1. **PROLIA (60 mg solution for injection)**

Prolia 60 mg solution for injection pre-filled syringe

Prolia 60 mg solution for injection pre-filled syringe with automatic needle guard

2. **QUALITATIVE AND QUANTITATIVE COMPOSITION**

Each 1 mL single-use pre-filled syringe contains 60 mg of denosumab in 1 mL (60 mg/mL).

Denosumab is produced in genetically engineered mammalian (Chinese Hamster Ovary, CHO) cells.

**Excipient(s) with known effect**

Each mL of solution contains 47 mg sorbitol (E420) (see section 4.4.).

For the full list of excipients, see section 6.1.

3. **PHARMACEUTICAL FORM**

Solution for injection.

Prolia® is supplied as a sterile, preservative-free, clear, colourless to slightly yellow solution for injection at pH 5.2. The solution should not be used if cloudy or discoloured. The solution may contain trace amounts of translucent to white proteinaceous particles.

4. **CLINICAL PARTICULARS**

4.1 **Therapeutic indications**

The treatment of osteoporosis in postmenopausal women. Prolia® significantly reduces the risk of vertebral, non-vertebral and hip fractures.

Treatment to increase bone mass in men with osteoporosis at increased risk of fracture.

4.2 **Dose and method of administration**

Administration should be performed by an individual who has been adequately trained in injection techniques.

**Dose**

The recommended dose of Prolia® is a single subcutaneous (SC) injection of 60 mg, once every 6 months. If Prolia® treatment is discontinued, consider transitioning to an alternative antiresorptive therapy.

To reduce the risk of hypocalcaemia, patients must be adequately supplemented with calcium and vitamin D (see section 4.4, Hypocalcaemia). In the major clinical trials of
Prolia®, daily supplementation with 1000 mg of calcium and at least 400 IU vitamin D was recommended.

**Elderly**

No dose adjustment is necessary in elderly patients (see section 4.4, Use in the Elderly).

**Renal Impairment**

No dose adjustment is necessary in patients with renal impairment (see section 4.4, Renal Impairment).

**Hepatic Impairment**

The safety and efficacy of Prolia® has not been studied in patients with hepatic impairment.

**Paediatric Population**

The safety and efficacy of Prolia® in paediatric patients have not been established. Prolia® is not recommended for use in paediatric patients (see section 4.4, Paediatric Population).

**Method of administration**

Prolia® is a sterile and preservative-free product. Before administration, the Prolia® solution should be inspected for particulate matter and discolouration. Do not use if the solution is cloudy or discoloured. Do not excessively shake the pre-filled syringe. To avoid discomfort at the site of injection, allow the pre-filled syringe to reach room temperature (up to 25°C) before injecting, and inject slowly. Inject the entire contents of the pre-filled syringe.

Product is for single-use in one patient only. Dispose of any medicinal product remaining in the pre-filled syringe.

**4.3 Contraindications**

Hypocalcaemia (See section 4.4).

Hypersensitivity to the active substance, to CHO-derived proteins or to any of the excipients listed in section 6.1.

Pregnancy and in women trying to get pregnant (See section 4.6).
4.4 Special warnings and precautions for use

Hypocalcaemia

Hypocalcaemia must be corrected prior to initiating therapy with Prolia®. In the post-marketing setting, severe symptomatic hypocalcaemia has been reported (see section 4.8), with most cases occurring in the first weeks of initiating therapy, but it can occur later. Clinical monitoring of calcium levels is recommended before each dose.

In patients predisposed to hypocalcaemia (e.g. history of hypoparathyroidism, thyroid surgery, parathyroid surgery, malabsorption syndromes, excision of small intestine, severe renal impairment [creatinine clearance < 30 mL/min] or receiving dialysis), clinical monitoring of calcium levels is recommended during treatment, especially in the first two weeks of initiating therapy.

Hypocalcaemia following Prolia® administration is a significant risk in patients with severe renal impairment [creatinine clearance < 30 mL/min] or receiving dialysis.

Instruct all patients about the symptoms of hypocalcaemia and the importance of maintaining calcium levels with adequate calcium and vitamin D supplementation.

Adequate intake of calcium and vitamin D is important in all patients (see section 4.2 and section 4.8).

Skin Infections

Patients receiving Prolia® may develop skin infections (predominantly cellulitis) leading to hospitalisation (see section 4.8). Patients should be advised to seek prompt medical attention if they develop signs or symptoms of cellulitis.

Osteonecrosis of the Jaw

Osteonecrosis of the jaw (ONJ) has been reported in patients treated with denosumab or bisphosphonates, another class of antiresorptive agents. Most cases have been in cancer patients; however some have occurred in patients with osteoporosis.

ONJ has been reported rarely in clinical studies in patients receiving denosumab at a dose of 60 mg every 6 months for osteoporosis. There have been reports of ONJ in clinical studies in patients with advanced cancer treated with denosumab at the studied dose of 120 mg administered monthly.

Known risk factors for ONJ include a diagnosis of cancer with bone lesions, concomitant therapies (e.g. chemotherapy, antiangiogenic biologics, corticosteroids, radiotherapy to head and neck), poor oral hygiene, invasive dental procedures (e.g. tooth extraction),
and co-morbid disorders (e.g. pre-existing dental disease, anaemia, coagulopathy, infection). The risk of ONJ may increase with duration of exposure to Prolia®.

It is important to evaluate patients for risk factors for ONJ before starting treatment. If risk factors are identified, a dental examination with appropriate preventive dentistry is recommended prior to treatment with Prolia®. Good oral hygiene practices should be maintained during treatment with Prolia®.

