | 2.67 | 7 | 220 ml | Nepro HP (Strawberry) Nepro HP (Vanilla) | |
| Initiation For patients with acute or chronic kidney disease. | | | | | | |
| LOW ELECTROLYTE ORAL FEED 2 KCAL/ML – Restricted see ter Liquid 9.1 g protein, 19 g carbohydrate and 10 g fat per 100 ml, c | arton | 3.31 | l | 237 ml | Novasource Renal (Vanilla) | |
| ↓ Liquid 3 g protein, 25.5 g carbohydrate and 9.6 g fat per 100 ml, 2 bottle ↓ Liquid 7.5 g protein, 20 g carbohydrate and 10 g fat per 100 ml, 1 carton → Restricted (RS1228) Initiation For patients with acute or chronic kidney disease. | | | | | e.g. Renilon 7.5 | |
| Respiratory Products | | | | | | |
| LOW CARBOHYDRATE ORAL FEED 1.5 KCAL/ML - Restricted se ↓ Liquid 6.2 g protein, 10.5 g carbohydrate and 9.32 g fat per 100 n (<i>Pulmocare (Vanilla) Liquid 6.2 g protein, 10.5 g carbohydrate and 9.3</i> → Restricted (RS1230) Initiation For patients with CORD and hypercapnia, defined as a CO2 value exercise | nl, bottle 32 g fat per | 1.66 100 i | nl, bo | 237 ml ttle to be | Pulmocare (Vanilla) delisted 1 October 2020) | |
| Surgical Products | | | | | | |
| HIGH ARGININE ORAL FEED 1.4 KCAL/ML − Restricted see terms Liquid 10.1 g protein, 15 g carbonhydrate, 4.5 g fat and 0 g fibre p 100 ml, carton | ber | 4.00 |) | 178 ml | Impact Advanced Recovery | |
| → Restricted (RS1231) | | | | | Hoovery | |
| Initiation Three packs per day for 5 to 7 days prior to major gastrointestinal, heap PREOPERATIVE CARBOHYDRATE FEED 0.5 KCAL/ML - Restrict ↓ Oral liq 0 g protein, 12.6 g carbohydrate and 0 g fat per 100 ml, 2 bottle | ed see terr 00 ml | ns be | low | 4 | preOp | |
| Initiation Maximum of 400 ml as part of an Enhanced Recovery After Surgery (| ERAS) pro | tocol | 2 to 3 | hours be | fore major abdominal | |

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Price (ex man. excl. GST)

\$

Per

Brand or Generic Manufacturer

Standard Feeds

➡ Restricted (RS1214)

Initiation

Any of the following:

- For patients with malnutrition, defined as any of the following:
- 1 Any of the following:
 - 1.1 BMI < 18.5; or
 - 1.2 Greater than 10% weight loss in the last 3-6 months; or
 - 1.3 BMI < 20 with greater than 5% weight loss in the last 3-6 months; or
- 2 For patients who have, or are expected to, eat little or nothing for 5 days; or
- 3 For patients who have a poor absorptive capacity and/or high nutrient losses and/or increased nutritional needs from causes such as catabolism; or
- 4 For use pre- and post-surgery; or
- 5 For patients being tube-fed; or
- 6 For tube-feeding as a transition from intravenous nutrition; or
- 7 For any other condition that meets the community Special Authority criteria.

ENTERAL FEED 1.5 KCAL/ML - Restricted see terms above

| L t | Liquid 6 g protein, 18.3 g carbohydrate and 5.8 g fat per 100 ml, bag7.00 | 1,000 ml | Nutrison Energy |
|--------|--|----------|---|
| t | Liquid 6 g protein, 18.4 g carbohydrate, 5.8 g fat and 1.5 g fibre per | | o a Nutricon Energy |
| | 100 ml, 1,000 ml bag | | e.g. Nutrison Energy Multi Fibre |
| t | Liquid 6.25 g protein, 20 g carbohydrate and 5 g fat per 100 ml, can | 250 ml | Ensure Plus HN |
| t | Liquid 6.27 g protein, 20.4 g carbohydrate and 4.9 g fat per 100 ml, bag7.00 | 1,000 ml | Ensure Plus HN RTH |
| t | Liquid 6.38 g protein, 21.1 g carbohydrate, 4.9 g fat and 1.2 g fibre per | , | |
| | 100 ml, bag | 1,000 ml | Jevity HiCal RTH |
| ΕN | ITERAL FEED 1 KCAL/ML - Restricted see terms above | | |
| t | Liquid 4 g protein, 13.6 g carbohydrate and 3.4 g fat per 100 ml, bottle | 1,000 ml | Osmolite RTH |
| t | Liquid 4 g protein, 14.1 g carbohydrate, 3.47 g fat and 1.76 g fibre per | | |
| | 100 ml, bottle | 1,000 ml | Jevity RTH |
| t | Liquid 4 g protein, 12.3 g carbohydrate and 3.9 g fat per 100 ml, | | |
| | 1,000 ml bag | | e.g. NutrisonStdRTH; |
| | | | NutrisonLowSodium |
| t | Liquid 4 g protein, 12.3 g carbohydrate and 3.9 g fat per 100 ml, | | |
| • | 1.000 ml bottle | | e.g. Nutrison Low |
| | | | Sodium |
| t | Liquid 4 g protein, 12.3 g carbohydrate, 3.9 g fat and 1.5 g fibre per | | e e a a a a a a a a a a a a a a a a a a |
| | 100 ml, 1000 ml bag | | e.g. Nutrison Multi Fibre |
| ΕN | ITERAL FEED 1.2 KCAL/ML – Restricted see terms above | | |
| t | Liquid 5.55 g protein, 15.1 g carbohydrate, 3.93 g fat and 2 g fibre per | | |
| | 100 ml, 1,000 ml bag | | e.g. Jevity Plus RTH |
| ΕN | ITERAL FEED WITH FIBRE 0.83 KCAL/ML - Restricted see terms above | | |
| t | Liquid 5.5 g protein, 8.8 g carbohydrate, 2.5 g fat and 1.5 g fibre per | | |
| | 100 ml, bottle | 1,000 ml | Nutrison 800 Complete Multi Fibre |

