PRODUCT MONOGRAPH

Risedronate Sodium (as the hemi-pentahydrate) Tablets, USP 5 mg, 30 mg, 35 mg, 75 mg and 150 mg

Risedronate Sodium (as the hemi-pentahydrate) Delayed-Release Tablets 35 mg
PART I: HEALTH PROFESSIONAL INFORMATION

SUMMARY PRODUCT INFORMATION

<table>
<thead>
<tr>
<th>Route of Administration</th>
<th>Dosage Form/Strength</th>
<th>Clinically Relevant Nonmedicinal Ingredients</th>
</tr>
</thead>
<tbody>
<tr>
<td>oral</td>
<td>ACTONEL film-coated tablet 5 mg, 30 mg, 35 mg, 75 mg and 150 mg</td>
<td>lactose monohydrate (5 mg, 30 mg and 35 mg)</td>
</tr>
<tr>
<td>oral</td>
<td>ACTONEL DR enteric-coated, delayed-release tablet 35 mg</td>
<td>Edetate disodium (EDTA)</td>
</tr>
</tbody>
</table>

For a complete listing see DOSAGE FORMS, COMPOSITION AND PACKAGING section.

INDICATIONS AND CLINICAL USE

ACTONEL (risedronate sodium hemi-pentahydrate) is indicated for:

- the treatment and prevention of osteoporosis in postmenopausal women
- the treatment of osteoporosis in men, to improve bone mineral density
- the treatment and prevention of glucocorticoid-induced osteoporosis in men and women
- Paget’s disease of bone

ACTONEL DR (risedronate sodium hemi-pentahydrate) is indicated for:

- the treatment of osteoporosis in postmenopausal women

Postmenopausal Osteoporosis: In the treatment of osteoporosis in postmenopausal women at risk of fracture, ACTONEL and ACTONEL DR prevent vertebral and nonvertebral osteoporosis-related (fragility) fractures and increase bone mineral density (BMD) at all measured skeletal sites of clinical importance for osteoporotic fractures, including spine, hip, and wrist.

Osteoporosis may be confirmed by the presence or history of osteoporotic fracture, or by the finding of low bone mass (e.g., at least 2 standard deviation [SD] below the premenopausal mean).

For the prevention of osteoporosis in postmenopausal women who are at risk of developing osteoporosis, ACTONEL preserves or increases BMD at sites of clinical importance.

ACTONEL may be considered in postmenopausal women who are at risk of developing osteoporosis and for whom the desired clinical outcome is to maintain bone mass and to reduce the risk of fracture.

Factors such as family history of osteoporosis (particularly maternal history), age, previous fracture, smoking, moderately low BMD, high bone turnover, thin body frame, Caucasian or Asian race, and early menopause are associated with an increased risk of developing osteoporosis and fractures.

The optimal duration of use has not been determined. Patients should have the need for continued therapy re-evaluated on a periodic basis.

Paget’s Disease of Bone: ACTONEL is indicated for patients with Paget’s disease of bone (osteitis deformans) having alkaline phosphatase levels at least two times the upper limit of normal, or who are asymptomatic, or who are at risk for future complications from their disease, to induce remission (normalization of serum alkaline phosphatase).

Geriatrics: In ACTONEL and ACTONEL DR osteoporosis studies, 26-46% of patients were between 65 and 75 years of age and 10-23% were over 75 years of age. No overall differences in efficacy or safety were observed between these patients and younger patients (< 65 years) in the above osteoporosis studies. (See CLINICAL TRIALS section).

Pediatrics: Safety and efficacy in children and growing adolescents have not been established.

CONTRAINDICATIONS

- Patients who are hypersensitive to this drug or to any ingredient in the formulation. For a complete listing, see DOSAGE FORMS, COMPOSITION AND PACKAGING.
- Hypocalcemia (see WARNINGS AND PRECAUTIONS, General).

WARNINGS AND PRECAUTIONS

General

Hypocalcemia and other disturbances of bone and mineral metabolism should be effectively treated before starting ACTONEL (risedronate sodium) therapy.

Adequate intake of calcium and vitamin D is important in all patients, especially in patients with Paget’s disease in whom bone turnover is significantly elevated (see DRUG INTERACTIONS).

ACTONEL DR delayed release tablets are formulated to release in the small intestine to provide effective absorption of risedronate when taken as directed with breakfast. Other ACTONEL formulations should be taken on an empty stomach at least 30 minutes before first food of the day. For this reason, ACTONEL 35 mg should not be substituted for ACTONEL DR 35 mg.

Detailed dosing instructions (see DOSAGE AND ADMINISTRATION) are provided to ensure correct dosing of each ACTONEL therapy.

Osteonecrosis of the Jaw: In post-marketing reporting, osteonecrosis of the jaw has been reported in patients treated with bisphosphonates. The majority of reports occurred following dental procedures such as tooth extractions and have involved cancer patients treated with intravenous bisphosphonates, but some occurred in patients receiving oral treatment for postmenopausal osteoporosis and other diagnoses. Many had signs of local infection, including osteomyelitis. Osteonecrosis has other well documented multiple risk factors. It is not possible to determine if these events are related to bisphosphonates, to concomitant drugs or other therapies, to the patient’s underlying disease or to other co-morbid risk factors (e.g., anemia, infection, pre-existing oral disease). A dental examination with appropriate preventative dentistry should be considered prior to treatment with bisphosphonates in patients with concomitant risk factors (e.g., cancer, immunosuppression, head and neck radiotherapy or poor oral hygiene). While on treatment, these patients should avoid invasive dental procedures if possible. For patients requiring dental procedures, there are no data available to suggest whether discontinuation of bisphosphonate treatment prior to the procedure reduces the risk of osteonecrosis of the jaw. Clinical judgment, based on individual risk assessment, should guide the management of patients undergoing dental procedures.
Atypical Subtrochanteric and Diaphyseal Femoral Fractures: Atypical, low-energy, or low trauma fractures of the femoral shaft have been reported in bisphosphonate-treated patients. These fractures can occur anywhere in the femoral shaft from just below the lesser trochanter to above the supracondylar flare and are transverse or short oblique in orientation without evidence of comminution. Atypical femur fractures most commonly occur with minimal or no impact trauma to the affected area. They may be bilateral and many patients report prodromal pain in the affected area, usually presenting as dull, aching thigh pain, weeks to months before a complete fracture occurs. Poor healing of these fractures was also reported. Any patient with a history of bisphosphonate exposure who presents with thigh or groin pain should be suspected of having an atypical fracture and should be evaluated to rule out an incomplete femur fracture. Patients presenting with an atypical fracture should also be assessed for symptoms and signs of fracture in the contra-lateral limb. Interruption of bisphosphonate therapy should be considered, pending a risk/benefit assessment. Although causality has not been established, the role of bisphosphonates cannot be ruled out. 

Musculoskeletal: In postmarketing experience, severe and occasionally incapacitating bone, joint, and/or muscle pain has been reported in patients taking bisphosphonates (see ADVERSE REACTIONS). The time to onset of symptoms varied from one day to several months after starting the drug. Most patients had relief of symptoms after stopping the medication. A subset of patients had recurrence of symptoms when rechallenged with the same drug or another bisphosphonate. Consider discontinuing use if severe symptoms develop.

Gastrointestinal Bisphosphonates may cause upper gastrointestinal (GI) disorders such as dysphagia, esophagitis, esophageal ulcer, and gastric ulcer (see ADVERSE REACTIONS). Since some bisphosphonates have been associated with esophagitis and esophageal ulcerations, to facilitate delivery to the stomach and minimize the risk of these events, patients should take ACTONEL and ACTONEL DR while in an upright position (i.e., sitting or standing) and with sufficient plain water (≥ 120 mL). Patients should not lie down for at least 30 minutes after taking the drug. Health professionals should be particularly careful to emphasize the importance of the dosing instructions to patients with a history of esophageal disorders (e.g., inflammation, stricture, ulcer, or disorders of motility).

Ophthalmologic: Ocular disturbances including conjunctivitis, uveitis, episcleritis, iritis, and scleritis have been reported with ACTONEL therapy. Patients with ocular events other than uncomplicated conjunctivitis should be referred to an ophthalmologist for evaluation. If ocular inflammatory symptoms are observed, treatment may have to be discontinued.

Renal Risedronate sodium is not recommended for use in patients with severe renal impairment (creatinine clearance < 30 mL/min).

Special Populations Pediatrics: The safety and efficacy of risedronate sodium in children and growing adolescents have not been established.

Pregnant Women: Risedronate sodium is not intended for use during pregnancy. There are no studies of risedronate sodium in pregnant women.

Nursing Women: Risedronate sodium is not intended for use with nursing mothers. It is not known whether risedronate is excreted in human milk. Risedronate was detected in feeding pups exposed to lactating rats for a 24-hour period post-dosing, indicating a small degree of lacteal transfer. Since many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from bisphosphonates, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

ADVERSE REACTIONS

Adverse Drug Reaction Overview Bisphosphonates may cause upper gastrointestinal disorders such as dysphagia, esophagitis, esophageal ulcer and gastric ulcer. It is therefore important to follow the recommended dosing instructions (see DOSAGE AND ADMINISTRATION).

Musculoskeletal pain, rarely severe, has been reported as a common adverse event in patients who received ACTONEL and ACTONEL DR for all indications and dosage forms. In ACTONEL and ACTONEL DR osteoporosis studies, the most commonly reported adverse reactions were abdominal pain, dyspepsia and nausea. In addition, diarrhea was the most commonly reported adverse reaction for the highest ACTONEL monthly dose.

In Paget’s disease studies with ACTONEL, the most commonly reported adverse reactions were diarrhea, nausea, abdominal pain and headache.

Clinical Trial Adverse Drug Reactions Because clinical trials are conducted under very specific conditions the adverse reaction rates observed in the clinical trials may not reflect the rates observed in practice and should not be compared to the rates in the clinical trials of another drug. Adverse drug reaction information from clinical trials is useful for identifying drug-related adverse events and approximate rates of occurrence.

Treatment and Prevention of Postmenopausal Osteoporosis: ACTONEL 5 mg daily has been studied for up to 3 years in over 5000 women enrolled in Phase III clinical trials for treatment or prevention of postmenopausal osteoporosis. Most adverse events reported in these trials were either mild or moderate in severity, and did not lead to discontinuation from the study. The distribution of severe adverse events was similar across treatment groups. In addition, the overall incidence of adverse events (AEs) was found to be comparable amongst ACTONEL and placebo-treated patients.
Table 1 lists adverse events considered possibly or probably drug-related, reported in ≥ 1% of ACTONEL 5 mg daily-treated patients, in Phase III postmenopausal osteoporosis trials. Discontinuation of therapy due to serious clinical adverse events occurred in 5.5% of ACTONEL 5 mg daily-treated patients and 6.0% of patients treated with placebo.

<table>
<thead>
<tr>
<th>Drug-Related* Adverse Events Reported in ≥ 1% of ACTONEL 5 mg Daily-Treated Patients in Combined Phase III Postmenopausal Osteoporosis Trials</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Adverse Event</strong></td>
</tr>
<tr>
<td>Body as a Whole</td>
</tr>
<tr>
<td>Abdominal Pain</td>
</tr>
<tr>
<td>Headache</td>
</tr>
<tr>
<td>Asthenia</td>
</tr>
<tr>
<td>Digestive System</td>
</tr>
<tr>
<td>Dyspepsia</td>
</tr>
<tr>
<td>Nausea</td>
</tr>
<tr>
<td>Constipation</td>
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<td>Diarrhea</td>
</tr>
<tr>
<td>Flatulence</td>
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<td>Gastritis</td>
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<td>Skin and Appendages</td>
</tr>
<tr>
<td>Rash</td>
</tr>
<tr>
<td>Pruritus</td>
</tr>
</tbody>
</table>

*Considered to be possibly or probably causally related by clinical study Investigators.

**Weekly Dosing:** In the 1-year, double-blind, multicentre study comparing ACTONEL 35 mg Once-a-Week to ACTONEL 5 mg daily for the treatment of osteoporosis in postmenopausal women, the overall safety and tolerability profiles of the 2 oral dosing regimens were similar.