Avoid invasive dental procedures during treatment with Prolia®. For patients in whom invasive dental procedures cannot be avoided, the clinical judgement of the treating physician should guide the management plan of each patient based on individual benefit/risk assessment.

Patients who are suspected of having or who develop ONJ while on Prolia® should receive care by a dentist or an oral surgeon. In patients who develop ONJ during treatment with Prolia®, a temporary interruption of treatment should be considered based on individual risk/benefit assessment until the condition resolves.

Atypical Femoral Fractures

Atypical femoral fractures have been reported in patients receiving Prolia®. Atypical femoral fractures may occur with little or no trauma in the subtrochanteric and diaphyseal regions of the femur and may be bilateral. Specific radiographic findings characterise these events. Atypical femoral fractures have also been reported in patients with certain co-morbid conditions (e.g. vitamin D deficiency, rheumatoid arthritis, hypophosphatasia) and with use of certain pharmaceutical agents (e.g. bisphosphonates, glucocorticoids, proton pump inhibitors). These events have also occurred without antiresorptive therapy. During Prolia® treatment, patients should be advised to report new or unusual thigh, hip, or groin pain. Patients presenting with such symptoms should be evaluated for an incomplete femoral fracture, and the contralateral femur should also be examined.

Multiple Vertebral Fractures (MVF) Following Discontinuation of Prolia® Treatment

Multiple vertebral fractures (MVF) may occur following discontinuation of treatment with Prolia®, particularly in patients with a history of vertebral fracture.

Patients being treated with Prolia®, should be advised not to interrupt Prolia® therapy without prior consultation with their treating physician. The individual benefit/risk should be evaluated before discontinuing treatment with Prolia®. If Prolia® treatment is discontinued, consider transitioning to an alternative antiresorptive therapy.
Medicines with Same Active Ingredient

Prolia® contains the same active ingredient found in Xgeva® (denosumab), used for the treatment of skeletal related events in patients with bone metastasis from solid tumours. Patients being treated with Prolia® should not be treated with Xgeva® concomitantly.

Excipients with Known Effects

Prolia® contains sorbitol. Patients with rare hereditary problems of fructose intolerance should not use Prolia®.

Prolia contains less than 1 mmol sodium (23 mg) per 60 mg, i.e. essentially ‘sodium free’.

Paediatric population

The safety and efficacy of Prolia® in paediatric patients have not been established. Prolia® is not recommended for use in paediatric patients. Adolescent primates dosed with denosumab at 27 and 150 times (10 and 50 mg/kg dose) the clinical exposure based on AUC had abnormal growth plates. In neonatal rats, inhibition of RANKL (target of denosumab therapy) with a construct of osteoprotegerin bound to immunoglobulin Fc segment (OPG-Fc) at high doses was associated with inhibition of bone growth and tooth eruption. Therefore, treatment with denosumab may impair bone growth in children with open growth plates and may inhibit eruption of dentition.

Use in renal impairment

No dose adjustment is necessary in patients with renal impairment.

In clinical studies, patients with severe renal impairment (creatinine clearance < 30 mL/min) or receiving dialysis were at greater risk of developing hypocalcaemia. Adequate intake of calcium and vitamin D is important in patients with severe renal impairment or receiving dialysis (see Hypocalcaemia).

Use in the Elderly

Of the total number of patients in clinical studies of Prolia®, 9,943 patients were ≥ 65 years, while 3,576 were ≥ 75 years. No overall differences in safety or efficacy were observed between these patients and younger patients.

Of the patients in the osteoporosis study in men, 133 patients (55%) were ≥ 65 years old, while 39 patients (16%) were ≥ 75 years old.
4.5 Interaction with other medicines and other forms of interaction

In an interaction study conducted on 17 postmenopausal women with osteoporosis, midazolam (2 mg oral) was administered two weeks after a single dose of denosumab (60 mg subcutaneous injection), which approximates the $T_{\text{max}}$ of denosumab. Prolia® did not affect the pharmacokinetics of midazolam, which is metabolised by cytochrome P450 3A4 (CYP3A4). This indicates that Prolia® should not alter the pharmacokinetics of medicines metabolised by CYP3A4.

No interactions with laboratory and diagnostic tests have been identified.

4.6 Fertility, pregnancy and lactation

Pregnancy

Pregnancy Category: D

There are no adequate and well-controlled studies of Prolia® in pregnant women. Prolia® is contraindicated for use during pregnancy and in women trying to get pregnant.

Developmental toxicity studies have been performed with denosumab in cynomolgus monkeys and have shown serious adverse events on the development (including fetal and infant lethality). Denosumab was shown to cross the placenta in monkeys.

In a study of cynomolgus monkeys with denosumab at subcutaneous doses up to 12.5 mg/kg/week, given during the period equivalent to the first trimester, and yielding AUC exposures up to 99-fold higher than the human exposure (60 mg every 6 months), there was no evidence of maternal or fetal harm. In this study fetal lymph nodes were not examined.

In another study of cynomolgus monkeys dosed with denosumab throughout pregnancy at 50 mg/kg/month, yielding AUC exposures 119-fold higher than the human exposure, there were increased stillbirths and postnatal mortality; abnormal bone growth resulting in reduced bone strength, almost complete obliteration of bone marrow spaces (leading to reduced haematopoiesis), and tooth malalignment, dental dysplasia and a shortened/straighter dental arch (although no effect on the pattern or date of tooth eruption); altered appearance of eyes (increased apparent size, exophthalmos); absence of peripheral lymph nodes; and decreased neonatal growth. Following a 6 month period after birth, bone-related changes showed incomplete recovery. The effects on lymph nodes, tooth malalignment and dental dysplasia persisted, and minimal to moderate mineralisation in multiple tissues was seen in one animal. There was no evidence of maternal harm prior to labour; adverse maternal effects occurred infrequently during
labour. Maternal mammary gland development was normal. A no observed adverse effect level has not been established in animal studies and the findings are attributable to the primary pharmacological activity of denosumab.