| Price (ex man. excl. GS | :Т) | Brand or Generic |
|---|--------|--|
| \$ | Per | Manufacturer |
| ORAL FEED – Restricted see terms on the previous page | | |
| t Powder 15.9 g protein, 57.4 g carbohydrate and 14 g fat per 100 g, can26.00 | 850 g | Ensure (Chocolate) Ensure (Vanilla) |
| t Powder 20.8 g protein, 61 g carbohydrate and 9.4 g fat per 100 g, can | 857 g | Fortisip (Vanilla) |
| t Powder 23 g protein, 65 g carbohydrate and 2.5 g fat per 100 g, can | 840 g | Sustagen Hospital Formula Active (Choc) Sustagen Hospital Formula Active (Van) |
| Note: Community subsidy of Sustagen Hospital Formula is subject to both Spec manufacturer's surcharge. Higher subsidy by endorsement is available for patie criteria; fat malabsorption, fat intolerance or chyle leak. | | criteria and a |
| ORAL FEED 1 KCAL/ML - Restricted see terms on the previous page | | |
| Liquid 3.8 g protein, 23 g carbohydrate and 12.7 g fibre per 100 ml, | | |
| 237 ml carton | | e.g. Resource Fruit Beverage |
| ORAL FEED 1.5 KCAL/ML - Restricted see terms on the previous page | | |
| Liquid 5.5 g protein, 21.1 g carbohydrate and 4.81 g fat per 100 ml, can | 237 ml | Ensure Plus (Vanilla) |
| carton1.26 | 200 ml | Ensure Plus (Banana) Ensure Plus (Chocolate) Ensure Plus (Fruit of the Forest) Ensure Plus (Vanilla) |
| Liquid 4 g protein and 33.5 g carbohydrate per 100 ml, 200 ml bottle Liquid 6 g protein, 18.4 g carbohydrate and 5.8 g fat per 100 ml, 200 ml | | e.g. Fortijuice |
| bottle | | e.g. Fortisip |
| Liquid 6 g protein, 18.4 g carbohydrate, 5.8 g fat and 2.3 g fibre per 100 ml, 200 ml bottle | | e.g. Fortisip Multi Fibre |

VACCINES

| | F (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|--|-----------------------------|-------------------------------|-----------------------|-------------------------------------|
| Bacterial and Viral Vaccines | | | | | |
| DIPHTHERIA, TETANUS, PERTUSSIS AND POLIO VACCINE – Res | stricted se | e terr | ns <mark>belo</mark> | W | |
| Inj 30 IU diphtheria toxoid with 30IU tetanus toxoid, 25 mcg pertus toxoid, 25 mcg pertussis filamentous haemagglutinin, 8 mcg pertactin and 80 D-antigen units poliomyelitis virus in 0.5 ml s - 0% DV Oct-20 to 2024. | sis yringe | | | 10 | Infanrix IPV |
| → Restricted (RS1387) Initiation | | | | | |
| Any of the following: 1 A single dose for children up to the age of 7 who have complete 2 A course of up to four vaccines is funded for catch up programmer primary immunisation; or 3 An additional four doses (as appropriate) are funded for (re-)im | mes for ch | ildren n for p | (to the atients | age of 1 post HS | CT, or chemotherapy; pre- |
| or post splenectomy; pre- or post solid organ transplant, renal or or | - | | er seve | rely imm | unosuppressive regimens; |
| 4 Five doses will be funded for children requiring solid organ tran | • | | | | |
| Note: Please refer to the Immunisation Handbook for appropriate sche DIPHTHERIA, TETANUS, PERTUSSIS, POLIO, HEPATITIS B AND H | | | | | |
| Restricted see terms below Inj 30 IU diphtheria toxoid with 40 IU tetanus toxoid, 25 mcg pertus toxoid, 25 mcg pertussis filamentous haemagglutinin, 8 mcg pertactin, 80 D-antigen units poliomyelitis virus, 10 mcg hepat – 0% DV Oct-20 to 2024 | ssis itis B | | | 10 | Infanrix-hexa |
| Initiation | | | | | |
| Any of the following: 1 Up to four doses for children up to and under the age of 10 for 2 An additional four doses (as appropriate) are funded for (re-)im are patients post haematopoietic stem cell transplantation, or c organ transplant, renal dialysis and other severely immunosupp 3 Up to five doses for children up to and under the age of 10 rece | munisatior hemothera pressive re | n for c apy; p egimei | hildren re or po ns; or | up to ar ost spler | ectomy; pre- or post solid |
| Note: A course of up-to four vaccines is funded for catch up programm complete full primary immunisation. Please refer to the Immunisation programmes. | nes for chi | ildren | (up to a | and unde | er the age of 10 years) to |
| Bacterial Vaccines | | | | | |
| ADULT DIPHTHERIA AND TETANUS VACCINE Inj 2 IU diphtheria toxoid with 20 IU tetanus toxoid in 0.5 ml syring | e | 0.00 |) | 5 | ADT Booster |
| → Restricted (RS1386) Initiation Any of the following: For vaccination of patients aged 45 and 65 years old; or For vaccination of previously unimmunised or partially immunis | ed patients | s; or | | | |
| 3 For revaccination following immunosuppression; or 4 For boosting of patients with tetanus-prone wounds; or 5 For use in testing for primary immunodeficiency diseases, on the paediatrician. | ne recomm | nenda | tion of a | an interr | al medicine physician or |