The proportion of patients who experienced an upper gastrointestinal adverse event and the pattern of those events were found to be similar between the ACTONEL 35 mg Once-a-Week and ACTONEL 5 mg daily-treated groups. In addition to the previously described adverse reactions reported in ACTONEL osteoporosis clinical trials, arthralgia (ACTONEL 35 mg, 2.1%; ACTONEL 5 mg, 1.3%) was reported in ≥ 1% of patients and in more ACTONEL 35 mg weekly treated patients than in ACTONEL 5 mg daily treated patients.

In the 1-year, double-blind, multicentre study comparing ACTONEL 35 mg Once-a-Week to placebo for the prevention of osteoporosis in postmenopausal women, the overall safety and tolerability profiles of the two groups were comparable with the exception of arthralgia. Specifically, 1.5% of patients taking ACTONEL 35 mg Once-a-Week experienced arthralgia compared to 0.7% of placebo patients. The overall safety profile observed in this study showed no substantive difference from that observed in the ACTONEL 5 mg daily versus ACTONEL 35 mg Once-a-Week treatment study.

**ACTONEL DR** – In a 2-year, double-blind, multicentre study comparing ACTONEL DR 35 mg weekly taken following breakfast to ACTONEL 5 mg daily for the treatment of osteoporosis in postmenopausal women, gastrointestinal adverse events were reported in 38.8% of patients taking ACTONEL DR 35 mg, compared to 34.9% of patients taking ACTONEL 5 mg. Abdominal pain, vomiting, and upper abdominal pain were reported more frequently by patients taking ACTONEL DR (6.2%, 4.9%, 3.6%) compared to patients taking ACTONEL 5 mg (3.3%, 3.3%, 2.6%). Other events reported more frequently by patients taking ACTONEL DR included diarrhea, constipation, nasopharyngitis, upper respiratory tract infection, and pharyngitis.

**Monthly Dosing: (Two Consecutive Days per Month)** – In a 1-year, double-blind, multicentre study for the treatment of osteoporosis in postmenopausal women comparing ACTONEL 75 mg on two consecutive days per month to ACTONEL 5 mg daily, the overall safety profiles of the dosing regimens were similar. The proportion of patients who experienced an upper gastrointestinal adverse event and the pattern of those events were found to be similar between the ACTONEL 75 mg two consecutive days per month and the ACTONEL 5 mg daily treated groups. In addition to the previously described adverse reactions, arthralgia (ACTONEL 75 mg, 1.5%; ACTONEL 5 mg, 1.0%), vomiting (ACTONEL 75 mg, 1.1%; ACTONEL 5 mg, 1.0%) and gastritis erosive (ACTONEL 75 mg, 1.0%; ACTONEL 5 mg, 0.3%) was reported in ≥ 1% of patients and in more ACTONEL 75 mg treated patients than in ACTONEL 5 mg daily treated patients.

Symptoms consistent with acute phase reactions have been reported. Based on reporting of any 33 acute phase reaction-like symptoms (without regard to causality) within the first 5 days of first dose, the overall incidence of acute phase reaction was 7.6% of patients on ACTONEL 75 mg two consecutive days per month and 3.6% of patients on ACTONEL 5 mg daily. Fever or influenza-like illness (without regard to causality) occurring within the first 5 days of first dose were reported by 0.6% of patients in the ACTONEL 75 mg two consecutive days per month and 0.0% in the ACTONEL 5 mg daily groups.

**(Once-a-Month)** – In a 1-year, double-blind, multicentre study for the treatment of osteoporosis in postmenopausal women comparing ACTONEL 150 mg Once-a-Month to ACTONEL 5 mg daily, the overall safety profiles of the dosing regimens were similar. The proportion of patients who experienced an upper gastrointestinal adverse event and the pattern of those events were found to be similar between the ACTONEL 150 mg Once-a-Month and the ACTONEL 5 mg daily treated groups. In addition to the previously described adverse reactions diarrhea (ACTONEL 150 mg, 3.1%; ACTONEL 5 mg, 0.5%), vomiting (ACTONEL 150 mg, 1.5%; ACTONEL 5 mg, 0.6%), arthralgia (ACTONEL 150 mg, 1.5%; ACTONEL 5 mg, 0.9%) and myalgia (ACTONEL 150 mg, 1.1%; ACTONEL 5 mg, 0.3%) were reported in ≥1% of patients and in more ACTONEL 150 mg treated patients than in ACTONEL 5 mg daily treated patients.

Symptoms consistent with acute phase reactions have been reported. Based on reporting of any 33 acute phase reaction-like symptoms (without regard to causality) within the first 3 days of first dose and lasting less than 7 days, the overall incidence of acute phase reaction was 5.2 % of patients in the ACTONEL 150 mg once-a-month group and 1.1% in the ACTONEL 5 mg daily group. Fever or influenza-like illness (without regard to causality) occurring within the first 3 days of first dose and lasting less than 7 days was reported by 1.4% of patients in the ACTONEL 150 mg Once-a-Month group and 0.2% of patients in the ACTONEL 5 mg daily group.

**Treatment of Osteoporosis in Men, to Improve Bone Mineral Density:** In a 2-year, double-blind, multicentre study using ACTONEL 35 mg Once-a-Week (n=191) and placebo (n=93) in men with osteoporosis, the overall safety and tolerability profiles of the two treatment groups were similar.

The proportion of patients who experienced an upper gastrointestinal adverse event and the pattern of those events were higher in placebo (18%) than in ACTONEL 35 mg Once-a-Week treated patients (8%).
In addition to the previously described adverse events, the following adverse events were reported in ≥2% of patients and in more ACTONEL-treated patients than placebo-treated patients in the male osteoporosis study (events are included without attribution of causality): hypoaesthesia (ACTONEL 35 mg, 2%; placebo, 1%), nephrolithiasis (ACTONEL 35 mg, 3%; placebo, 0%), benign prostatic hyperplasia (ACTONEL 35 mg, 5%; placebo, 3%) and arrhythmia (ACTONEL 35 mg, 2%; placebo, 0%).

Glucocorticoid-Induced Osteoporosis: ACTONEL 5 mg daily has been studied in two Phase III glucocorticoid-induced osteoporosis trials enrolling more than 500 patients. The adverse event profile of this population was similar to that seen in postmenopausal osteoporosis trials.

The overall incidence of adverse events was found to be comparable between the ACTONEL 5 mg daily and placebo treatment groups, with the exception of back and joint pain. Back pain was reported in 8.8% of placebo-treated patients and 17.8% of ACTONEL-treated patients; joint pain occurred in 14.7% of placebo patients and 24.7% of ACTONEL patients. Most adverse experiences reported were either mild or moderate in severity, and did not lead to discontinuation from the study. Discontinuation of therapy due to serious clinical adverse events occurred in 2.9% of ACTONEL 5 mg daily-treated patients and 5.3% of patients treated with placebo. The occurrence of adverse events does not appear to be related to patient age, gender or race.

Table 2 lists adverse events considered possibly or probably drug-related, reported in ≥1% of ACTONEL 5 mg daily-treated patients, in Phase III glucocorticoid-induced osteoporosis studies.

### Table 2

<table>
<thead>
<tr>
<th>Adverse Event</th>
<th>ACTONEL 5 mg N = 174 (%)</th>
<th>Placebo Control N = 170 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Body as a Whole</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal Pain</td>
<td>4.0</td>
<td>4.7</td>
</tr>
<tr>
<td>Headache</td>
<td>1.1</td>
<td>1.2</td>
</tr>
<tr>
<td><strong>Digestive System</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>5.7</td>
<td>2.9</td>
</tr>
<tr>
<td>Nausea</td>
<td>5.7</td>
<td>5.3</td>
</tr>
<tr>
<td>Constipation</td>
<td>2.9</td>
<td>3.5</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>2.9</td>
<td>3.5</td>
</tr>
<tr>
<td>Dry Mouth</td>
<td>1.1</td>
<td>0.6</td>
</tr>
<tr>
<td>Duodenitis</td>
<td>1.1</td>
<td>0.0</td>
</tr>
<tr>
<td>Esophagitis</td>
<td>1.1</td>
<td>0.0</td>
</tr>
<tr>
<td>Flatulence</td>
<td>1.1</td>
<td>1.8</td>
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<tr>
<td>Gastrointestinal Disorder</td>
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<tr>
<td><strong>Nervous System</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dizziness</td>
<td>1.1</td>
<td>1.2</td>
</tr>
<tr>
<td><strong>Skin and Appendages</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rash</td>
<td>1.1</td>
<td>2.4</td>
</tr>
<tr>
<td>Skin Disorder</td>
<td>1.1</td>
<td>0.0</td>
</tr>
</tbody>
</table>

*Considered to be possibly or probably causally related by clinical study Investigators.*

Endoscopic Findings: ACTONEL 5 mg daily clinical studies enrolled over 5700 patients for the treatment and prevention of postmenopausal and glucocorticoid-induced osteoporosis, many with pre-existing gastrointestinal disease and concomitant use of NSAIDs or ASA. Investigators were encouraged to perform endoscopies in any patients with moderate-to-severe gastrointestinal complaints while maintaining the blind. These endoscopies were ultimately performed on equal numbers of patients between the treated and placebo groups (75 ACTONEL; 75 placebo).

Across treatment groups, the percentage of patients with normal esophageal, gastric and duodenal mucosa on endoscopy was similar (21% ACTONEL; 20% placebo). Positive findings on endoscopy were also generally comparable across treatment groups. There were a higher number of reports of mild duodenitis in the ACTONEL group; however, there were more duodenal ulcers in the placebo group. Clinically important findings (perforations, ulcers or bleeding) among this symptomatic population were similar between groups (39% ACTONEL; 51% placebo).

At the 1-year time point in studies, comparing ACTONEL 35 mg Once-a-Week to ACTONEL 5 mg daily and ACTONEL DR 35 mg weekly to ACTONEL 5 mg daily in the treatment of postmenopausal osteoporosis, endoscopies performed during the studies revealed no dose dependent pattern in the number of patients with positive endoscopic findings or in the anatomical location of abnormalities detected. Endoscopies were conducted only on consenting patients experiencing moderate to severe gastrointestinal complaints.

In two, 1-year studies for the treatment of osteoporosis in postmenopausal women comparing ACTONEL 75 mg on two consecutive days per month and ACTONEL 150 mg Once-a-Month respectively to ACTONEL 5 mg daily, a similar percentage of patients for each of the intermittent regimens had at least one abnormal endoscopic finding when compared to the daily regimen (ACTONEL 75 mg, 3.2%; ACTONEL 5 mg, 3.1% and ACTONEL 150 mg, 3.4%; ACTONEL 5 mg, 4.2%).

Paget's Disease of Bone: ACTONEL has been studied in over 390 patients with Paget's disease of bone. The adverse experiences reported have usually been mild or moderate and generally have not required discontinuation of treatment. The occurrence of adverse experiences does not appear to be related to patient age, gender or race.

In a Phase III clinical study, ACTONEL and Didronel® (etidronate disodium tablets) showed similar adverse event profiles: 6.6% (4/61) of the patients treated with ACTONEL 30 mg daily for 2 months discontinued treatment due to adverse experiences, compared with 8.2% (5/61) of the patients treated with Didronel 400 mg daily for 6 months.
Table 3 lists adverse events considered possibly or probably drug-related, reported in ≥ 1% of ACTONEL 30 mg daily-treated patients, in the Phase III Paget’s trial.