Preclinical studies in RANK/RANKL knockout mice suggest absence of RANKL could interfere with the development of lymph nodes in the fetus. Knockout mice lacking RANK or RANKL also exhibited decreased body weight, reduced bone growth and a lack of tooth eruption. Similar phenotypic changes (inhibition of bone growth and tooth eruption) were observed in a study in neonatal rats using a surrogate for denosumab, the RANKL inhibitor osteoprotegerin bound to Fc (OPG-Fc). Therefore, treatment with denosumab may impair bone growth in children with open growth plates and may inhibit eruption of dentition. The reversibility of the effects of OPG-Fc has not been examined.

Preclinical studies in RANK/RANKL knockout mice suggest absence of RANKL during pregnancy may interfere with maturation of the mammary gland leading to impaired lactation post-partum.

**Breast-feeding**

It is unknown whether denosumab is excreted in human milk. Only limited excretion of denosumab in milk was observed in a study in monkeys. A decision on whether to abstain from breast-feeding or to abstain from therapy with Prolia® should be made, taking into account the benefit of breast-feeding to the newborn/infant and the benefit of Prolia® therapy to the woman.

**Fertility**

No data are available on the effect of denosumab on human fertility. Denosumab had no effect on female fertility or male reproductive organs or sperm motility in cynomolgus monkeys at subcutaneous doses up to 12.5 mg/kg/week (females) or 50 mg/kg/month (males), yielding exposures that were approximately 150-fold higher than the human exposure at 60 mg subcutaneous administered once every 6 months.

**4.7 Effects on ability to drive and use machines**

No studies on the effects on the ability to drive or use machinery have been performed.
4.8 Undesirable effects

Summary of safety profile

Treatment of Postmenopausal Osteoporosis

Prolia® has been studied in over 10,500 women with postmenopausal osteoporosis in clinical trials representing up to 8 years of continued Prolia® treatment.

The safety of Prolia® in the treatment of postmenopausal osteoporosis was assessed in FREEDOM, a large, 3-year, randomised, double-blind, placebo-controlled, multinational phase III study of 7,808 postmenopausal women aged 60 to 91 years with osteoporosis. A total of 3,886 women were exposed to Prolia® and 3,876 women were exposed to placebo administered once every 6 months as a single 60 mg subcutaneous dose.

The safety of Prolia® was also assessed in a second phase 3 study of similar design. A total of 322 postmenopausal women aged 43 to 83 years with low bone mass were enrolled in this 2-year study. A total of 164 women were exposed to Prolia® and 165 women were exposed to placebo administered once every 6 months as a single 60 mg subcutaneous dose.

In both studies, all women received at least 1000 mg of calcium and 400 IU of vitamin D supplementation per day.

Across the two phase III studies the incidence of all-cause mortality was 1.7% (n = 70) in the Prolia® group and 2.2% (n = 90) in the placebo group. The incidence of serious adverse events was 25.3% in the Prolia® group and 24.3% in the placebo group. The percentage of patients who withdrew from the studies due to adverse events was 2.3% and 2.1% for the Prolia® and placebo groups, respectively.

The most common adverse events reported in studies of women with postmenopausal osteoporosis or low bone mass (n = 8,091), occurring in ≥ 10% of patients either in the Prolia®-treated or placebo group, were back pain (34.1% Prolia®, 34.0% placebo), arthralgia (20.4% in each group), hypertension (15.3% Prolia®, 16.1% placebo), nasopharyngitis (14.8% Prolia®, 15.6% placebo), pain in extremity (11.8% Prolia®, 11.2% placebo) and osteoarthritis (10.9% Prolia®, 11.1% placebo).

Adverse events reported in at least 2% of postmenopausal women with osteoporosis or low bone mass (n = 8,091) and at least 1% more frequently in the Prolia®-treated women than in the placebo-treated women were: hypercholesterolemia (7.0% Prolia®, 5.9% placebo) and eczema (includes dermatitis, allergic dermatitis, atopic dermatitis and contact dermatitis) (3.1% Prolia®, 1.7% placebo).
In STAND, a double-blind, randomised, alendronate-controlled, study in postmenopausal women with low bone mass who had received alendronate for at least 6 months preceding study entry, patients received either Prolia® 60 mg Q6M SC (n = 253) or alendronate orally 70 mg weekly for 12 months (n = 249). The safety profile was similar for patients transitioning from alendronate to denosumab and those continuing on alendronate therapy, including the overall incidence of adverse events and serious adverse events. Eight patients (3.2%) in the Prolia® group and 4 patients (1.6%) in the alendronate group reported adverse events of fracture.

Tabulated list of adverse reactions

The data in table 1 describe adverse reactions reported from Phase II and III clinical trials in patients with osteoporosis and breast or prostate cancer patients receiving hormone ablation, and/or spontaneous reporting.