Note: Please refer to the Immunisation Handbook for the appropriate schedule for catch up programmes.

Products with Hospital Supply Status (HSS) are in **bold**

Expiry date of HSS period is 30 June of the year indicated unless otherwise stated.

| (ex n | nan. | Price excl. \$ | GST) | Per | | Brand or Generic Manufacturer |
|--|---------------|----------------------|---------------------|---------------------|--------------|-------------------------------------|
| (ADT Booster Inj 2 IU diphtheria toxoid with 20 IU tetanus toxoid in 0.5 ml sy | ringe | e to b | e delis | ted 1 C | Octob | per 2020) |
| BACILLUS CALMETTE-GUERIN VACCINE - Restricted see terms below Inj Mycobacterium bovis BCG (Bacillus Calmette-Guerin), Danish strain 1331, live attenuated, vial Danish strain 1331, live attenuated, vial with diluent - 0% DV Oct-20 to 2024 → Restricted (RS1233) Initiation | | 0.00 | 0 | 10 | | BCG Vaccine |
| All of the following: | | | | | | |
| For infants at increased risk of tuberculosis defined as: Living in a house or family with a person with current or past history of Having one or more household members or carers who within the las equal to 40 per 100,000 for 6 months or longer; and During their first 5 years will be living 3 months or longer in a country | t 5 y | ears a rat | lived in e of TE | 3 > or e | equal | to 40 per 100,000. |
| Note: A list of countries with high rates of TB are available at http://www.hea www.bcgatlas.org/index.php | uun.g | JONI'L | IZ/IUDe | rculosi | 5 (56 | Parch for Downloads) of |
| DIPHTHERIA, TETANUS AND PERTUSSIS VACCINE – Restricted see te Inj 2 IU diphtheria toxoid with 20 IU tetanus toxoid, 8 mcg pertussis toxoid, 8 mcg pertussis filamentous haemagglutinin and 2.5 mcg pertactin in 0.5 ml syringe – 0% DV Oct-20 to 2024 | | | | 1 10 | | Boostrix Boostrix |
| ➡ Restricted (RS1766) | | | | 10 | | DOOSITIX |
| Initiation | | | | | | |
| Any of the following: | | | | | | |
| A single dose for pregnant women in the second or third trimester of a A single dose for parents or primary caregivers of infants admitted to Baby Unit for more than 3 days, who had not been exposed to materi A course of up to four doses is funded for children from age 7 up the immunisation; or | a Ne nal v | eonat accir | al Inter ation a | nsive C at least | Care 14 d | lays prior to birth; or; or |
| 4 An additional four doses (as appropriate) are funded for (re-)immunis transplantation or chemotherapy; pre or post splenectomy; pre- or po severely immunosuppressive regimens; or 5 A single dose for vaccination of patients aged 65 years old; or | | | | | | |
| 6 A single dose for vaccination of patients aged 45 years old who have 7 For vaccination of previously unimmunised or partially immunised pat 8 For revaccination following immunosuppression; or 9 For boosting of patients with tetanus-prone wounds. | | | 1 previo | ous teta | anus | doses; or |
| Note: Tdap is not registered for patients aged less than 10 years. Please re schedule for catch up programmes. | fer t | to the | Immui | nisatio | n Ha | ndbook for the appropriate |
| HAEMOPHILUS INFLUENZAE TYPE B VACCINE – Restricted see terms | oelov | w | | | | |
| Haemophilus Influenzae type B polysaccharide 10 mcg conjugated to tetanus toxoid as carrier protein 20-40 mcg; prefilled syringe plus | | | | | | |
| vial 0.5 ml → Restricted (RS1520) Initiation | | 0.00 | 0 | 1 | | Hiberix |
| Therapy limited to 1 dose Any of the following: | | | | | | |
| For primary vaccination in children; or An additional dose (as appropriate) is funded for (re-)immunisation fo | r pat | tients | post h | aemat | opoie | etic stem cell |

e.g. Brand indicates brand example only. It is not a contracted product.