<table>
<thead>
<tr>
<th>Adverse Event</th>
<th>ACTONEL 30 mg/day x 2 months N = 61 (%)</th>
<th>Didronel 400 mg/day x 6 months N = 61 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Body as a Whole</strong></td>
<td></td>
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</tr>
<tr>
<td>Abdominal Pain</td>
<td>6.6</td>
<td>3.3</td>
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<tr>
<td>Headache</td>
<td>4.9</td>
<td>6.6</td>
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<tr>
<td>Infection</td>
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<td>6.6</td>
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<td>Flu Syndrome</td>
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<tr>
<td>Neck Rigidity</td>
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<td>Neoplasm</td>
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<td>Pain</td>
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<td>Chest Pain</td>
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<td>Diarrhea</td>
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<tr>
<td>Nausea</td>
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<td>4.9</td>
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<td>Constipation</td>
<td>3.3</td>
<td>1.6</td>
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<tr>
<td>Flatulence</td>
<td>3.3</td>
<td>4.9</td>
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<td>Colitis</td>
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<td><strong>Metabolic and Nutritional</strong></td>
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<td>Peripheral Edema</td>
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<td>Hypocalcemia</td>
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<td>Weight Decreased</td>
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<tr>
<td><strong>Musculoskeletal System</strong></td>
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<tr>
<td>Arthralgia</td>
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<td>Leg Cramps</td>
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<td>Myasthenia</td>
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<tr>
<td>Bone Pain</td>
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<tr>
<td><strong>Nervous System</strong></td>
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<tr>
<td>Dizziness</td>
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<tr>
<td><strong>Respiratory System</strong></td>
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<tr>
<td>Apnea</td>
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<td>Bronchitis</td>
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<td>Sinusitis</td>
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<td><strong>Skin</strong></td>
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<td>Rash</td>
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<td><strong>Special Senses</strong></td>
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</table>

* Considered to be possibly or probably causally related by clinical study Investigators.

In the Phase III comparative study versus Didronel, patients with a history of upper GI disease or abnormalities were not excluded. Patients were also not excluded based on NSAID or ASA use. The proportion of ACTONEL 30 mg daily-treated patients with mild or moderate upper GI experiences was similar to that in the Didronel-treated group, with no severe upper GI experiences observed in either treatment group.

**Less Common Clinical Trial Adverse Drug Reactions**

The following adverse drug reactions were reported in ≥ 1% of patients who received ACTONEL for all indications:

- Uncommon (0.1-1.0%): duodenitis, iritis
- Rare (< 0.1%): abnormal liver function tests, glossitis

**Abnormal Hematologic and Clinical Chemistry Findings**

Asymptomatic mild decreases in serum calcium and phosphorus levels have been observed in some patients. Asymptomatic elevations in PTH levels were observed in some patients receiving ACTONEL DR (see ACTION AND CLINICAL PHARMACOLOGY, Pharmacodynamics).

Rare cases of leukemia have been reported following therapy with bisphosphonates. Any causal relationship to either the treatment or to the patients’ underlying disease has not been established.
**Post-Market Adverse Drug Reactions**

**Hypersensitivity and Skin Reactions:** Reported rarely, angioedema, generalized rash and bullous skin reactions, some severe.

**Musculoskeletal and Connective tissue:** Reported very rarely, low-energy femoral shaft fractures (see WARNINGS AND PRECAUTIONS)

**Osteonecrosis of the Jaw:** Osteonecrosis of the jaw has been reported rarely (see WARNINGS AND PRECAUTIONS).

**Ophthalmologic:** Reported rarely, conjunctivitis, episcleritis, iritis, scleritis and uveitis (see WARNINGS AND PRECAUTIONS).

**DRUG INTERACTIONS**

**Overview**

No specific drug-drug interaction studies were performed with risedronate sodium film-coated tablets. Animal studies have demonstrated that risedronate is highly concentrated in bone and is retained only minimally in soft tissue. No metabolites have been detected systemically or in bone. The binding of risedronate to plasma proteins in humans is low (24%), resulting in minimal potential for interference with the binding of other drugs. In an additional animal study, there was also no evidence of hepatic microsomal enzyme induction. In summary, risedronate sodium is not systemically metabolized, does not induce cytochrome P450 enzymes and has low protein binding.

Risedronate sodium is therefore not expected to interact with other drugs based on the effects of protein binding displacement, enzyme induction or metabolism of other drugs.

**Drug-Drug Interactions**

Patients in the clinical trials were exposed to a wide variety of commonly used concomitant medications (including NSAIDs, H2-blockers, proton pump inhibitors, antacids, calcium channel blockers, beta-blockers, thiazides, glucocorticoids, anticoagulants, anticonvulsants, cardiac glycosides) without evidence of clinically relevant interactions.

The drugs listed in Table 4 are based on either drug interaction case reports or predicted interactions due to the expected magnitude and seriousness of the interaction (i.e., those identified as contraindicated).

### Table 4

<table>
<thead>
<tr>
<th>Reference</th>
<th>Effect</th>
<th>Clinical Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antacids and calcium supplements which contain polyvalent cations (e.g., calcium, magnesium, aluminum and iron)</strong></td>
<td>CT/T</td>
<td>Interference with the absorption of ACTONEL and ACTONEL DR. Co-administration of ACTONEL DR with calcium supplement after breakfast reduced bioavailability of ACTONEL DR by approximately 38%.</td>
</tr>
<tr>
<td><strong>Hormone replacement therapy (HRT)</strong></td>
<td>CT</td>
<td>No clinically significant effect for ACTONEL.</td>
</tr>
<tr>
<td><strong>H2-blockers and proton pump inhibitors (PPIs)</strong></td>
<td>CT</td>
<td>Among H2-blockers and PPIs users, the incidence of upper gastrointestinal adverse events was similar between the ACTONEL-treated patients and placebo-treated patients.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Among H2-blockers and PPIs users, the incidence of upper gastrointestinal adverse experiences was found to be similar between the weekly- and daily-treated groups.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Drugs that raise stomach pH (e.g. H2 blockers, PPIs, or antacids) may affect the enteric coating on ACTONEL DR tablets and thereby reduce bioavailability of ACTONEL DR. The effects of concomitant administration of H2 blockers, PPIs, and antacids on bioavailability of ACTONEL DR have not been evaluated.</td>
</tr>
</tbody>
</table>

**CT:** Clinical Trial; **T:** Theoretical

---

Of over 5700 patients enrolled in the ACTONEL 5 mg daily Phase III osteoporosis studies, ASA use was reported by 31% of patients and NSAID use by 48%. Among these ASA or NSAID users, the incidence of upper gastrointestinal adverse events was similar between the ACTONEL-treated patients and placebo-treated patients.

In the 1-year study comparing ACTONEL 35 mg Once-a-Week to ACTONEL 5 mg daily, ASA use was reported by 56% and NSAID use by 41%. The incidence of upper gastrointestinal adverse events was similar between the ACTONEL- and daily-treated groups.

In the Phase 3 study comparing ACTONEL DR 35 mg weekly immediately following breakfast and ACTONEL 5 mg daily, 22% of NSAID/ASA users in both groups developed upper gastrointestinal adverse reactions. Among non-users, 16% of patients taking ACTONEL DR 35 mg weekly immediately following breakfast developed upper gastrointestinal adverse reactions, compared to 13% taking ACTONEL 5 mg daily.

In two, 1-year studies comparing ACTONEL 75 mg two consecutive days per month or ACTONEL 150 mg once-a-month to ACTONEL 5 mg daily in postmenopausal women, 55% (75 mg) and 46% (150 mg) of patients reported the use of ASA and/or NSAIDs. Among these ASA or NSAID users, the incidence of upper gastrointestinal adverse events was similar in the ACTONEL monthly-treated groups when compared to the daily-treated groups respectively.
Drug-Food Interactions
Clinical benefits may be compromised by failure to take ACTONEL on an empty stomach.

ACTONEL DR should be taken with food. When compared with ACTONEL 5 mg, treatment with ACTONEL DR resulted in a higher incidence of upper abdominal pain when administered before breakfast under fasting conditions. For dosing information see DOSAGE AND ADMINISTRATION.

Drug-Herb Interactions
Interactions with herbs have not been studied.

Drug-Laboratory Interactions
Bisphosphonates are known to interfere with the use of bone-imaging agents. Specific studies with ACTONEL and ACTONEL DR have not been performed.

DOSAGE AND ADMINISTRATION

Dosing Considerations
Patients should receive supplemental calcium and vitamin D if dietary intake is inadequate (see WARNINGS AND PRECAUTIONS, General).

ACTONEL (risedronate sodium) film-coated tablets
- ACTONEL should be taken on an empty stomach at least 30 minutes before consuming the first food, drink (other than plain water) and/or any other medication of the day. Food, medication or drink other than plain water can interfere with the absorption of ACTONEL. (See Recommended Dose and Dosage Adjustment).
- Each ACTONEL tablet should be swallowed whole while the patient is in an upright position and with sufficient plain water (≥ 120 mL) to facilitate delivery to the stomach. ACTONEL tablets should not be chewed, cut, or crushed (see WARNINGS AND PRECAUTIONS, General).
- Patients taking ACTONEL should not lie down for at least 30 minutes after taking the medication (see WARNINGS AND PRECAUTIONS, General).
- Medications containing polyvalent cations (e.g. calcium, magnesium, aluminum, and iron) can interfere with the absorption of ACTONEL. These medications should be administered at a different time of the day than ACTONEL.

ACTONEL DR (risedronate sodium) delayed-release tablets
- ACTONEL DR should be taken in the morning, with breakfast, (this may include high fat foods, coffee, tea, milk, orange juice, etc.) (See Recommended Dose and Dosage Adjustment section). A higher incidence of upper abdominal pain was seen when ACTONEL DR was taken in a fasted state before breakfast (see WARNINGS AND PRECAUTIONS – Drug-Food interactions).
- Each ACTONEL DR tablet should be swallowed whole while the patient is in an upright position and with sufficient plain water (≥ 120 mL) to facilitate delivery to the stomach.
- Patients taking ACTONEL DR should not lie down for at least 30 minutes after taking the medication (see WARNINGS AND PRECAUTIONS, General).
- ACTONEL DR tablets should not be chewed, cut, or crushed. Care should be taken not to break the outer coating which is designed to remain intact until the tablet reaches the small intestine where the tablet coating dissolves and releases the active ingredient (see WARNINGS AND PRECAUTIONS, General).
- Calcium supplements and antacids can interfere with the absorption of ACTONEL DR. These medications should be administered at a different time of the day than ACTONEL DR.

Recommended Dose and Dosage Adjustment
For all indications and doses: The patient should be informed to pay particular attention to the dosing instructions as clinical benefits may be compromised by failure to take the drug according to instructions.

Treatment of Postmenopausal Osteoporosis: The recommended regimens are daily (5 mg), weekly (35 mg Once-a-Week film-coated and delayed-release tablets), monthly duet (75 mg on two consecutive days per month, on the same calendar days each month) or monthly (1 tablet of 150 mg once-a-month on the same calendar day each month), taken orally.

Prevention of Postmenopausal Osteoporosis: The recommended regimens are daily (5 mg) or weekly (35 mg Once-a-Week film-coated tablets), taken orally.

Treatment of Osteoporosis in Men, to Improve Bone Mineral Density: The recommended regimen is 35 mg Once-a-Week film-coated tablets, taken orally.

Treatment and Prevention of Glucocorticoid-Induced Osteoporosis: The recommended regimen is 5 mg daily, taken orally.

Treatment of Paget's Disease of Bone: The recommended regimen is 30 mg daily for 2 months, taken orally. Re-treatment may be considered (following post-treatment observation of at least 2 months) if relapse has occurred, or if treatment fails to normalize serum alkaline phosphatase. For re-treatment, the dose and duration of therapy are the same as for initial treatment. There are no data available on more than one course of re-treatment.

Renal Impairment: No dosage adjustment is necessary in patients with a creatinine clearance ≥ 30 mL/min or in the elderly. Not recommended for use in patients with severe renal impairment (creatinine clearance < 30 mL/min).

Geriatrics: No dosage adjustment is necessary in elderly patients (see INDICATIONS AND CLINICAL USE, Geriatrics).

Missed Dose
Daily: Patients should be instructed that if they miss a dose of ACTONEL 5 mg or 30 mg, they should take 1 tablet of ACTONEL as they normally would for their next dose. Patients should not double their next dose or take 2 tablets on the same day.

Weekly: Patients should be instructed that if they miss a dose of ACTONEL or ACTONEL DR 35 mg Once-a-Week on their regularly scheduled day, they should take 1 tablet on the day they first remember missing their dose. Patients should then return to taking 1 tablet once a week as originally scheduled on their chosen day. Patients should not take 2 tablets on the same day.