<table>
<thead>
<tr>
<th>System Organ Class</th>
<th>Very common ≥ 10%</th>
<th>Common ≥ 1% and &lt; 10%</th>
<th>Uncommon ≥ 0.1% and &lt; 1%</th>
<th>Rare ≥ 0.01% and &lt; 0.1%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immune system disorders</td>
<td></td>
<td></td>
<td></td>
<td>Hypersensitivity reactions\a</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td></td>
<td></td>
<td>Skin infections\b leading to hospitalization</td>
<td></td>
</tr>
<tr>
<td>Metabolism and nutrition disorders</td>
<td></td>
<td></td>
<td></td>
<td>Hypocalcaemia</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Severe symptomatic hypocalcaemia\c</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Eczema\d</td>
<td></td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>Pain in extremity Musculoskeletal pain\e</td>
<td>Multiple vertebral fractures following discontinuation of Prolia\f</td>
<td>Osteonecrosis of the jaw Atypical femoral fracture</td>
<td></td>
</tr>
</tbody>
</table>

\a including rash, urticarial, facial swelling, erythema and anaphylactic reactions
\b predominantly cellulitis
\c reported in patients at increased risk of hypocalcaemia receiving Prolia
\d includes dermatitis, allergic dermatitis, atopic dermatitis, and contact dermatitis
\e including severe cases
\f particularly in those with a history of vertebral fracture
Description of selected adverse reactions

**Hypocalcaemia**

In two phase III placebo-controlled clinical trials in postmenopausal women with osteoporosis, approximately 0.05% (2 of 4,050) of patients had declines of serum calcium levels (less than 1.88 mmol/L) following Prolia® administration.

**Skin Infections**

In two phase III placebo-controlled clinical trials in postmenopausal women with osteoporosis, skin infections leading to hospitalisation were reported more frequently in the Prolia® (0.4%, 16 of 4,050) versus the placebo (0.1%, 3 of 4,041) groups, respectively. These cases were predominantly cellulitis. The overall incidence of skin infections was similar between the Prolia® (1.5%, 59 of 4,050) and placebo groups (1.2%, 50 of 4,041).

**Pancreatitis**

Pancreatitis was reported in 4 patients (0.1%) in the placebo and 8 patients (0.2%) in the Prolia® groups. Several patients had a prior history of pancreatitis or a confounding event (e.g. gallstones). The time from product administration to event occurrence was variable.

**Osteonecrosis of the Jaw (ONJ)**

In the osteoporosis clinical trial program, ONJ was reported rarely in patients treated with Prolia®.

**Atypical Femoral Fractures**

In the osteoporosis clinical trial program, atypical femoral fractures were reported very rarely in patients treated with Prolia®.

**Multiple Vertebral Fractures (MVF) Following Discontinuation of Prolia® Treatment**

In the osteoporosis clinical trial program, MVF were reported in patients following discontinuation of treatment with Prolia®, particularly in those with a history of vertebral fracture.

**Long Term Safety in Postmenopausal Osteoporosis**

A total of 4550 women who completed FREEDOM (Study 20030216, N = 7808) enrolled into FREEDOM Extension (Study 20060289), a 7-year, multinational, multicenter, open-label, single-arm extension study to evaluate the long-term safety and efficacy of Prolia®.
All patients in the extension study receive Prolia® every 6 months as a single SC 60 mg dose, as well as daily calcium (1000 mg) and vitamin D (at least 400 IU).

During the FREEDOM Extension study, the rates of adverse events observed through month 60 have not shown an increase over time and were similar to those observed in the initial 3 years of FREEDOM. Eight adjudicated cases of osteonecrosis of the jaw (ONJ) and two atypical fractures of the femur have occurred during the extension study.

**Treatment of Osteoporosis in Men**

The safety of Prolia® in the treatment of men with osteoporosis was assessed in ADAMO, a randomised, double-blind, placebo-controlled study; a 1 year double-blind phase followed by a 1 year open-label extension. During the double-blind phase, a total of 120 men were exposed to Prolia® and 120 men were exposed to placebo administered subcutaneously once every 6 months as a single 60 mg dose. All men were instructed to take at least 1000 mg of calcium and 800 IU of vitamin D supplementation per day.

The incidence of all-cause mortality was 0.8% (n = 1) in the Prolia® group and 0.8% (n = 1) in the placebo group. The incidence of serious adverse events was 9.2% in the Prolia® group and 8.3% in the placebo group. The percentage of patients who withdrew from the study due to adverse events was 2.5% and 0% for the Prolia® and placebo groups, respectively.

Adverse events in men with osteoporosis (n=240) occurring in at least 5% of Prolia®-treated men and more frequently than in the placebo-treated patients were: back pain (8.3% Prolia®, 6.7% placebo), arthralgia (6.7% Prolia®, 5.8% placebo), and nasopharyngitis (6.7% Prolia®, 5.8% placebo).

**Postmarketing Experience**

Rare events of medicine-related hypersensitivity reactions: rash, urticaria, facial swelling, erythema and anaphylactic reactions.

Rare events of severe symptomatic hypocalcaemia have been reported in patients at increased risk of hypocalcaemia. Examples of the clinical manifestations of severe symptomatic hypocalcaemia have included QT prolongation, tetany, seizures and altered mental status (see PRECAUTIONS, Hypocalcaemia). Symptoms of hypocalcaemia in denosumab clinical studies included paraesthesia, muscle stiffness, twitching, spasms and muscle cramps.
Musculoskeletal pain, including severe cases, has been reported in patients receiving Prolia.

**Reporting of suspected adverse reactions**

Reporting suspected adverse reactions after authorization of the medicine is important. It allows continued monitoring of the benefit/risk balance of the medicine. Healthcare professionals are asked to report any suspected adverse reactions to [https://nzphvc.otago.ac.nz/reporting/](https://nzphvc.otago.ac.nz/reporting/)

### 4.9 Overdose

There is no experience with overdosage with Prolia®. Prolia® has been administered in clinical studies using doses up to 180 mg every 4 weeks (cumulative doses up to 1080 mg over 6 months), and no additional adverse effects were observed.