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VACCINES

| | | | VACCINES |
|--|--|-------------------------------|---|
| | Price (ex man. excl. G \$ | aST) Per | Brand or Generic Manufacturer |
| ontinued transplantation, or chemotherapy; functional asplenic; pre or p post cochlear implants, renal dialysis and other severely immu 3 For use in testing for primary immunodeficiency diseases, on paediatrician. | inosuppressive reg | gimens; or | |
| IENINGOCOCCAL (A, C, Y AND W-135) CONJUGATE VACCINE | - Restricted see t | erms below | |
| Inj 4 mcg or each meningococcal polysaccharide conjugated to a | | | |
| approximately 48 mcg of diphtheria toxoid carrier per 0.5 ml 0% DV Oct-20 to 2024 → Restricted (RS1719) nitiation Either: | | 1 | Menactra |
| 1 Any of the following: | | | |
| 1.1 Up to three doses and a booster every five years for pa complement deficiency (acquired or inherited), function or 1.2 One dose for close contacts of meningococcal cases; of 1.3 A maximum of two doses for bone marrow transplant p 1.4 A maximum of two doses for patients following immuno | nal or anatomic asp or patients; or | | |
| 2 Both: | | | |
| Person is aged between 13 and 25 years, inclusive; an 2.2 Either: | 10 | | |
| 2.2.1 One dose for individuals who are entering within boarding school hostels, tertiary education halls2.2.2 One dose for individuals who are currently living residence, military barracks, or prisons, from 1 | of residence, milit g in boarding schoo | ary barrack ol hostels, te | s, or prisons; or ertiary education halls of |
| lotes: children under seven years of age require two doses 8 weeks ind then five yearly. | | | |
| Immunosuppression due to steroid or other immunosuppressive ther IENINGOCOCCAL C CONJUGATE VACCINE - Restricted see te | | perioù or gr | ealer man 20 uays. |
| Inj 10 mcg in 0.5 ml syringe | | 1 | Neisvac-C |
| nitiation – Children under 9 months of age | | | |
| Any of the following: 1 Up to three doses for patients pre- and post splenectomy and inherited), functional or anatomic asplenia or pre or post solid 2 Two doses for close contacts of meningococcal cases; or 3 A maximum of two doses for bone marrow transplant patients; 4 A maximum of two doses for patients pre- and post-immunosu | organ transplant; o ; or | | nent deficiency (acquired or |
| Notes: children under nine months of age require two doses 8 weeks schedules with meningococcal ACWY vaccine. | s apart. Refer to th | | |
| Immunosuppression due to steroid or other immunosuppressive their NEUMOCOCCAL (PCV10) CONJUGATE VACCINE – Restricted a mcg of pneumococcal polysaccharide serotypes 1, 5, 6B, 7F, 9V | see terms below | period of gr | eater than 28 days. |
| 14 and 23F; 3 mcg of pneumococcal polysaccharide serotyp 18C and 19F in 0.5 ml prefilled syringe - 0% DV Oct-20 to → Restricted (RS1768) | | 10 | Synflorix |
| nitiation A primary course of three doses for previously unvaccinated individua Note: Please refer to the Immunisation Handbook for the appropriate | | | |

| (ex r | Price nan. excl. GST \$ |) Per | Brand or Generic Manufacturer |
|--|-------------------------------|----------|-------------------------------------|
| PNEUMOCOCCAL (PCV13) CONJUGATE VACCINE – Restricted see ten Inj 30.8 mcg of pneumococcal polysaccharide serotypes 1, 3, 4, 5, 6A, | ns below | | |
| 6B, 7F, 9V, 14, 18C, 19A, 19F and 23F in 0.5 ml syringe | 0.00 | 1 10 | Prevenar 13 Prevenar 13 |

➡ Restricted (RS1769)

Initiation – High risk children who have received PCV10

Therapy limited to 1 dose

Two doses are funded for high risk children (over the age of 12 months and under 18 years) who have previously received two doses of the primary course of PCV10.

Initiation - High risk children aged under 5 years

Therapy limited to 4 doses

Both:

- 1 Up to an additional four doses (as appropriate) are funded for children aged under 5 years for (re-)immunisation; and
- 2 Any of the following:
 - 2.1 On immunosuppressive therapy or radiation therapy, vaccinate when there is expected to be a sufficient immune response; or
 - 2.2 With primary immune deficiencies; or
 - 2.3 With HIV infection; or
 - 2.4 With renal failure, or nephrotic syndrome; or
 - 2.5 Who are immune-suppressed following organ transplantation (including haematopoietic stem cell transplant); or
 - 2.6 With cochlear implants or intracranial shunts; or
 - 2.7 With cerebrospinal fluid leaks; or
 - 2.8 Receiving corticosteroid therapy for more than two weeks, and who are on an equivalent daily dosage of prednisone of 2 mg/kg per day or greater, or children who weigh more than 10 kg on a total daily dosage of 20 mg or greater; or
 - 2.9 With chronic pulmonary disease (including asthma treated with high-dose corticosteroid therapy); or
 - 2.10 Pre term infants, born before 28 weeks gestation; or
 - 2.11 With cardiac disease, with cyanosis or failure; or
 - 2.12 With diabetes; or
 - 2.13 With Down syndrome; or
 - 2.14 Who are pre-or post-splenectomy, or with functional asplenia.