Monthly Duet: If one or both tablets of ACTONEL 75 mg monthly duet are missed, and the next month’s scheduled doses are more than 7 days away, the patient should be instructed as follows:
- If both tablets are missed, take 1 ACTONEL 75 mg tablet in the morning after the day it is remembered and then the other tablet on the next consecutive morning.
- If only 1 ACTONEL 75 mg tablet is missed, take the missed tablet in the morning after the day it is remembered.
Patients should then return to taking their ACTONEL 75 mg monthly duet on two consecutive days each month as originally scheduled. Patients should not take more than two 75 mg tablets within 7 days. If one or both tablets of ACTONEL 75 mg are missed, and the next month’s scheduled doses are within 7 days, patients should wait until their next month’s scheduled doses and then continue taking ACTONEL 75 mg monthly duet on two consecutive days each month as originally scheduled.

**Once-a-Month:** Patients should be instructed that if they miss a 150 mg dose of ACTONEL (1 tablet of 150 mg), and the next month’s scheduled dose is more than 7 days away, they should take the missed tablet in the morning after the day it is remembered. Patients should then return to taking their ACTONEL 150 mg as originally scheduled.

If a dose of ACTONEL 150 mg is missed, and the next month’s scheduled dose is within 7 days, patients should be instructed to wait until their next month’s scheduled dose and then continue taking ACTONEL 150 mg. Patients should not take more than 150 mg of ACTONEL within 7 days.

**OVERDOSAGE**

Decreases in serum calcium following substantial overdose may be expected in some patients. Signs and symptoms of hypocalcemia may also occur in some of these patients.

Milk or antacids containing calcium, magnesium, and aluminum may be given to bind ACTONEL (film-coated tablets) and reduce absorption of the drug; the impact of this intervention for ACTONEL DR (delayed-release tablets) has not been evaluated. The ACTONEL DR formulation is less sensitive to the binding effects of divalent cations. In cases of substantial overdose, gastric lavage may be considered to remove unabsorbed drug if performed within 30 minutes of ingestion. Standard procedures that are effective for treating hypocalcemia, including the administration of calcium intravenously, would be expected to restore physiologic amounts of ionized calcium and to relieve signs and symptoms of hypocalcemia.

For management of a suspected drug overdose, contact your regional Poison Control Centre.

**ACTION AND CLINICAL PHARMACOLOGY**

**Mechanism of Action**

Risedronate sodium, a pyridinyl-bisphosphonate in the form of hemi-pentahydrate with small amounts of monohydrate, inhibits osteoclast bone resorption and modulates bone metabolism. Risedronate has a high affinity for hydroxyapatite crystals in bone and is a potent antiresorptive agent. At the cellular level, risedronate inhibits osteoclasts. The osteoclasts adhere normally to the bone surface, but show evidence of reduced active resorption (e.g., lack of ruffled border). Histomorphometry in rats, dogs, minipigs and humans showed that risedronate treatment reduces bone turnover (i.e., activation frequency, the rate at which bone remodelling sites are activated) and bone resorption at remodelling sites.

**Pharmacodynamics**

**Treatment and Prevention of Osteoporosis in Postmenopausal Women:** Osteoporosis is a degenerative and debilitating bone disease characterized by decreased bone mass and increased fracture risk at the spine, hip, and wrist. The diagnosis can be confirmed by the finding of low bone mass, evidence of fracture on x-ray, a history of osteoporotic fracture, or height loss or kyphosis indicative of vertebral fracture. Osteoporosis occurs in both men and women but is more common among women following menopause.

In healthy humans, bone formation and resorption are closely linked; old bone is resorbed and replaced by newly-formed bone. In postmenopausal osteoporosis, bone resorption exceeds bone formation, leading to bone loss and increased risk of bone fracture. After menopause, the risk of fractures of the spine and hip increases dramatically; approximately 40% of 50-year-old women will experience an osteoporosis-related fracture of the spine, hip, or wrist during their remaining lifetimes. After experiencing one osteoporosis-related fracture, the risk of future fracture increases 5-fold compared to the risk among a non-fractured population. One in five men older than 50 years will have an osteoporotic fracture, most commonly at the spine, hip and wrist.

Risedronate sodium treatment decreases the elevated rate of bone turnover and corrects the imbalance of bone resorption relative to bone formation that is typically seen in postmenopausal osteoporosis. In clinical trials, administration of ACTONEL to postmenopausal women resulted in dose-dependent decreases in biochemical markers of bone turnover, including urinary markers of bone resorption and serum markers of bone formation, at doses as low as 2.5 mg daily. At the 5 mg daily dose, decreases in resorption markers were evident within 14 days of treatment. Changes in bone formation markers were observed later than changes in resorption markers, as expected, due to the coupled nature of bone formation and bone resorption; decreases in bone formation of about 20% were evident within 3 months of treatment. Bone turnover markers (BTMs) reached a nadir of about 40% below baseline values by the sixth month of treatment and remained stable with continued treatment for up to 3 years.

These data demonstrate that ACTONEL 5 mg administered daily to postmenopausal women produces a rapid reduction in bone resorption without over-suppression of bone formation. Bone turnover is decreased as early as 2 weeks and maximally within about 6 months of treatment, with achievement of a new steady-state which more nearly approximates the rate of bone turnover seen in premenopausal women.

In weekly and monthly ACTONEL postmenopausal osteoporosis dosing studies, consistent decreases in bone resorption (50-60%) and bone formation (30-40%) markers were observed at Month 12. Similarly, in a 2-year study for the treatment of osteoporosis in postmenopausal women comparing ACTONEL DR 35 mg weekly to baseline, consistent decreases in bone resorption (47-50%, 49-54%) and bone formation (33-34%, 35-37%) markers were observed at Month 12 and Month 24, respectively.

As a result of the inhibition of bone resorption, asymptomatic and usually transient decreases from baseline in serum calcium (about 2%) and serum phosphate levels (about 5%) and compensatory increases in serum parathyroid hormone (PTH) levels were observed within 6 months in ACTONEL 5 mg daily-treated patients in postmenopausal osteoporosis trials. No further decreases in serum calcium or phosphate, or increases in PTH were observed in postmenopausal women treated for up to 3 years.

In two 1-year studies for the treatment of osteoporosis in postmenopausal women comparing ACTONEL 35 mg Once-a-Week and ACTONEL 150 mg Once-a-Month respectively to ACTONEL 5 mg daily, similar mean changes from baseline in serum calcium, phosphate and PTH were found for each of the intermittent regimens when compared to the daily dosage regimen. In the 1-year study comparing ACTONEL 75 mg on two consecutive days per month to ACTONEL 5 mg daily, the mean percent changes from baseline were for serum calcium (0.8% and 0.2%), phosphate (-1.1% and -1.9%) and PTH (-11.7% and -3.0%), respectively.

In a 2-year study for the treatment of osteoporosis in postmenopausal women comparing ACTONEL DR 35 mg weekly to ACTONEL 5 mg daily, similar mean percent changes from baseline to 2 years were found between the 2 oral dosing regimens in serum calcium and phosphate. The effect of ACTONEL DR 35 mg weekly and ACTONEL 5 mg daily on PTH was evaluated in postmenopausal women with osteoporosis. At 2 years, in subjects with normal levels at baseline, PTH levels greater than 65 ng/L (upper limit of normal) were noted in 12% of subjects receiving ACTONEL DR 35 mg weekly immediately following breakfast and 6% of subjects receiving ACTONEL 5 mg daily. In subjects with normal levels at baseline, PTH levels greater than 97 ng/L (1.5 times the upper limit of normal) at 2 years were seen in 3% of subjects receiving ACTONEL DR 35 mg weekly immediately following breakfast and 0 subjects receiving ACTONEL 5 mg daily. There were no clinically significant differences between treatment groups for levels of calcium, phosphorus and magnesium.

Consistent with the effects of ACTONEL on biochemical markers of bone turnover, daily oral doses as low as 2.5 mg produced dose dependent, significant increases in lumbar spine bone mineral density (BMD) (ACTONEL 2.5 mg, 3% to 3.7%; ACTONEL 5 mg, 4% to 4.5%) after 12 months of treatment in large-scale postmenopausal osteoporosis trials.
A dose-dependent response to treatment was also observed in the BMD of the femoral neck over the same time (ACTONEL 2.5 mg, 0.7% to 0.9%; ACTONEL 5 mg, 1.5% to 2%). In three 1-year weekly and monthly dosing studies for the treatment of osteoporosis in postmenopausal women, comparing ACTONEL 35 mg Once-a-Week, ACTONEL 75 mg on two consecutive days per month and ACTONEL 150 mg Once-a-Month respectively to ACTONEL 5 mg daily, similar mean changes from baseline in BMD of the lumbar spine, total proximal femur, femoral neck and femoral trochanter were found for each of the intermittent regimens when compared to the daily regimen. In the two year study of ACTONEL DR 35 mg, it was shown that at 1 year and 2 years, ACTONEL DR 35 mg weekly was non-inferior to the ACTONEL 5 mg daily regimen for the primary efficacy variable of percent change from baseline of lumbar spine BMD. The two treatment groups were also similar with regard to percent change from baseline BMD at the total proximal femur, greater trochanter and femoral neck. Non-inferiority was observed with ACTONEL DR relative to ACTONEL 5 mg. At 2 years, the mean percent change from baseline in lumbar spine BMD was 4.1% for ACTONEL 5 mg and 5.2% for the ACTONEL DR 35 mg (upper limit CI = -0.355%). (see CLINICAL TRIALS, Treatment of Osteoporosis in Postmenopausal Women).

The ACTONEL DR tablet has an enteric coating, which delays the formulation of risedronate until the small intestine. The other formulations of ACTONEL are film coated.

**Treatment of Osteoporosis in Men, to Improve Bone Mineral Density:** In a 2-year clinical trial in the treatment of osteoporosis in men, ACTONEL 35 mg Once-a-Week decreased urinary collagen cross-linked N-telopeptide (NTX) (a marker of bone resorption), and serum bone specific alkaline phosphatase (BAP) (a marker of bone formation) by approximately 40% and 30%, below baseline values, respectively, within 12 months. The BTMs all had statistically significant decreases in bone turnover from baseline compared to placebo at all time points. The decreases in bone turnover were observed within 3 months after initiation of therapy and maintained throughout the 2-year study.

**Glucocorticoid-Induced Osteoporosis:** Chronic exposure to glucocorticoids (≥ 7.5 mg/day prednisone or its equivalent) induces rapid bone loss by decreasing bone formation and increasing bone resorption. The bone loss occurs most rapidly during the first 7 months of therapy with persistent but slowing bone loss for as long as glucocorticoid therapy continues.

Glucocorticoid-induced osteoporosis is characterized by low bone mass that leads to an increased risk of fracture (especially vertebral, hip and rib). It occurs in both men and women, and approximately 50% of patients on chronic glucocorticoid treatment will experience fractures. The relative risk of a hip fracture in patients on > 7.5 mg/day prednisone is more than doubled (RR = 2.27); the relative risk of vertebral fracture is increased five-fold (RR = 5.18).

ACTONEL treatment decreases bone resorption without directly inhibiting bone formation. In 1-year clinical trials in the treatment and prevention of glucocorticoid-induced osteoporosis, ACTONEL 5 mg daily produced rapid and statistically significant reductions in biochemical markers of bone turnover, similar to those seen in postmenopausal osteoporosis. Urinary collagen cross-linked N-telopeptide (a marker of bone resorption) and serum bone specific alkaline phosphatase (a marker of bone formation) were decreased by 50% to 55% and 25% to 30%, respectively, within 3 to 6 months after initiation of therapy. The reduction was evident within 14 days and BTMs remained decreased throughout the duration of ACTONEL treatment.

Consistent with the changes in biochemical markers of bone turnover, ACTONEL 5 mg daily provides a beneficial effect on bone mineral density and reduces the risk of vertebral fractures by approximately 70% when compared to placebo (see CLINICAL TRIALS, Glucocorticoid-Induced Osteoporosis).

**Paget’s Disease of Bone:** Paget’s disease of bone is a chronic focal skeletal disorder characterized by greatly increased and disordered bone remodelling. Excessive osteoclastic bone resorption is followed by osteoblastic new bone formation, leading to the replacement of the normal bone architecture by disorganized, enlarged and weakened bone structure.