For advice on the management of overdose please contact the National Poisons Centre on 0800 POISON (0800 764766).

### 5. PHARMACOLOGICAL PROPERTIES

#### 5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Drugs for the treatment of bone diseases – Other drugs affecting bone structure and mineralisation, ATC code: M05BX04

Denosumab is a fully human IgG2 monoclonal antibody with high affinity and specificity for RANK ligand (RANKL). Denosumab has an approximate molecular weight of 147 kDa.

CAS number: 615258-40-7

Prolia® is the Amgen Inc. trademark for denosumab (rch).
Pharmacodynamic effects

Mechanism of Action

RANKL exists as a transmembrane or soluble protein. RANKL is essential for the formation, function and survival of osteoclasts, the sole cell type responsible for bone resorption. Osteoclasts play an important role in bone loss associated with postmenopausal osteoporosis and hormone ablation. Denosumab binds with high affinity and specificity to RANKL, preventing RANKL from activating its only receptor, RANK, on the surface of osteoclasts and their precursors, independent of bone surface. Prevention of RANKL/RANK interaction inhibits osteoclast formation, function and survival, thereby decreasing bone resorption and increasing bone mass and strength in both cortical and trabecular bone.

Pharmacodynamics

In clinical studies, treatment with 60 mg of Prolia® resulted in rapid reduction in the bone resorption marker serum type 1 C-telopeptides (CTX) within 6 hours of SC administration by approximately 70%, with reductions of approximately 85% occurring by 3 days. CTX reductions were maintained over the 6-month dosing interval. At the end of each dosing interval, CTX reductions were partially attenuated from maximal reduction of ≥ 87% to ≥ 45% (range 45% to 80%), reflecting the reversibility of the effects of Prolia® on bone remodelling once serum denosumab levels diminish. These effects were sustained with continued treatment. Consistent with the physiological coupling of bone formation and
resorption in skeletal remodeling, subsequent reductions in bone formation markers (e.g. bone specific alkaline phosphatase [BSAP] and serum N-terminal propeptide of type 1 collagen [P1NP]) were observed beginning 1 month after the first dose of Prolia®.

Bone turnover markers (bone resorption and formation markers) generally reached pretreatment levels within 9 months after the last 60 mg subcutaneous dose. Upon re-initiation, the degree of inhibition of CTX by Prolia® was similar to those observed in patients initiating Prolia®.

In a clinical study of postmenopausal women with low bone mass (n = 504) who were previously treated with alendronate for a median of 3 years, those transitioning to receive Prolia® experienced additional reductions in serum CTX, compared with women who remained on alendronate. In this study, the changes in serum calcium were similar between the two groups.

Clinical efficacy and safety

Treatment of osteoporosis in postmenopausal women

Independent risk factors, for example, low bone mineral density (BMD), age, the existence of previous fracture, family history of hip fractures, high bone turnover and low body mass index (BMI) should be considered in order to identify women at increased risk of osteoporotic fractures who could benefit from treatment.

Fracture REduction Evaluation of Denosumab in Osteoporosis every 6 Months (FREEDOM): The efficacy and safety of Prolia® in the treatment of postmenopausal osteoporosis was demonstrated in FREEDOM (Study 20030216), a 3-year, randomised, double-blind, placebo-controlled, multinational study of women with baseline BMD T-scores at the lumbar spine or total hip between -2.5 and -4.0. 7,808 women aged 60 to 91 years were enrolled of whom 23.6% had prevalent vertebral fractures. Women with other diseases or on therapies that may affect bone (e.g. rheumatoid arthritis, osteogenesis imperfecta and Paget’s disease) were excluded from this study.

BMD and other individual risk factors were collected for women enrolled in the FREEDOM study. The mean absolute 10-year fracture probability for women enrolled was 18.60% (deciles: 7.9-32.4%) for major osteoporotic fracture and 7.22% (deciles: 1.4-14.9%) for hip fracture, as derived from FRAX®, the WHO Fracture Risk Assessment Tool algorithm.

Women were randomised to receive subcutaneous injections of either Prolia® 60 mg (n = 3,902) or placebo (n = 3,906) once every 6 months. Women received calcium (at least
1,000 mg) and vitamin D (at least 400 IU) supplementation daily. The primary efficacy variable was the incidence of new vertebral fractures. Secondary efficacy variables included the incidence of non-vertebral fractures and hip fractures, assessed at 3 years.

**Effect on vertebral fractures**

Prolia® significantly reduced the risk of new vertebral fractures at 1, 2 and 3 years (p < 0.0001) (see Table 2).

<table>
<thead>
<tr>
<th></th>
<th>Proportion of women with fracture (%)</th>
<th>Absolute risk reduction (%) (95% CI)</th>
<th>Relative risk reduction (%) (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolia</td>
<td>n = 3,902 (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>n = 3,906 (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0-1 Year</td>
<td>0.9 (2.2)</td>
<td>1.4 (0.8, 1.9)</td>
<td>61 (42, 74)*</td>
</tr>
<tr>
<td>0-2 Years</td>
<td>1.4 (5.0)</td>
<td>3.5 (2.7, 4.3)</td>
<td>71 (61, 79)*</td>
</tr>
<tr>
<td>0-3 Years</td>
<td>2.3 (7.2)</td>
<td>4.8 (3.9, 5.8)</td>
<td>68 (59, 74)*</td>
</tr>
</tbody>
</table>

*p < 0.0001

The reductions in the risk of new vertebral fractures by Prolia® over 3 years were consistent and significant regardless of whether or not women had a prevalent vertebral fracture or history of a non-vertebral fracture, and regardless of baseline age, BMD, bone turnover level and prior use of a medicinal product for osteoporosis.