Initiation - High risk adults and children 5 years and over

Therapy limited to 4 doses

Up to an additional four doses (as appropriate) are funded for (re-)immunisation of patients 5 years and over with HIV, for patients pre or post haematopoietic stem cell transplantation, or chemotherapy; pre- or post splenectomy; functional asplenia, pre- or post-solid organ transplant, renal dialysis, complement deficiency (acquired or inherited), cochlear implants, or primary immunodeficiency.

Initiation – Testing for primary immunodeficiency diseases

For use in testing for primary immunodeficiency diseases, on the recommendation of an internal medicine physician or paediatrician.

Note: Please refer to the Immunisation Handbook for the appropriate schedule for catch up programmes

PNEUMOCOCCAL (PPV23) POLYSACCHARIDE VACCINE - Restricted see terms below

Inj 575 mcg in 0.5 ml prefilled syringe (25 mcg of each 23 pneumococcal serotype) - 0% DV Oct-20 to 2024......0.00
 Pneumovax 23

→ Restricted (RS1587)

Initiation – High risk patients

Therapy limited to 3 doses

For patients with HIV, for patients post haematopoietic stem cell transplant, or chemotherapy; pre- or post-splenectomy; or with functional asplenia, pre- or post-solid organ transplant, renal dialysis, complement deficiency (acquired or inherited), cochlear

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST |) | Generic |
| \$ | Per | Manufacturer |

continued...

implants, or primary immunodeficiency.

Initiation – High risk children

Therapy limited to 2 doses

Both:

- 1 Patient is a child under 18 years for (re-)immunisation; and
- 2 Any of the following:
 - 2.1 On immunosuppressive therapy or radiation therapy, vaccinate when there is expected to be a sufficient immune response; or
 - 2.2 With primary immune deficiencies; or
 - 2.3 With HIV infection; or
 - 2.4 With renal failure, or nephrotic syndrome; or
 - 2.5 Who are immune-suppressed following organ transplantation (including haematopoietic stem cell transplant); or
 - 2.6 With cochlear implants or intracranial shunts; or
 - 2.7 With cerebrospinal fluid leaks; or
 - 2.8 Receiving corticosteroid therapy for more than two weeks, and who are on an equivalent daily dosage of prednisone of 2 mg/kg per day or greater, or children who weigh more than 10 kg on a total daily dosage of 20 mg or greater; or
 - 2.9 With chronic pulmonary disease (including asthma treated with high-dose corticosteroid therapy); or
 - 2.10 Pre term infants, born before 28 weeks gestation; or
 - 2.11 With cardiac disease, with cyanosis or failure; or
 - 2.12 With diabetes; or
 - 2.13 With Down syndrome; or
 - 2.14 Who are pre-or post-splenectomy, or with functional asplenia.

Initiation - Testing for primary immunodeficiency diseases

For use in testing for primary immunodeficiency diseases, on the recommendation of an internal medicine physician or paediatrician.

SALMONELLA TYPHI VACCINE - Restricted see terms below

Inj 25 mcg in 0.5 ml syringe

➡ Restricted (RS1243)

Initiation

For use during typhoid fever outbreaks.

Viral Vaccines

| HEPATITIS A VACCINE - Restricted see terms below Inj 720 ELISA units in 0.5 ml syringe - 0% DV Oct-20 to 20240.00 Inj 1440 ELISA units in 1 ml syringe - 0% DV Oct-20 to 20240.00 → Restricted (RS1638) Initiation | 1 1 | Havrix Junior Havrix | |
|--|--------|-------------------------|--|
| Any of the following: 1 Two vaccinations for use in transplant patients; or | | | |
| 2 Two vaccinations for use in children with chronic liver disease; or3 One dose of vaccine for close contacts of known hepatitis A cases. | | | |
| HEPATITIS B RECOMBINANT VACCINE ↓ Inj 5 mcg in 0.5 ml vial0.00 → Restricted (RS1588) Initiation Any of the following: | 1 | HBvaxPRO | |