Clinical manifestations of Paget’s disease range from no symptoms to severe morbidity due to bone pain, bone deformity, pathological fractures, and neurological and other complications. Serum alkaline phosphatase, the most frequently used biochemical marker of disease activity, provides an objective measure of disease severity and response to therapy.

ACTONEL is a bisphosphonate that acts primarily to inhibit bone resorption. This effect is related to its inhibitory effect on osteoclasts. In the Phase III clinical trial, ACTONEL 30 mg daily for 2 months produced significant (p < 0.001) reductions of 81% to 88% in serum alkaline phosphatase excess, as well as significant reductions in bone-specific serum alkaline phosphatase (Ostase, 67% to 70%) and urinary deoxypyridinoline/creatinine (47% to 51%). Reductions were evident as early as 1 month after the start of treatment, and progressively increased in magnitude (following completion of the 2 month treatment) when measured at monthly intervals over a 6 month period. Clinically meaningful reductions in serum alkaline phosphatase were observed starting at 1 month with levels maintained through 12 months.

Asymptomatic and mild decreases in serum calcium and phosphorus levels have been observed in some patients. These decreases in calcium are associated with increases in serum intact PTH and 1,25-dihydroxy vitamin D, resulting in an increase in tubular reabsorption of calcium.

Markers of bone resorption (such as urinary deoxypyridinoline/creatinine or hydroxyproline/creatinine) usually decrease before markers of bone formation (such as serum alkaline phosphatase). This difference is indicative of the primary antiresorptive effect of ACTONEL.

Bone turnover marker levels continue to decrease when ACTONEL treatment is stopped. Therefore, to assess the full effect of response, patients should be followed for at least 2 HI months following the 2 month treatment period.

### Pharmacokinetics

<table>
<thead>
<tr>
<th>Dosage Form</th>
<th>t&lt;sub&gt;max&lt;/sub&gt; (hours)</th>
<th>t&lt;sub&gt;1/2&lt;/sub&gt; (hours)</th>
<th>AUC&lt;sub&gt;0-&lt;infty&gt;&lt;/sub&gt; (ng·h·mL&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>Clearance (L/h/kg)</th>
<th>V&lt;sub&gt;z&lt;/sub&gt; (L/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 mg tablet; single dose</td>
<td>0.93&lt;sup&gt;a&lt;/sup&gt;</td>
<td>206.1</td>
<td>3.45</td>
<td>19.94</td>
<td>5542</td>
</tr>
<tr>
<td>30 mg tablet; single dose</td>
<td>4.2</td>
<td>226.1</td>
<td>17.1</td>
<td>23.60</td>
<td>7542</td>
</tr>
<tr>
<td>35 mg tablet; multiple dose, steady state</td>
<td>10.6</td>
<td>0.49</td>
<td>nd</td>
<td>53.3</td>
<td>nd</td>
</tr>
<tr>
<td>35 mg DR tablet; single dose</td>
<td>14.1</td>
<td>3.0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>nd</td>
<td>34.2&lt;sup&gt;b&lt;/sup&gt;</td>
<td>nd</td>
</tr>
<tr>
<td>75 mg tablet; multiple dose, steady state</td>
<td>19.3&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.66&lt;sup&gt;d&lt;/sup&gt;</td>
<td>299.7&lt;sup&gt;d&lt;/sup&gt;</td>
<td>180.7&lt;sup&gt;d&lt;/sup&gt;</td>
<td>14.8&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>150 mg tablet; single dose</td>
<td>74.6&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0.66&lt;sup&gt;e&lt;/sup&gt;</td>
<td>349.6&lt;sup&gt;e&lt;/sup&gt;</td>
<td>332.4&lt;sup&gt;e&lt;/sup&gt;</td>
<td>6.94&lt;sup&gt;e&lt;/sup&gt;</td>
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</table>

<sup>a</sup> arithmetic mean; <sup>b</sup> administered weekly; <sup>c</sup> administered on two consecutive days per month (150 mg total monthly dose); <sup>d</sup> geometric mean; <sup>e</sup> t<sub>1/2</sub>, z: is the half-life of the terminal exponential phase; V<sub>z</sub>: is the terminal volume of distribution uncorrected for bioavailability; nd: not determined; * AUClast.

**Absorption:** Absorption after an oral dose is relatively rapid (t<sub>max</sub> ~ 1 hour) for the film-coated tablet and occurs throughout the upper gastrointestinal tract. Absorption is independent of dose up to 75 mg two consecutive days per month; systemic exposure increases disproportionally at 150 mg (about 2 fold greater than expected based on dose). Steady-state conditions in the serum are observed within 57 days of dosing. The mean oral bioavailability of the 30 mg film-coated tablet is 0.63% and is bioequivalent to a solution. Extent of absorption when administered 30 minutes before breakfast is reduced by 55% compared to dosing in the fasting state (i.e., no food or drink for 10 hours prior to...
or 4 hours after dosing). Dosing 1 hour prior to breakfast reduces extent of absorption by 30% compared to dosing in the fasting state. Dosing either 30 minutes prior to breakfast or 2 hours after a meal results in a similar extent of absorption.

ACTONEL DR (risedronate sodium) 35 mg delayed-release tablet achieved a peak serum concentration at approximately 3 hours. Urinary excretion data showed that the fraction of the dose absorbed from ACTONEL DR is independent of the dose over the range studied (single dose, from 20 mg to 100 mg).

In a crossover pharmacokinetic study that evaluated food effect, the bioavailability of ACTONEL DR 35 mg delayed-release tablets decreased by ~30% when administered immediately after a high-fat breakfast compared to administration 4 hours before a meal. The bioavailability of the 35 mg ACTONEL DR tablet administered after a high fat breakfast was ~2 to 4-fold greater than the 35 mg risedronate film-coated tablet administered 30 minutes prior to a high-fat breakfast. Across different studies, the bioavailability of ACTONEL DR was not affected by breakfast meals with varying amount of fat and calories.

In a separate study, ACTONEL DR administered after dinner exhibited approximately 87% increase in exposure compared to administration following a breakfast. The safety and efficacy of dosing ACTONEL DR after dinner has not been evaluated (see DOSAGE AND ADMINISTRATION).

**Distribution:** The mean steady-state volume of distribution is 6.3 L/kg in humans. Human plasma protein binding of drug is about 24%. Preclinical studies in rats and dogs showed intravenous with single doses of [14C] risedronate indicate that approximately 60% of the dose is distributed to bone. The remainder of the dose is excreted in the urine. After multiple oral dosing in rats, the uptake of risedronate in soft tissues was found to be minimal (in the range of 0.001% to 0.01%), with drug levels quickly decreasing after the final dose.

**Metabolism:** There is no evidence that risedronate is systemically metabolized.

**Excretion:** Approximately half of the absorbed dose is excreted in urine within 24 hours, and 85% of an intravenous dose is recovered in the urine over 28 days. The mean renal clearance is 105 mL/min (CV = 34%) and mean total clearance is 122 mL/min (CV = 19%), with the difference primarily reflecting non-renal clearance or clearance due to adsorption to bone. The renal clearance is not concentration dependent, and there is a linear relationship between renal clearance and creatinine clearance. Unabsorbed drug is eliminated unchanged in feces. Once risedronate is absorbed, the serum concentration-time profile is multi-phasic with an initial half-life of about 1.5 hours and a terminal exponential half-life of 480 hours. Although the elimination rate of bisphosphonates from human bone is unknown, the 480 hour half-life is hypothesized to represent the dissociation of risedronate from the surface of bone.

**Special Populations and Conditions**

**Pediatrics:** Risedronate pharmacokinetics have not been studied in patients < 18 years of age.

**Geriatrics:** Bioavailability and disposition are similar in elderly (> 65 years of age) and younger subjects. No dosage adjustment is necessary.

**Gender:** Bioavailability and disposition following oral administration are similar in men and women.

**Race:** Pharmacokinetic differences due to race have not been studied.

**Hepatic Insufficiency:** No studies have been performed to assess risedronate’s safety or efficacy in patients with hepatic impairment. Risedronate is not metabolized in rat, dog, and human liver preparations. Insignificant amounts (< 0.1% of intravenous dose) of drug are excreted in the bile in rats. Therefore, dosage adjustment is unlikely to be needed in patients with hepatic impairment.

**Renal Insufficiency:** Risedronate is excreted intact primarily via the kidney. Patients with mild-to-moderate renal impairment (creatinine clearance > 30 mL/min) do not require a dosage adjustment. Exposure to risedronate was estimated to increase by 44% in patients with creatinine clearance of 20 mL/min. ACTONEL and ACTONEL DR are not recommended for use in patients with severe renal impairment (creatinine clearance < 30 mL/min) because of a lack of clinical experience.

**Genetic Polymorphism:** No data are available.

**STORAGE AND STABILITY**

Store at controlled room temperature 20°C - 25°C.

**DOSAGE FORMS, COMPOSITION AND PACKAGING**

**ACTONEL**

**Medicinal Ingredients:** Each risedronate sodium tablet for oral administration contains the equivalent of 5, 30, 35, 75 or 150 mg of anhydrous risedronate sodium in the form of the hemi-pentahydrate with small amounts of monohydrate.

**Nonmedicinal Ingredients (Film-coated Tablets):** Crospovidone, ferric oxide red (35 and 75 mg), ferric oxide yellow (5 and 35 mg), hydroxypropyl cellulose, hypromellose, indigo carmine (150 mg), lactose monohydrate (5, 30 and 35 mg), magnesium stearate, microcrystalline cellulose, polyethylene glycol, silicon dioxide and titanium dioxide.

<table>
<thead>
<tr>
<th>Dosage Strength</th>
<th>Description</th>
<th>Packaging</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 mg</td>
<td>film-coated, oval-shaped, yellow tablets with “RSN” engraved on one face and “5 mg” engraved on the other</td>
<td>carton of 28 blister packaged tablets</td>
</tr>
<tr>
<td>30 mg</td>
<td>film-coated, oval-shaped, white tablets with “RSN” engraved on one face and “30 mg” engraved on the other</td>
<td>bottle of 30 tablets</td>
</tr>
<tr>
<td>35 mg</td>
<td>film-coated, oval-shaped, orange tablets with “RSN” engraved on one face and “35 mg” engraved on the other</td>
<td>carton of 4 blister packaged tablets</td>
</tr>
<tr>
<td>75 mg</td>
<td>film-coated, oval-shaped, pink tablets with “RSN” engraved on one face and “75 mg” engraved on the other</td>
<td>carton of 2 blister packaged tablets</td>
</tr>
<tr>
<td>150 mg</td>
<td>film-coated, oval-shaped, blue tablets with “RSN” engraved on one face and “150 mg” engraved on the other</td>
<td>carton of 1 blister packaged tablet</td>
</tr>
</tbody>
</table>
ACTONEL DR

Medicinal Ingredients: Each risedronate sodium tablet for oral administration contains the equivalent 35 mg of anhydrous risedronate sodium in the form of the hemi-pentahydrate with small amounts of monohydrate.

Nonmedicinal Ingredients (Delayed-release tablets): Edetate disodium, ferric oxide yellow, magnesium stearate, methacrylic acid copolymer dispersion, silicified microcrystalline cellulose, polysorbate 80, simethicone, sodium starch glycolate, stearic acid, talc, and triethyl citrate.

<table>
<thead>
<tr>
<th>Dosage Strength</th>
<th>Description</th>
<th>Packaging</th>
</tr>
</thead>
<tbody>
<tr>
<td>35 mg</td>
<td>delayed-release, oval-shaped, yellow tablets with “EC 35” engraved on one face</td>
<td>carton of 4 blister packaged tablets</td>
</tr>
</tbody>
</table>

The ACTONEL DR tablet has an enteric coating, which delays the release of risedronate until the small intestine. The other formulations of ACTONEL are film coated, and must be taken before the first food of the day.

PART II: SCIENTIFIC INFORMATION

PHARMACEUTICAL INFORMATION

Drug Substance

- **Proper Name:** risedronate sodium hemi-pentahydrate
- **Chemical Name:** Phosphonic acid, [1-hydroxy-2-(3-pyridinyl)ethylidene]bis-, monosodium salt.
- **Molecular Formula:** C,H,NO,P:Na·2.5H2O
- **Structural Formula:**

![Structural Formula](image)

- **Molecular Weight:** Anhydrous: 305.10
  Hemi-pentahydrate: 350.13

- **Solubility:** Risedronate sodium is soluble in pH 7.0 potassium phosphate dibasic solution, 0.1 N sodium hydroxide, and water; very slightly soluble in 0.1 N hydrochloric acid, practically insoluble in ethanol, and insoluble in isopropanol.