Prolia® also reduced the risk of new vertebral fracture by 65% (6.5% absolute risk reduction, p < 0.0001) in patients at high risk of fractures (defined as women who met ≥ 2 of the 3 following criteria at baseline: age ≥ 70 years, BMD T-score ≤ -3.0 [at lumbar spine, total hip, or femoral neck] or prevalent vertebral fracture).

Prolia® also reduced the risk of new and worsening vertebral fractures (67% relative risk reduction, 4.8% absolute risk reduction) as well as multiple vertebral fractures (61% relative risk reduction, 1.0% absolute risk reduction) at 3 years, when compared to placebo (all p < 0.0001).

**Effect on hip fractures**

Prolia® demonstrated a 40% relative reduction (0.5% absolute risk reduction) in the risk of hip fracture over 3 years (p < 0.05) (see Figure 1). The incidence of hip fracture was 0.7% in the Prolia® group compared to 1.2% in the placebo group at 3 years.
In women with high fracture risk as defined above by baseline age, BMD and prevalent vertebral fracture, a 48% relative risk reduction was observed with Prolia® (1.1% absolute risk reduction, p < 0.05).

**Effect on all clinical fractures**

Prolia® demonstrated superiority to placebo in reducing the risk of any clinical fractures, clinical (symptomatic) vertebral fractures, non-vertebral fractures (including hip), major non-vertebral fractures and major osteoporotic fractures (see Table 3).
Table 3. The effect of Prolia® on the risk of clinical fractures over 3 years

<table>
<thead>
<tr>
<th>Event</th>
<th>Prolia n = 3,902 (%)</th>
<th>Placebo n = 3,906 (%)</th>
<th>Absolute risk reduction (%) (95% CI)</th>
<th>Relative risk reduction (%) (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any clinical fracture¹</td>
<td>7.2</td>
<td>10.2</td>
<td>2.9 (1.6, 4.2)</td>
<td>30 (19, 41)***</td>
</tr>
<tr>
<td>Clinical vertebral fracture</td>
<td>0.8</td>
<td>2.6</td>
<td>1.8 (1.2, 2.4)</td>
<td>69 (53, 80)***</td>
</tr>
<tr>
<td>Non-vertebral fracture²</td>
<td>6.5</td>
<td>8.0</td>
<td>1.5 (0.3, 2.7)</td>
<td>20 (5, 33)**</td>
</tr>
<tr>
<td>Major non-vertebral fracture³</td>
<td>5.2</td>
<td>6.4</td>
<td>1.2 (0.1, 2.2)</td>
<td>20 (3, 34)*</td>
</tr>
<tr>
<td>Major osteoporotic fracture⁴</td>
<td>5.3</td>
<td>8.0</td>
<td>2.7 (1.6, 3.9)</td>
<td>35 (22, 45)***</td>
</tr>
</tbody>
</table>

*p ≤ 0.05; **p = 0.0106, ***p ≤ 0.0001
+ Event rates based on Kaplan-Meier estimates at 3 years
(1) Includes clinical vertebral fractures and non-vertebral fractures
(2) Excludes those of the vertebrae (cervical, thoracic, and lumbar), skull, facial, mandible, metacarpus, and finger and toe phalanges
(3) Includes pelvis, distal femur (i.e. femur excluding hip), proximal tibia (i.e. tibia excluding ankle), ribs, proximal humerus (i.e. humerus excluding elbow), forearm, and hip
(4) Includes clinical vertebral, hip, forearm, and humerus fractures, as defined by the WHO

Women in the FREEDOM study had a mean baseline BMD T-score of -2.2 at the femoral neck. In women with baseline femoral neck BMD ≤ -2.5, Prolia® reduced the incidence of non-vertebral fracture (35% relative risk reduction, 4.1% absolute risk reduction, p < 0.001).

The reduction in the incidence of new vertebral fractures, hip fractures and non-vertebral fractures by Prolia® over 3 years were consistent regardless of the 10-year baseline fracture risk as assessed by FRAX.

**Effect on bone mineral density**

Prolia® significantly increased BMD at all clinical sites measured, relative to treatment with placebo at 1, 2 and 3 years in FREEDOM. Prolia® increased BMD by 9.2% at the lumbar spine, 6.0% at the total hip, 4.8% at the femoral neck, 7.9% at the hip trochanter, 3.5% at the distal 1/3 radius and 4.1% at the total body over 3 years (all p < 0.0001).

Increases in BMD at lumbar spine, total hip and hip trochanter were observed as early as
1 month after the initial dose. Prolia® increased lumbar spine BMD from baseline in 95% of postmenopausal women at 3 years. Consistent effects on BMD were observed at the lumbar spine regardless of baseline age, race, weight/BMI, BMD and bone turnover level. The effects of Prolia® on bone architecture were evaluated using quantitative computed tomography (QCT) in postmenopausal women with BMD T-score below -2.5 at the lumbar spine or total hip. Treatment with Prolia® increased volumetric trabecular BMD at the lumbar spine, volumetric BMD at the total hip and the volumetric cortical BMD and cortical thickness at the distal radius.