| | l (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|--|-------------------------|----------------------------------|-----|-------------------------------------|
| Portinued For household or sexual contacts of known acute hepatitis B For children born to mothers who are hepatitis B surface antig For children up to and under the age of 18 years inclusive wh and require additional vaccination or require a primary course For HIV positive patients; or For hepatitis C positive patients; or for patients following non-consensual sexual intercourse; or For solid organ transplant patients; or For post-haematopoietic stem cell transplant (HSCT) patients Following needle stick injury. | yen (HBsAg o are consi of vaccina |) pos dered | itive; or not to | | |
| Inj 10 mcg in 1 ml vial • Restricted (RS1588) itiation ny of the following: | | 0.0 | 0 | 1 | HBvaxPRO |
| For household or sexual contacts of known acute hepatitis B For children born to mothers who are hepatitis B surface antig For children up to and under the age of 18 years inclusive wh and require additional vaccination or require a primary course For HIV positive patients; or For hepatitis C positive patients; or for patients following non-consensual sexual intercourse; or For patients following immunosuppression; or For post-haematopoietic stem cell transplant (HSCT) patients Following needle stick injury. | yen (HBsAg o are consi of vaccina |) pos dered | itive; or not to | | |
| Inj 20 mcg per 1 ml prefilled syringe – 0% DV Oct-20 to 2024 • Restricted (RS1671) iitiation ny of the following: 1 For household or sexual contacts of known acute hepatitis B j 2 For children born to mothers who are hepatitis B surface antig 3 For children up to and under the age of 18 years inclusive wh and require additional vaccination or require a primary course 4 For HIV positive patients; or 5 For hepatitis C positive patients; or 6 for patients following non-consensual sexual intercourse; or 7 For patients following immunosuppression; or 8 For solid organ transplant patients; or 9 For post-haematopoietic stem cell transplant (HSCT) patients 10 Following needle stick injury; or 11 For dialysis patients; or 12 For liver or kidney transplant patients. | patients or I gen (HBsAg o are consi of vaccina | nepat) pos dered | itis B ca itive; or not to | | |
| Inj 40 mcg per 1 ml vial * Restricted (RS1413) ititation oth: 1 For dialysis patients; and 2 For liver or kidney transplant patient. | | 0.0 | 0 | 1 | HBvaxPRO |

| | F (ex man. | Price excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|---|-----------------|----------------------|------|------------------|-------------------------------------|
| (HBvaxPRO Inj 5 mcg in 0.5 ml vial to be delisted 1 October 2020) (HBvaxPRO Inj 10 mcg in 1 ml vial to be delisted 1 October 2020) (HBvaxPRO Inj 40 mcg per 1 ml vial to be delisted 1 October 2020) | | | | | |
| HUMAN PAPILLOMAVIRUS (6, 11, 16, 18, 31, 33, 45, 52 AND 58) \ Inj 270 mcg in 0.5 ml syringe – 0% DV Oct-20 to 2024 | | | | ricted see 10 | e terms below Gardasil 9 |
| Up to 3 doses for people aged 15 to 26 years inclusive; or 2 Both: People aged 9 to 26 years inclusive; and People aged 9 to 26 years inclusive; and Any of the following: Up to 3 doses for confirmed HIV infection; or Up to 3 doses for transplant (including stem ce Up to 4 doses for Post chemotherapy. | ll) patients; (| or | | | |
| Initiation – Recurrent Respiratory Papillomatosis All of the following: 1 Either: 1.1 Maximum of two doses for children aged 14 years and 1.2 Maximum of three doses for people aged 15 years and 2 The patient has recurrent respiratory papillomatosis; and 3 The patient has not previously had an HPV vaccine. | | | | | |
| NFLUENZA VACCINE Inj 30 mcg in 0.25 ml syringe (paediatric quadrivalent vaccine) | | 9.00 | 0 | 1 | Afluria Quad Junior |
| → Restricted (RS1675) Initiation – cardiovascular disease for patients aged 6 months to Any of the following: Ischaemic heart disease; or Congestive heart failure; or Rheumatic heart disease; or Congenital heart disease; or Cerebro-vascular disease. | o 35 month | S | | | (2020 Formulation) |
| Note: hypertension and/or dyslipidaemia without evidence of end-or Initiation – chronic respiratory disease for patients aged 6 mont Either: 1 Asthma, if on a regular preventative therapy; or | | | | from fund | ding. |
| 2 Other chronic respiratory disease with impaired lung function. Note: asthma not requiring regular preventative therapy is excluded initiation – Other conditions for patients aged 6 months to 35 me Any of the following: | from fundin | g. | | | |
| Diabetes; or Chronic renal disease; or Any cancer, excluding basal and squamous skin cancers if no | ot invasive; o | or | | | |
| | | | | | continued. |

VACCINES

| Price | | Brand or |
|--------------------|-----|--------------|
| (ex man. excl. GST |) | Generic |
| \$ | Per | Manufacturer |

| cor | ntir | าน | ed | | |
|-----|------|----|----|--|--|
| | | | | | |

- 4 Autoimmune disease; or
- 5 Immune suppression or immune deficiency; or
- 6 HIV; or
- 7 Transplant recipient; or
- 8 Neuromuscular and CNS diseases/ disorders; or
- 9 Haemoglobinopathies; or
- 10 Is a child on long term aspirin; or
- 11 Has a cochlear implant; or
- 12 Errors of metabolism at risk of major metabolic decompensation; or
- 13 Pre and post splenectomy; or
- 14 Down syndrome; or
- 15 Child who has been hospitalised for respiratory illness or has a history of significant respiratory illness.

| t | Inj 60 mcg in 0.5 ml syringe (quadrivalent vaccine) | | 10 | Afluria Quad |
|---|---|------|----|--|
| | | 9.00 | 1 | (2020 Formualtion) Influvac Tetra (2020 formulation) |

→ Restricted (RS1674)

Initiation - People over 65

The patient is 65 years of age or over.