- **Solution pH:** The pH of a 1.0% aqueous solution of risedronate sodium is 4.15.

- **Dissociation Constants:** The five pKₐ values for risedronate sodium are as follows:
  pK₁ = 1.6 ± 0.2, pK₂ = 2.2 ± 0.2, pK₃ = 5.9 ± 0.1, pK₄ = 7.1 ± 0.1 and pK₅ = 11.7 ± 0.3.

- **Description:** Risedronate sodium is a fine white to off-white crystalline powder. Risedronate sodium is present in the form of hemi-pentahydrate with small amounts of monohydrate.
## CLINICAL TRIALS
### Treatment of Osteoporosis in Postmenopausal Women

#### Study Demographics and Trial Design

<table>
<thead>
<tr>
<th>Study Number</th>
<th>Trial Designa</th>
<th>Dosage</th>
<th>Duration</th>
<th>Patients N = number</th>
<th>Age Range (Age Mean)</th>
<th>Daily Supplement**</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>R, PC, DB, MC, PG</td>
<td>2.5 mg/day 5 mg/day Placebo</td>
<td>2 years 3 years 3 years</td>
<td>1226</td>
<td>48-85 (71.0)</td>
<td>≤500 IU</td>
</tr>
<tr>
<td>2</td>
<td>R, PC, DB, MC, PG</td>
<td>2.5 mg/day 5 mg/day Placebo</td>
<td>1 year 3 years 3 years</td>
<td>2458</td>
<td>28-85 (68.6)</td>
<td>≤500 IU</td>
</tr>
<tr>
<td>3</td>
<td>R, PC, DB, MC, PG</td>
<td>2.5 mg/day 5 mg/day Placebo</td>
<td>2 years</td>
<td>543</td>
<td>45-80 (64.7)</td>
<td>-</td>
</tr>
<tr>
<td>4</td>
<td>R, PC, DB, MC, PG</td>
<td>2.5 mg/day 5 mg/day Placebo</td>
<td>12 – 18 months</td>
<td>648</td>
<td>39-80 (62.5)</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>R, AC, DB, MC, PG</td>
<td>5 mg/day 35 mg/week* 50 mg/week*</td>
<td>12 months</td>
<td>1456</td>
<td>48-95 (67.9)</td>
<td>≤500 IU</td>
</tr>
<tr>
<td>6</td>
<td>R, AC, DB, MC, PG</td>
<td>5 mg/day 35 mg/week*</td>
<td>24 months</td>
<td>922</td>
<td>50-87 (66.7)</td>
<td>800-1000 IU</td>
</tr>
<tr>
<td>7</td>
<td>R, AC, DB, MC, PG</td>
<td>5 mg/day 75 mg x 2 days/ month*</td>
<td>12 months</td>
<td>1229</td>
<td>50-86 (64.6)</td>
<td>400-800 IU</td>
</tr>
<tr>
<td>8</td>
<td>R, AC, DB, MC, PG</td>
<td>5 mg/day 150 mg once/month*</td>
<td>12 months</td>
<td>1292</td>
<td>50-88 (64.9)</td>
<td>400-500 to 1000 IU</td>
</tr>
</tbody>
</table>

*a* R: randomized; AC: active-controlled; PC: placebo-controlled; DB: double-blind; MC: multicentre; PG: parallel-group

* Placebo other days of treatment. † 35 mg enteric-coated following breakfast and before breakfast.

** Patients in these studies were supplemented with 1000 mg elemental calcium/day

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In Studies 1 and 2, patients were selected on the basis of radiographic evidence of previous vertebral fracture, and had established disease. The average number of prevalent vertebral fractures per patient at study entry was 4 in Study 1, and 2.5 in Study 2, with a broad range of baseline BMD levels. All fractures (symptomatic/painful/clinical vertebral fractures and asymptomatic/nonpainful/silent vertebral fractures) were systematically captured and measured by annual radiographs.

In Studies 3 to 5 postmenopausal women were recruited on the basis of low lumbar spine bone mass (i.e., more than 2 SD below the premenopausal mean) rather than a history of vertebral fracture.

In Studies 5 to 8, patients had either lumbar spine bone mass more than 2.5 SD below the premenopausal mean, or lumbar spine bone mass more than 2.0 SD below, and a prevalent vertebral fracture.

Patients with active or a history of upper gastrointestinal disorders at baseline and those taking ASA, NSAIDs or drugs usually used for the treatment of peptic ulcers were not specifically excluded from participating in the ACTONEL daily, weekly or monthly or ACTONEL DR weekly dosing osteoporosis studies.
Study Results

Results of Studies 1 and 2:

The pivotal studies of ACTONEL (risedronate sodium) in the treatment of postmenopausal osteoporosis clearly demonstrate that ACTONEL 5 mg daily reduces vertebral fracture incidence in patients with low bone mass and vertebral fractures, regardless of age, years since menopause or disease severity at baseline. ACTONEL 5 mg daily significantly reduced the risk of new vertebral fractures in each of the two large treatment studies. When measured by annual radiographs, the effect of ACTONEL 5 mg daily on vertebral fracture incidence was seen at the first year of treatment in each study. In the North American study, treatment with ACTONEL 5 mg daily for 1 year significantly reduced the risk of new vertebral fractures by 65% compared to treatment with placebo (p < 0.001). In the Multinational study, a similar significant reduction of 61% was seen (p = 0.001). Treatment with ACTONEL 5 mg daily also significantly reduced the proportion of patients experiencing new and worsening vertebral fractures in each of the studies. Figures 1 and 2 below display the cumulative incidence of vertebral and nonvertebral fractures (i.e., hip, wrist, humerus, clavicle, pelvis, and leg). In both figures, the cumulative incidence of these types of fractures is lower with ACTONEL compared with placebo at all time points, consistent with the positive effect of ACTONEL on bone strength.

Table 7
Effect of ACTONEL on Fracture, Height and Bone Mineral Density in the Treatment of Osteoporosis in Postmenopausal Women

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 5 mg</th>
<th>Placebo</th>
<th>Mean Difference from Placebo</th>
<th>Relative Risk Reduction %</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Study 1: VERT-MN</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cumulative incidence of new vertebral fracture over 3 years (% of patients)</td>
<td>18.1</td>
<td>29.0</td>
<td>49</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Median annual height change (mm/yr)</td>
<td>-1.33</td>
<td>-2.4</td>
<td></td>
<td>0.003</td>
<td></td>
</tr>
<tr>
<td>Mean increase in BMD (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 months Lumbar Spine</td>
<td>3.3</td>
<td>-0.1</td>
<td>3.4</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>36 months Lumbar Spine</td>
<td>7.1</td>
<td>1.3</td>
<td>5.9</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>2.0</td>
<td>-1.0</td>
<td>3.1</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Trochanter</td>
<td>5.1</td>
<td>-1.3</td>
<td>6.4</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>36 months Midshaft Radius</td>
<td>0.5</td>
<td>-1.9</td>
<td>2.4</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Study 2: VERT-NA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cumulative incidence of new vertebral Fracture over 3 years (% of patients)</td>
<td>11.3</td>
<td>16.3</td>
<td>41</td>
<td>0.003</td>
<td></td>
</tr>
<tr>
<td>Median annual height change (mm/yr)</td>
<td>-0.67</td>
<td>-1.14</td>
<td></td>
<td>0.001</td>
<td></td>
</tr>
<tr>
<td>Mean increase in BMD (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 months Lumbar Spine</td>
<td>2.7</td>
<td>0.4</td>
<td>2.2</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>36 months Lumbar Spine</td>
<td>5.4</td>
<td>1.1</td>
<td>4.3</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>1.6</td>
<td>-1.2</td>
<td>2.8</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Trochanter</td>
<td>3.3</td>
<td>-0.7</td>
<td>3.9</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>36 months Midshaft Radius</td>
<td>0.2</td>
<td>-1.4</td>
<td>1.6</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Prospectively Combined Studies 1 and 2: VERT-MN and VERT-NA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cumulative incidence of non-vertebral fracture over 3 years (% of patients)</td>
<td>7.1</td>
<td>11.0</td>
<td>36</td>
<td>0.005</td>
<td></td>
</tr>
</tbody>
</table>

a Measured by stadiometer
b Osteoporosis-related non-vertebral fractures (hip, wrist, humerus, clavicle, pelvis, and leg)
Figure 1
Cumulative New Vertebral Fracture Incidence in Postmenopausal Women with Osteoporosis

Study 1: “VERT-MN”

Study 2: “VERT-NA”

ACTONEL 5 mg daily was associated with a significant reduction of about 50% in the annual rate of height loss compared to treatment with placebo.

ACTONEL 5 mg daily produced increases in lumbar spine BMD which were progressive over the 3 years of treatment, and were statistically significant relative to baseline and to placebo at 6 months and at all later time points (12, 18, 24 and 36 months).
Results of Studies 3 and 4:

Table 8

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 5 mg Daily Mean Increase in BMD %</th>
<th>Placebo Mean Increase in BMD %</th>
<th>Mean Difference from Placebo %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Study 3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 months</td>
<td>Lumbar Spine</td>
<td>3.3</td>
<td>0.4</td>
</tr>
<tr>
<td>24 months</td>
<td>Lumbar Spine</td>
<td>4.1</td>
<td>-0.0</td>
</tr>
<tr>
<td></td>
<td>Femoral Neck</td>
<td>1.3</td>
<td>-1.0</td>
</tr>
<tr>
<td></td>
<td>Trochanter</td>
<td>2.7</td>
<td>-0.6</td>
</tr>
<tr>
<td>Study 4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 months</td>
<td>Lumbar Spine</td>
<td>3.3</td>
<td>0.7</td>
</tr>
<tr>
<td>18 months</td>
<td>Lumbar Spine</td>
<td>5.2</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>Femoral Neck</td>
<td>3.1</td>
<td>0.2</td>
</tr>
<tr>
<td></td>
<td>Trochanter</td>
<td>4.8</td>
<td>1.4</td>
</tr>
</tbody>
</table>

vs. placebo: *p<0.01; **p<0.001

In Studies 3 and 4, ACTONEL 5 mg daily produced significant mean increases in BMD of the lumbar spine compared to placebo at 6 months in women with low bone mass. Compared to placebo after 1.5 to 2 years, further significant mean increases in BMD were seen at the lumbar spine, femoral neck and trochanter.

The results of four large, randomized, placebo-controlled trials (Studies 1 to 4) in women with postmenopausal osteoporosis separately and together demonstrate that ACTONEL 5 mg daily reverses the progression of disease, increasing BMD at the spine, hip and wrist compared to the effects seen with placebo.

Results of Study 5:

Table 9

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 5 mg Daily Mean Increase in BMD % (95% Confidence Interval)</th>
<th>ACTONEL 35 mg Once-a-Week Mean Increase in BMD % (95% Confidence Interval)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 months</td>
<td>Lumbar Spine</td>
<td>n = 391</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(4.0, 4.3)</td>
</tr>
</tbody>
</table>

The results of the intent-to-treat analysis with the last observation carried forward were consistent with the primary efficacy analysis of completers. There were also no statistically significant differences between the two treatment groups at 1 year in regards to BMD increases from baseline at other skeletal sites (total proximal femur, femoral neck and femoral trochanter). Based on these BMD outcomes, ACTONEL 35 mg Once-a-Week was concluded to be non-inferior to ACTONEL 5 mg daily.

In trials with ACTONEL 5 mg daily, changes in BMD of this magnitude were associated with a significant decrease in fracture incidence relative to placebo (see Table 7). This is further supported by the fact that within the 1-year study comparing ACTONEL 35 mg Once-a-Week to ACTONEL 5 mg daily, no statistically significant differences amongst these treatment groups were seen with respect to the number of patients with at least 1 new fractured vertebra at 1 year. ACTONEL 35 mg taken once a week is similar in safety and efficacy to ACTONEL 5 mg daily for the treatment of postmenopausal osteoporosis.