Study of Transitioning from Alendronate to Denosumab (STAND, Study 20050234) was a double-blind, randomised, alendronate-controlled, study in postmenopausal women with low BMD (T score between -2.0 and -4.0 at the lumbar spine or total hip) who had received alendronate (70 mg weekly [or equivalent] orally) for at least 6 months preceding study entry. Patients received either Prolia® 60 mg Q6M SC (n = 253) or alendronate orally 70 mg weekly for 12 months (n = 251).

Women who transitioned to receive Prolia® had greater increases in BMD at the total hip (1.9% versus 1.1%, p < 0.001; primary efficacy endpoint) after 1 year, compared to those who continued to receive alendronate therapy. Consistently greater increases in BMD were also seen at the lumbar spine, femoral neck, hip trochanter, and distal 1/3 radius in women treated with Prolia®, compared to those who continued to receive alendronate therapy (all p < 0.05).

In clinical studies examining the effects of discontinuation of Prolia®, BMD returned to approximately pre-treatment levels and remained above placebo within 18 months of the last dose. These data indicate that continued treatment with Prolia® is required to maintain the effect of the medicine. Re-initiation of Prolia® resulted in gains in BMD similar to those when Prolia® was first administered.

**Open-label extension study in the treatment of postmenopausal osteoporosis (FREEDOM Extension Study)**

A total of 4550 women, (2343 Prolia® and 2207 placebo) who missed no more than one dose of Prolia® in the FREEDOM pivotal study (Study 20030216, N = 7808) and completed the month 36 study visit, enrolled in FREEDOM Extension (Study 20060289), a 7-year, multinational, multicenter, open-label, single-arm extension study to evaluate the long-term safety and efficacy of Prolia®. All women in the FREEDOM Extension study receive Prolia® every 6 months in an open-label manner as a single 60 mg SC dose, as well as daily calcium (at least 1000 mg) and vitamin D (at least 400 IU). Safety
was the primary endpoint; BMD and fracture incidence were two of the many secondary endpoints. At month 60 of the extension study, after 8 years of Prolia® treatment, the long-term group increased BMD by 18.4% (95% CI: 18.0, 18.8) at the lumbar spine, 8.3% (8.0, 8.5) at the total hip, 7.8% (7.5, 8.1) at the femoral neck and 11.6% (11.2, 12.0) at the trochanter from the pivotal FREEDOM study baseline. In years 4 through 8 of Prolia treatment, the rates of new vertebral and non-vertebral fractures did not increase over time; annualised rates were approximately 1.1% and 1.3% respectively.

**Bone Histology**

Fifty-three trans-iliac crest bone biopsy specimens were obtained at either 2 years and/or 3 years from 47 postmenopausal women with osteoporosis treated with Prolia® in the FREEDOM study. Fifteen bone biopsy specimens were also obtained after 1 year of treatment with Prolia® from 15 postmenopausal women with low bone mass who had transitioned from previous alendronate therapy. Histology assessments in both studies showed bone of normal architecture and quality, as well as the expected decrease in bone turnover relative to placebo treatment. There was no evidence of mineralisation defects, woven bone or marrow fibrosis.

Forty-one women participated in the bone biopsy sub-study at month 24 of the FREEDOM extension study, representing up to 5 years of treatment with Prolia®. Bone biopsy results showed bone of normal architecture and quality with no evidence of mineralisation defects, woven bone or marrow fibrosis as well as the expected decrease in bone turnover.

**Treatment of Osteoporosis in Men**

A Multicenter Randomised Double-blind Placebo Controlled Study to Compare the Efficacy and Safety of Denosumab versus Placebo in Males with Osteoporosis (ADAMO):

The efficacy and safety of Prolia® in the treatment of men with osteoporosis was demonstrated in ADAMO (Study 20080098), a 1-year, multinational study of men with low bone mass, who had a baseline BMD T-score between -2.0 and -3.5 at the lumbar spine or femoral neck. Men with a BMD T-score between -1.0 and -3.5 at the lumbar spine or femoral neck and with history of prior fragility fracture were also enrolled. Men with other diseases (such as rheumatoid arthritis, osteogenesis imperfecta, and Paget’s disease), or with significantly impaired renal function (GFR of ≤ 30 mL/min), or on therapies that may affect bone were excluded from this study.
### Table 4. Baseline BMD T-scores (Randomised Subjects)

<table>
<thead>
<tr>
<th></th>
<th>Prolia® (N = 121)</th>
<th>Placebo (N = 121)</th>
<th>All (N = 242)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum BMD T-score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>at lumbar spine or</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>femoral neck</td>
<td>n (%)</td>
<td>n (%)</td>
<td>n (%)</td>
</tr>
<tr>
<td>≤ -2.5</td>
<td>61 (50)</td>
<td>56 (46)</td>
<td>117 (48)</td>
</tr>
<tr>
<td>&gt; -2.5</td>
<td>60 (50)</td>
<td>65 (54)</td>
<td>125 (52)</td>
</tr>
</tbody>
</table>

N = number of subjects randomised.

The 242 men enrolled in the ADAMO study ranged in age from 31 to 84 years and were randomised to receive subcutaneous injections of either Prolia® 60 mg (n = 121) or placebo (n = 121) once every 6 months. Men received calcium (at least 1000 mg) and vitamin D (at least 800 IU) supplementation daily.

The primary efficacy variable was percent change in lumbar spine BMD at 1 year. Secondary efficacy variables included percent change in total hip, hip trochanter, femoral neck, and distal 1/3 radius BMD at 1 year, and change in CTX at day 15.