Initiation - cardiovascular disease for patients 3 years and over

Any of the following:

- 1 Ischaemic heart disease; or
- 2 Congestive heart failure; or
- 3 Rheumatic heart disease; or
- 4 Congenital heart disease; or
- 5 Cerebro-vascular disease.

Note: hypertension and/or dyslipidaemia without evidence of end-organ disease is excluded from funding. Initiation – chronic respiratory disease for patients 3 years and over

Either:

- 1 Asthma, if on a regular preventative therapy; or
- 2 Other chronic respiratory disease with impaired lung function.

Note: asthma not requiring regular preventative therapy is excluded from funding.

Initiation - Other conditions for patients 3 years and over

Either:

- 1 Any of the following:
 - 1.1 Diabetes; or
 - 1.2 chronic renal disease; or
 - 1.3 Any cancer, excluding basal and squamous skin cancers if not invasive; or
 - 1.4 Autoimmune disease; or
 - 1.5 Immune suppression or immune deficiency; or
 - 1.6 HIV; or
 - 1.7 Transplant recipient; or
 - 1.8 Neuromuscular and CNS diseases/ disorders; or
 - 1.9 Haemoglobinopathies; or
 - 1.10 Is a child on long term aspirin; or
 - 1.11 Has a cochlear implant; or
 - 1.12 Errors of metabolism at risk of major metabolic decompensation; or

VACCINES

| | (ex man. | Price . excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|---|---------------|------------------------|----------|------------|-------------------------------------|
| ontinued | | | | | |
| 1.13 Pre and post splenectomy; or | | | | | |
| 1.14 Down syndrome; or | | | | | |
| 1.15 Is pregnant; or | | | | | |
| 1.16 Is a child aged four and under who has been hospital | ised for resp | pirator | y illnes | s or has a | a history of significant |
| respiratory illness; or | | | | | |
| 2 Patients in a long-stay inpatient mental health care unit or wh | no are comp | ulsori | ly detai | ned long- | term in a forensic unit withi |
| a DHB hospital. | | | | | |
| IEASLES, MUMPS AND RUBELLA VACCINE - Restricted see to | erms below | | | | |
| Injection, measles virus 1,000 CCID50, mumps virus 5,012 CCI | D50, | | | | |
| Rubella virus 1,000 CCID50; prefilled syringe/ampoule of d | iluent | | | | |
| 0.5 ml - 0% DV Oct-20 to 2024 | | 0.0 | 0 | 10 | Priorix |
| Restricted (RS1487) nitiation – first dose prior to 12 months | | | | | |
| Therapy limited to 3 doses | | | | | |
| any of the following: | | | | | |
| 1 For primary vaccination in children; or | | | | | |
| 2 For revaccination following immunosuppression; or | | | | | |
| 3 For any individual susceptible to measles, mumps or rubella. | | | | | |
| nitiation – first dose after 12 months | | | | | |
| Therapy limited to 2 doses | | | | | |
| ny of the following: | | | | | |
| 1 For primary vaccination in children; or | | | | | |
| 2 For revaccination following immunosuppression; or | | | | | |
| 3 For any individual susceptible to measles, mumps or rubella. | | | | | |
| lote: Please refer to the Immunisation Handbook for appropriate se | chedule for o | catch | up prog | grammes. | |
| OLIOMYELITIS VACCINE – Restricted see terms below | | | | | |
| Inj 80 D-antigen units in 0.5 ml syringe – 0% DV Oct-20 to 202 | 4 | 0.0 | 0 | 1 | IPOL |
| → Restricted (RS1398) | | | | | |
| nitiation | | | | | |
| Fherapy limited to 3 doses | | | | | |
| ither: | | | | | |
| 1 For partially vaccinated or previously unvaccinated individual | is; or | | | | |
| 2 For revaccination following immunosuppression. lote: Please refer to the Immunisation Handbook for the appropria | to cohodulo | for or | toh un | programm | 200 |
| | | | lich up | programm | nes. |
| ABIES VACCINE Inj 2.5 IU vial with diluent | | | | | |
| , | | | | | |
| ROTAVIRUS ORAL VACCINE – Restricted see terms below | | | | | |
| Oral susp live attenuated human rotavirus 1,000,000 CCID50 p | | • • | ~ | 40 | Detecto |
| prefilled oral applicator – 0% DV Oct-20 to 2024 → Restricted (RS1590) | | 0.0 | U | 10 | Rotarix |
| nitiation | | | | | |
| Therapy limited to 2 doses | | | | | |
| Both: | | | | | |
| | | | | | |
| 1 First dose to be administered in infants aged under 14 weeks | s of age; and | b | | | |