Results of Study 6:

Table 10

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 5 mg Daily Mean Increase in BMD % (95% Confidence Interval)</th>
<th>ACTONEL DR 35 mg Weekly following breakfast Mean Increase in BMD % (95% Confidence Interval)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 months*</td>
<td>Lumbar Spine</td>
<td>n=307</td>
</tr>
<tr>
<td></td>
<td>3.1**</td>
<td>3.3**</td>
</tr>
<tr>
<td></td>
<td>(2.7, 3.5)</td>
<td>(2.9, 3.7)</td>
</tr>
<tr>
<td>24 months†</td>
<td>Lumbar Spine</td>
<td>4.1**</td>
</tr>
<tr>
<td></td>
<td>(3.7, 4.6)</td>
<td>(4.7, 5.7)</td>
</tr>
</tbody>
</table>

*Last available observation on or prior to month 12
†Last available observation on or prior to month 24

In a 2-year, double-blind, multicentre study of postmenopausal women with osteoporosis, ACTONEL DR 35 mg weekly was statistically shown to be non-inferior to ACTONEL 5 mg administered daily. At all time points, increases in BMD were statistically significant (p<0.05) compared to baseline for all sites measured.
At 1 year, ACTONEL DR 35 mg weekly was shown to be non-inferior to the ACTONEL 5 mg daily regimen for the primary efficacy variable of percent change from baseline of lumbar spine BMD. The two treatment groups were also similar with regard to percent change from baseline BMD at the total proximal femur, greater trochanter and femoral neck.

At 2 years, there were statistically significant greater increases (p<0.05) in mean percent change from baseline BMD at the greater trochanter for ACTONEL DR 35 mg weekly following breakfast (3.7) compared to ACTONEL 5 mg daily (2.8). This was also observed at the lumbar spine (see Table 10). The treatment groups were similar with regard to percent change from baseline BMD at the total proximal femur and femoral neck.

At 2 years, a statistically significant greater (p<0.05) percentage of patients in the ACTONEL DR 35 mg weekly group (89%) were considered responders (i.e., change from baseline in lumbar spine >0%) compared to the ACTONEL 5 mg group (82%).

Results of Study 7:

<table>
<thead>
<tr>
<th>Table 11</th>
<th>Comparison of ACTONEL on Two Consecutive Days Per Month vs. Daily Dosing in the Treatment of Osteoporosis in Postmenopausal Women – Primary Efficacy Analysis*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endpoints</td>
<td>ACTONEL 5 mg Daily Mean Increase in BMD % (95% Confidence Interval)</td>
</tr>
<tr>
<td>12 months Lumbar Spine</td>
<td>3.6 (3.3, 3.9)</td>
</tr>
<tr>
<td></td>
<td>N = 527</td>
</tr>
</tbody>
</table>

* The results of the intent-to-treat analysis with the last observation carried forward were consistent with the primary efficacy population analysis.

In the first year of a 2-year, double-blind, multicentre study of postmenopausal women with osteoporosis, ACTONEL 75 mg monthly duet, given on two consecutive days per month was shown to be non-inferior to ACTONEL 5 mg daily.

ACTONEL 75 mg on two consecutive days per month was statistically shown to be non-inferior to the ACTONEL 5 mg daily regimen for the primary efficacy variable of percent change from baseline to 1 year in lumbar spine BMD. The two treatment groups were also similar with regard to BMD increases at other skeletal sites. The incidences of vertebral and non-vertebral fractures, reported as adverse events, were similar in the two treatment groups. ACTONEL 75 mg taken on two consecutive days per month is similar in safety and efficacy to ACTONEL 5 mg daily for the treatment of postmenopausal osteoporosis. The safety and efficacy of ACTONEL 75 mg, on two consecutive days per month, has not yet been assessed beyond one year of treatment.

Results of Study 8:

<table>
<thead>
<tr>
<th>Table 12</th>
<th>Comparison of ACTONEL Once-a-Month vs. Daily Dosing in the Treatment of Osteoporosis in Postmenopausal Women – Primary Efficacy Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endpoints</td>
<td>ACTONEL 5 mg Daily Mean Increase in BMD % (95% Confidence Interval)</td>
</tr>
<tr>
<td>12 months (using LOCF*) Lumbar Spine</td>
<td>3.4 (3.0, 3.8)</td>
</tr>
<tr>
<td></td>
<td>n = 561</td>
</tr>
</tbody>
</table>

* LOCF: last observation carried forward

In the first year of a 2-year, double-blind, multicentre study of postmenopausal women with osteoporosis, ACTONEL 150 mg Once-a-Month was shown to be non-inferior to ACTONEL 5 mg daily. ACTONEL 150 mg Once-a-Month was statistically shown to be non-inferior to the ACTONEL 5 mg daily regimen for the primary efficacy variable of percent change from baseline to 1 year in increasing lumbar spine BMD. The two treatment groups were similar with regard to BMD increases at the lumbar spine, total proximal femur, femoral neck and femoral trochanter. The incidence of vertebral and non-vertebral fractures, reported as adverse events, was similar in the two treatment groups. ACTONEL 150 mg Once-a-Month is similar in safety and efficacy to ACTONEL 5 mg daily for the treatment of postmenopausal osteoporosis. The safety and efficacy of ACTONEL 150 mg Once-a-Month is currently being assessed beyond one year of treatment.

Histology/Histomorphometry: Histomorphometric evaluation of 278 bone biopsy samples from 204 postmenopausal women who received ACTONEL 5 mg or placebo once daily for 2 to 3 years (including 74 pairs of biopsies, 43 from ACTONEL-treated patients) showed a moderate and expected decrease in bone turnover in ACTONEL-treated women.

Histologic assessment showed no osteomalacia, impaired bone mineralization, or other adverse effects on bone in ACTONEL-treated women. These findings demonstrate that the bone formed during ACTONEL administration is of normal quality.
Prevention of Osteoporosis in Postmenopausal Women

Study Demographics and Trial Design

Table 13

<table>
<thead>
<tr>
<th>Study Number</th>
<th>Trial Design</th>
<th>Dosage</th>
<th>Duration</th>
<th>Patients</th>
<th>Age Range</th>
<th>Daily Supplement</th>
<th>Elemental Calcium</th>
<th>Vitamin D</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>R, PC, DB, MC, PG</td>
<td>2.5 mg/day 5 mg/day</td>
<td>2 years</td>
<td>383</td>
<td>42-63 (52.7)</td>
<td>1000 mg</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>R, DB, PC, MC, PG</td>
<td>35 mg/week Placebo</td>
<td>1 year</td>
<td>280</td>
<td>44-64 (53.6)</td>
<td>1000 mg</td>
<td>400 IU</td>
<td></td>
</tr>
</tbody>
</table>

R: randomized; PC: placebo-controlled; DB: double-blind; MC: multicentre; PG: parallel-group

Women in Study 9 were within 3 years of menopause and all patients in this study received supplemental calcium 1000 mg/day. Study 10 included women who were 0.5 to 5 years postmenopausal without osteoporosis. All patients were supplemented with 1000 mg elemental calcium and 400 IU vitamin D per day.

Results of Study 9:

Table 14

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 5 mg Mean Increase in BMD %</th>
<th>Placebo Mean Increase in BMD</th>
<th>Mean Difference from Placebo %</th>
</tr>
</thead>
<tbody>
<tr>
<td>24 months</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbar Spine</td>
<td>2.0</td>
<td>-2.5</td>
<td>4.5*</td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>1.0</td>
<td>-2.3</td>
<td>3.3*</td>
</tr>
<tr>
<td>Trochanter</td>
<td>2.3</td>
<td>-2.0</td>
<td>4.3*</td>
</tr>
</tbody>
</table>

* vs. placebo: p<0.001

Increases in BMD were observed as early as 3 months following initiation of ACTONEL treatment. Prevention of spinal bone loss was observed in the vast majority of women who received ACTONEL treatment. In contrast, most placebo-treated women experienced significant and progressive bone loss, despite receiving supplemental calcium 1000 mg/day. ACTONEL 5 mg daily was similarly effective in patients with lower baseline BMD (i.e., more than 1 SD below the premenopausal mean) and in those with higher BMD.

Results of Study 10:

Table 15

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 35 mg Once-a-Week Mean Increase in BMD %</th>
<th>Placebo Mean Increase in BMD</th>
<th>Mean Difference from Placebo %</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 months</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbar Spine</td>
<td>1.7</td>
<td>-0.5</td>
<td>2.2*</td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>1.0</td>
<td>-1.0</td>
<td>1.4*</td>
</tr>
<tr>
<td>Trochanter</td>
<td>0.4</td>
<td>-1.0</td>
<td>1.4*</td>
</tr>
<tr>
<td>12 months</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbar Spine</td>
<td>1.9</td>
<td>-1.1</td>
<td>3.0*</td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>0.3</td>
<td>-1.0</td>
<td>1.3**</td>
</tr>
<tr>
<td>Trochanter</td>
<td>1.0</td>
<td>-0.7</td>
<td>1.7*</td>
</tr>
</tbody>
</table>

vs. placebo: *p<0.0001; **p=0.0041
### Combined Administration with Hormone Replacement Therapy

#### Study Demographics and Trial Design

<table>
<thead>
<tr>
<th>Study Number</th>
<th>Trial Design</th>
<th>Dosage</th>
<th>Duration</th>
<th>Patients N = number</th>
<th>Age Range (Age Mean)</th>
<th>Gender</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>R, PC, DB, MC, PG, Stratified</td>
<td>ACTONEL 5 mg/day and conjugated estrogen 0.625 mg/day Placebo and conjugated estrogen 0.625 mg/day</td>
<td>1 year</td>
<td>524</td>
<td>37-82 (58.9)</td>
<td>Postmenopausal female</td>
</tr>
</tbody>
</table>

R: randomized; PC: placebo-controlled; DB: double-blind; MC: multicentre; PG: parallel-group

For inclusion in Study 11 women had a mean lumbar spine BMD 1.3 SD below the pre-menopausal mean and had recently initiated conjugated estrogen treatment (i.e., not taken estrogen for more than 1 month in the past year).

#### Results of Study 11:

<table>
<thead>
<tr>
<th>Endpoints</th>
<th>ACTONEL 5 mg Daily and Conjugated Estrogen Mean increase in BMD (%)</th>
<th>Conjugated Estrogen Mean increase in BMD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lumbar Spine</td>
<td>5.2</td>
<td>4.6</td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>2.7*</td>
<td>1.8</td>
</tr>
<tr>
<td>Trochanter</td>
<td>3.7</td>
<td>3.2</td>
</tr>
<tr>
<td>Midshaft Radius</td>
<td>0.7*</td>
<td>0.4</td>
</tr>
</tbody>
</table>

All values represent significant (p<0.05) change vs. baseline. vs. conjugated estrogen alone: *p<0.05

Consistent with the changes in BMD, the reduction in bone turnover, as measured by urinary deoxypyridinoline/creatinine, was significantly greater in the combined ACTONEL 5 mg daily plus estrogen group compared to the estrogen alone group (45-50% vs. 40%) and remained within the premenopausal range.

Histomorphometric evaluation of 93 bone biopsy samples from 61 women on testrogen therapy who received either placebo or ACTONEL 5 mg daily for 1 year (including 32 pairs of biopsies, 16 from ACTONEL-treated patients) found decreases in bone turnover in the ACTONEL-treated patients that were consistent with the changes in BTMs. Bone histology demonstrated that the bone of patients treated with ACTONELplus estrogen was of normal lamellar structure and normal mineralization.

### Treatment of Osteoporosis in Men, to Improve Bone Mineral Density

#### Study Demographics and Trial Design

<table>
<thead>
<tr>
<th>Study Number</th>
<th>Trial Design</th>
<th>Dosage</th>
<th>Duration</th>
<th>Patients N = number</th>
<th>Age Range (Age Mean)</th>
<th>Daily Supplement</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>R, DB, PC, MC, PG</td>
<td>ACTONEL 35 mg/week Placebo</td>
<td>2 years</td>
<td>191</td>
<td>36-84 (60.8)</td>
<td>1000 mg 400-500 IU</td>
</tr>
</tbody>
</table>

R: randomized; DB: double-blind; PC: placebo-controlled; MC: multicentre; PG: parallel-group

Patients with active or a history of upper gastrointestinal disorders at baseline and those taking ASA, NSAIDs, or drugs traditionally used for the treatment of peptic ulcers were not specifically excluded from participating in the 2-year male osteoporosis study.