Prolia® significantly increased BMD at all clinical sites measured, relative to treatment with placebo at 1 year in men with osteoporosis. Prolia® increased BMD by 4.8% at the lumbar spine, 2.0% at the total hip, 2.3% at the hip trochanter, 2.2% at the femoral neck and 0.9% at the distal 1/3 radius, relative to placebo. Consistent effects on BMD were observed at the lumbar spine regardless of baseline age, race, weight/body mass index (BMI), BMD, and level of bone turnover.

**Bone Histology**

A total of 29 trans-iliac crest bone biopsy specimens were obtained from men with osteoporosis at 12 months (17 specimens in Prolia® group, 12 specimens in placebo group). Qualitative histology assessments showed normal architecture and quality with no evidence of mineralisation defects, woven bone, or marrow fibrosis.

### 5.2 Pharmacokinetic properties

Following a 60 mg subcutaneous dose of denosumab, bioavailability was 61% and maximum serum denosumab concentrations (C\text{max}) of 6 μg/mL (range 1-17 μg/mL) occurred in 10 days (range 2-28 days). After C\text{max}, serum levels declined with a half-life of 26 days (range 6-52 days) over a period of 3 months (range 1.5-4.5 months). Fifty-three percent of patients had no measurable amounts of denosumab detected at 6 months post-dose.
No accumulation or change in denosumab pharmacokinetics over time was observed upon subcutaneous multiple-dosing of 60 mg once every 6 months. Denosumab pharmacokinetics was not affected by the formation of binding antibodies to denosumab and was similar in men and women.

Pharmacokinetic analysis was performed to evaluate the effects of demographic characteristics. This analysis showed no notable difference in pharmacokinetics with age (28 to 87 years), race or body weight (36 to 140 kg).

Denosumab is composed solely of amino acids and carbohydrates as native immunoglobulin and is unlikely to be eliminated via hepatic metabolic mechanisms. Based on nonclinical data, its metabolism and elimination are expected to follow the immunoglobulin clearance pathways, resulting in degradation to small peptides and individual amino acids.

**Immunogenicity**

In clinical studies, no neutralising antibodies for denosumab have been observed. Using a sensitive immunoassay < 1% of patients treated with denosumab for up to 5 years tested positive for non neutralising binding antibodies with no evidence of altered pharmacokinetics, toxicity, or clinical response.

**Paediatric population**

The pharmacokinetic profile has not been assessed in those ≤ 18 years.

**Elderly**

The pharmacokinetics of denosumab was not affected by age (28 to 87 years).

**Impaired hepatic function**

The pharmacokinetic profile has not been assessed in patients with impaired hepatic function.

**Impaired renal function**

In a study of 55 patients with varying degrees of renal function, including patients on dialysis, the degree of renal impairment had no effect on the pharmacokinetics of denosumab (see section 4.4, Hypocalcaemia and section 4.2, Renal Impairment).
5.3 Preclinical safety data

Carcinogenicity

The carcinogenic potential of denosumab has not been evaluated in long-term animal studies. In view of the mechanism of action of denosumab, it is unlikely that the molecule would be capable of inducing tumor development or proliferation.

Genotoxicity

The genotoxic potential of denosumab has not been evaluated. Denosumab is a recombinant protein comprised entirely of naturally occurring amino acids and contains no inorganic or synthetic organic linkers or other non-protein portions. Therefore, it is unlikely that denosumab or any of its derived fragments would react with DNA or other chromosomal material.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Sorbitol

Acetate

Polysorbate 20

Sodium hydroxide for adjusting pH

Water for Injection (USP)

Prolia® is a sterile, preservative-free, clear, colourless to slightly yellow solution for injection. The solution may contain trace amounts of translucent to white proteinaceous particles.

6.2 Incompatibilities

In the absence of compatibility studies, this medicine must not be mixed with other medicines.

6.3 Shelf life

36 months

If removed from the refrigerator, Prolia® should be kept at room temperature (up to 25°C) in the original container and must be used within 30 days.
6.4 Special precautions for storage
It is recommended to store pre-filled syringes in a refrigerator at 2° to 8°C in the original carton. Do not freeze. Protect from direct light. Do not excessively shake the pre-filled syringe. Do not expose to temperatures above 25°C.

6.5 Nature and contents of container
Pre-filled syringe with automatic needle guard;
Pre-filled syringe*

Pack size of one, presented in blistered (pre-filled syringe with or without an automatic needle guard) or unblistered packaging (pre-filled syringe only).

The needle cover of the pre-filled syringe contains dry natural rubber (a derivative of latex), which may cause allergic reactions.

* Not available in New Zealand.

6.6 Special precautions for disposal and other handling
Any unused medicine or waste material should be disposed of in accordance with local requirements.

7. MEDICINE SCHEDULE
Prescription Medicine

8. SPONSOR
Amgen (New Zealand) Limited
Level 20, Lumley Centre
88 Shortland Street
Auckland
New Zealand
Telephone: 0800 443 885
Email: medinfo.JAPAC@amgen.com

9. DATE OF FIRST APPROVAL
Date of publication in the New Zealand Gazette of consent to distribute the medicine:
11 August 2011
**DATE OF REVISION OF THE TEXT**
8 March 2018

**SUMMARY TABLE OF CHANGES**

<table>
<thead>
<tr>
<th>Section changed</th>
<th>Summary of new information</th>
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<tbody>
<tr>
<td>Multiple</td>
<td>Reformatted and updated to align with new data sheet template requirements</td>
</tr>
<tr>
<td>Availability status</td>
<td>Deletion of text ‘This medicine is not currently marketed in New Zealand’.</td>
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