| Price (ex man. excl. \$ | GST) | Per | Brand or Generic Manufacturer |
|--|-----------|-----------|-------------------------------------|
| VARICELLA VACCINE [CHICKENPOX VACCINE] – Restricted see terms below Inj 1350 PFU prefiiled syringe – 0% DV Oct-20 to 20240.0 | 0 | 1 10 | Varivax Varivax |
| Inj 2000 PFU prefilled syringe plus vial0.0 (Varilrix Inj 2000 PFU prefilled syringe plus vial to be delisted 1 October 2020) ⇒ Restricted (RS1591) Initiation - primary vaccinations Therapy limited to 1 dose Either: | 0 | 1 | Varilrix |
| Any infant born on or after 1 April 2016; or For previously unvaccinated children turning 11 years old on or after 1 July 20 infection (chickenpox). | 17, who | have no | ot previously had a varicella |
| Initiation – other conditions Therapy limited to 2 doses Any of the following: | | | |
| 1 Any of the following: | | | |
| for non-immune patients: | | | |
| 1.1 With chronic liver disease who may in future be candidates for transpla1.2 With deteriorating renal function before transplantation; or | intation; | or | |
| 1.3 Prior to solid organ transplant; or | | | |
| 1.4 Prior to any elective immunosuppression*; or | | | |
| 1.5 For post exposure prophylaxis who are immune competent inpatients; | | | |
| 2 For patients at least 2 years after bone marrow transplantation, on advice of th | | , | |
| 3 For patients at least 6 months after completion of chemotherapy, on advice of 4 For HIV positive patients non immune to varicella with mild or moderate immu 5 For patients with inborn errors of metabolism at risk of major metabolic decomvaricella; or | nosupp | ression o | on advice of HIV specialist; o |
| For household contacts of paediatric patients who are immunocompromised, immune compromise where the household contact has no clinical history of varice For household contacts of adult patients who have no clinical history of varice | aricella; | or | |
| immunocompromised or undergoing a procedure leading to immune comprom clinical history of varicella. | nise whe | ere the h | ousehold contact has no |
| Note: * immunosuppression due to steroid or other immunosuppressive therapy mus 28 days | | a treatm | ent period of greater than |
| VARICELLA ZOSTER VACCINE [SHINGLES VACCINE] – Restricted see terms be Varicella zoster virus (Oka strain) live attenuated vaccine [shingles | IOW | | |
| Various 2000 mile (one one in the construction vacous [one in the construction one one one one one one one one one o | 0 | 1 10 | Zostavax Zostavax |
| Initiation – people aged 65 years | | | |
| Therapy limited to 1 dose | | | |
| One dose for all people aged 65 years. | | | |
| Initiation – people aged between 66 and 80 years | | | |
| Therapy limited to 1 dose One dose for all people aged between 66 and 80 years inclusive from 1 April 2018 ar | nd 31 De | ecember | 2020. |
| Diagnostic Agents | | | |
| | | | |
| TUBERCULIN PPD [MANTOUX] TEST Inj 5 TU per 0.1 ml, 1 ml vial – 0% DV Oct-20 to 2024 0.0 | 0 | 1 | Tubersol |

PART III: OPTIONAL PHARMACEUTICALS

| Price (ex man. excl. GST) \$ | Per | Brand or Generic Manufacturer |
|------------------------------------|-----|-------------------------------------|
| | | |

Optional Pharmaceuticals

NOTE:

In addition to the products expressly listed here in Part III: Optional Pharmaceuticals, a range of hospital medical devices are listed in an addendum to Part III which is available at <u>www.pharmac.govt.nz</u>. The Optional Pharmaceuticals listed in the addendum are deemed to be listed in Part III, and the Rules of the Pharmaceutical Schedule applying to products listed in Part III apply to them.

| BLOOD GLUCOSE DIAGNOSTIC TEST METER | | |
|--|----------|--|
| 1 meter with 50 lancets, a lancing device, and 10 diagnostic test strips20.00 10.00 | 1 | CareSens N Premier Caresens N Caresens N POP |
| BLOOD GLUCOSE DIAGNOSTIC TEST STRIP | | |
| Blood glucose test strips10.56 | 50 test | CareSens N |
| Test strips 10.56 | 50 test | CareSens PRO |
| BLOOD KETONE DIAGNOSTIC TEST STRIP | | |
| Test strips | 10 strip | KetoSens |
| DUAL BLOOD GLUCOSE AND BLOOD KETONE DIAGNOSTIC TEST METER | | |
| Meter with 50 lancets, a lancing device, and 10 blood glucose diagnostic | | |
| test strips | 1 | CareSens Dual |
| MASK FOR SPACER DEVICE | | |
| Small | 1 | e-chamber Mask |
| PEAK FLOW METER | • | |
| Low Range | 1 | Mini-Wright AFS Low |
| Low Hange | I | Range |
| Normal Range9.54 | 1 | Mini-Wright Standard |
| PREGNANCY TEST - HCG URINE | | inin Might Clandard |
| Cassette | 40 test | Smith BioMed Rapid |
| Casselle | 40 1851 | Pregnancy Test |
| | | Tregnancy rest |
| SODIUM NITROPRUSSIDE | EQ atria | Kataatiw |
| Test strip22.00 | 50 strip | Ketostix |
| SPACER DEVICE | | |
| 220 ml (single patient) | 1 | e-chamber Turbo |
| 510 ml (single patient)5.12 | 1 | e-chamber La Grande |
| 800 ml | 1 | Volumatic |

- Symbols -

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| Adalat Oros |
| Adalimumab |
| Adapalene |
| Aderocor |
| Adenosine |
| Adenuric |
| Adrenaline |
| ADT Booster |
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