#### Results of Study 12:

ACTONEL 35 mg Once-a-Week demonstrated efficacy in men with osteoporosis, as measured by change in BMD. All patients in this study received supplemental calcium 1000 mg/day and vitamin D 400-500 IU/day. ACTONEL 35 mg produced significant mean increases in BMD at the lumbar spine, femoral neck, trochanter and total hip compared to placebo in a 2 year study (lumbar spine, 4.5%; femoral neck, 1.1%; trochanter, 2.2%; total hip, 1.5%). Statistically significant increases in lumbar spine BMD were observed within 6 months following initiation of ACTONEL treatment. Lumbar spine BMD percent change from baseline at Months 6, 12 and 24 showed that the ACTONEL 35 mg Once-a-Week group had a statistically significant increase in mean percent change from baseline versus placebo at all time points (see Figure 3).
Glucocorticoid-Induced Osteoporosis

Study Demographics and Trial Design

Table 19
Summary of Patient Demographics for Clinical Trials of ACTONEL in the Prevention and Treatment of Glucocorticoid-Induced Osteoporosis

<table>
<thead>
<tr>
<th>Study Number</th>
<th>Trial Design</th>
<th>Dosage</th>
<th>Duration</th>
<th>Patients N = number</th>
<th>Age Range (Age Mean)</th>
<th>Gender</th>
</tr>
</thead>
<tbody>
<tr>
<td>13</td>
<td>Recent GC</td>
<td>DB, PC</td>
<td>5 mg/day</td>
<td>Placebo</td>
<td>1 year</td>
<td>228</td>
</tr>
<tr>
<td>14</td>
<td>Long-term GC</td>
<td>DB, PC</td>
<td>5 mg/day</td>
<td>Placebo</td>
<td>1 year</td>
<td>290</td>
</tr>
</tbody>
</table>

GC: glucocorticoid; DB: double-blind; PC: placebo-controlled

In Study 13, each patient had initiated glucocorticoid therapy (> 7.5 mg/day of prednisone or equivalent) within the previous 3 months for rheumatic, skin and pulmonary diseases. The mean lumbar spine BMD was normal at baseline. All patients in this study received supplemental calcium 500 mg/day.

Long-term use in Study 14 was defined as > 6 months of glucocorticoids for rheumatic, skin and pulmonary diseases. The baseline mean lumbar spine BMD was low (1.63 SD below the young healthy population mean), with 28% of the patients more than 2.5 SD below the mean. All patients in this study received supplemental calcium 1000 mg/day and supplemental vitamin D 400 IU/day.

Results of Studies 13 and 14:

Table 20
Change in Bone Mineral Density at 12 months from Baseline in Patients taking Glucocorticoid Therapy

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>ACTONEL 5 mg</th>
<th>Placebo</th>
<th>Least Squares</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean Change in BMD %</td>
<td>Mean Change in BMD %</td>
<td>Mean Difference from Placebo %</td>
</tr>
<tr>
<td>Study 13: Recent GC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbar Spine</td>
<td>N = 58-60</td>
<td>0.6</td>
<td>-2.8</td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>0.8</td>
<td>-3.1</td>
<td></td>
</tr>
<tr>
<td>Trochanter</td>
<td>1.4</td>
<td>-3.1</td>
<td></td>
</tr>
<tr>
<td>Study 14: Long-term GC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbar Spine</td>
<td>N = 77-79</td>
<td>2.9</td>
<td>0.4</td>
</tr>
<tr>
<td>Femoral Neck</td>
<td>1.8</td>
<td>-0.3</td>
<td>1.9*</td>
</tr>
<tr>
<td>Trochanter</td>
<td>2.4***</td>
<td>1.0</td>
<td>1.4*</td>
</tr>
</tbody>
</table>

GC: glucocorticoid; *p<0.01 vs. placebo; **p<0.001 vs. placebo; ***p<0.05 vs. baseline
By the third month of treatment, and continuing through treatment, the placebo group experienced losses in BMD at the lumbar spine, femoral neck and trochanter, while BMD was maintained or increased in the ACTONEL 5 mg group. At each skeletal site there were statistically significant differences between the ACTONEL 5 mg group and the placebo group at all time points (Months 3, 6, 9, 12). The treatment differences increased with continued treatment. The results at these skeletal sites were also statistically significant when the subgroups of men and postmenopausal women were analyzed separately.

ACTONEL was effective and prevented bone loss regardless of underlying disease, age, gender, glucocorticoid dose or baseline BMD.

**Vertebral Fractures:** Vertebral fractures were monitored for safety in the two placebo-controlled studies.

<table>
<thead>
<tr>
<th>Table 21</th>
<th>Incidence of Vertebral Fracture in Patients Initiating or Continuing Glucocorticoid Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endpoints</td>
<td>ACTONEL 5 mg Daily</td>
</tr>
<tr>
<td>N</td>
<td>% of patients</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Study 13: Recent GC</td>
<td>53</td>
</tr>
<tr>
<td>Study 14: Long-term GC</td>
<td>58</td>
</tr>
<tr>
<td>Combined Studies 13 and 14</td>
<td>111</td>
</tr>
</tbody>
</table>

vs. placebo: *p<0.05

The statistically significant reduction in vertebral fracture incidence in the analysis of the combined studies corresponded to a relative risk reduction of 70%.

**Histology/Histomorphometry:** Histomorphometric evaluation of 70 bone biopsy samples from 48 patients on glucocorticoid therapy who received either placebo or ACTONEL 5 mg daily for 1 year (including 22 pairs of biopsies, 16 from ACTONEL-treated patients) indicated that ACTONEL reduces bone resorption and produces a mild-to-moderate decrease in the rate of bone turnover. The rate of bone formation was preserved or increased and there was no evidence of impaired mineralization. The structure of the cortical bone (cortical thickness and porosity) was maintained in the ACTONEL-treated patients; cortical porosity increased, however, in the placebo group. These findings indicate that bone formed during ACTONEL treatment is of normal quality.

Bone histology demonstrated that bone formed during treatment with ACTONEL was of normal lamellar structure and normal mineralization, with no bone or marrow abnormalities observed.

**Paget’s Disease of Bone**

**Study Demographics and Trial Design**

<table>
<thead>
<tr>
<th>Table 22</th>
<th>Summary of Patient Demographics for Clinical Trials in the Treatment of Paget’s Disease of Bone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Study Number</td>
<td>Trial Design</td>
</tr>
<tr>
<td>15</td>
<td>DB, AC</td>
</tr>
<tr>
<td>16</td>
<td>AC</td>
</tr>
<tr>
<td>17</td>
<td>OL</td>
</tr>
<tr>
<td>18</td>
<td>OL</td>
</tr>
<tr>
<td>19</td>
<td>OL</td>
</tr>
<tr>
<td>20</td>
<td>OL</td>
</tr>
</tbody>
</table>

DB: double-blind; AC: active-controlled; OL: open-label

Patients in Study 15 had moderate-to-severe Paget’s disease (i.e., serum alkaline phosphatase levels of at least two times the upper limit of normal). The efficacy of ACTONEL 30 mg daily was demonstrated in six clinical studies involving over 390 male and female patients.
Results of Study 15:

Figure 4 below shows that at Day 180, 77% (43/56) of ACTONEL-treated patients achieved normalization of serum alkaline phosphatase levels compared to 10.5% of patients treated with Didronel (p < 0.001). For 33 of these 43 patients (77%), the remission (i.e., normalization of serum alkaline phosphatase) induced by ACTONEL treatment was maintained through at least 300 days of post-treatment observation.

During the first 180 days of the active-controlled study, 85% (51/60) of ACTONEL-treated patients demonstrated a ≥75% reduction from baseline in serum alkaline phosphatase excess (difference between measured level and midpoint of the normal range) with 2 months of treatment compared to 20% (12/60) in the Didronel-treated group with 6 months of treatment (p < 0.001). Changes in serum alkaline phosphatase excess over time (shown in Figure 5 below) reveal that the onset of the effect of ACTONEL is significant following only 30 days of treatment, with a 36% reduction in serum alkaline phosphatase excess at that time compared to only 6% seen with Didronel treatment at the same time point (p < 0.001).

Response to ACTONEL therapy was independent of age, gender, or race and was similar in patients with mild to very severe Paget’s disease. Table 23 below shows the maximum mean percent reduction from baseline in excess serum alkaline phosphatase in patients with mild, moderate, or severe disease.
atrophy was also noted in male rats after 13 weeks of treatment at oral doses of 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²).

In female rats, ovulation was inhibited at an oral dose of 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²).

In a 1-year daily repeat dose toxicity study in dogs, the limiting toxicity of risedronate was observed at 8 mg/kg/day (160 mg/m²).

In 6 month and 1-year monthly repeat dose toxicity studies in dogs, the limiting systemic toxicity of risedronate was observed at 32 mg/kg (640 mg/m²) and consisted of liver, testicular, and renal toxicities. Gastric lesions were observed at 16 mg/kg (320 mg/m²). These doses are equivalent to approximately 3.5 and 7 times the human 10 mg dose based on surface area, mg/m².

A 13-week oral dog study was performed to evaluate the gastric and lower gastrointestinal toxicity and toxicokinetics of risedronate (8 and 16 mg/kg) when dosed with or without EDTA (2.5 and 12.5 mg/kg) following 14 once-weekly oral doses. No additional GI toxicity was observed with the addition of either dose of EDTA to either dose of risedronate. No new organs of toxicity were identified when dogs were treated with risedronate in combination with EDTA (vs risedronate alone). Treatment with EDTA alone was not associated with any treatment-related changes.

Co-administration of EDTA with 8 and/or 16 mg/kg risedronate was associated with potentiation of risedronate-mediated histologic alterations in the liver, kidneys, and testes (incidence and/or severity). Potentiation of toxicity was more evident at 12.5 mg/kg EDTA when compared with 2.5 mg/kg EDTA. With respect to expected pharmacological effects (e.g. increased bone), 12.5 mg/kg EDTA potentiated the severity of rib hypertrophy and the incidence of increased bone in nasal turbinates when administered in combination with 8 and 16 mg/kg risedronate (vs risedronate alone). These findings may be related to the observed increase in exposure noted when risedronate was administered in combination with EDTA.

Carcinogenicity: Three carcinogenicity studies in two species (mouse and rat) have been completed. All studies clearly showed dose-dependent bone pharmacologic effects. Risedronate was not carcinogenic in male or female rats dosed daily by gavage for 104 weeks at doses up to 24 mg/kg/day (12 times the human 30 mg dose based on surface area, mg/m²). Similarly, there was no evidence of a carcinogenic potential in male or female mice dosed daily by gavage for 80 weeks at doses up to 32 mg/kg/day (5 times the human 30 mg dose based on surface area, mg/m²).

Mutagenesis: In a series of seven in vitro and in vivo mutagenicity assays, risedronate was not genotoxic. An in vitro chromosomal aberration assay in Chinese hamster ovary cells was weakly positive at highly cytotoxic doses (> 675 µg/mL). However, when the assay was repeated at doses exhibiting increased cell survival (300 µg/mL), risedronate was negative.

Reproduction: In female rats, ovulation was inhibited at an oral dose of 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²). Decreased implantation was noted in female rats treated with doses ≥ 7 mg/kg/day (approximately 2.3 times the 30 mg/day human dose based on surface area, mg/m²). In male rats, testicular and epididymal atrophy and inflammation were noted at 40 mg/kg/day (approximately 13 times the 30 mg/day human dose based on surface area, mg/m²). Testicular atrophy was also noted in male rats after 13 weeks of treatment at oral doses of 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²). There was moderate-to-severe spermatid maturation block after 13 weeks in male dogs at an oral dose of 8 mg/kg/day (approximately 8 times the 30 mg/day human dose based on surface area, mg/m²). These findings tended to increase in severity with increased dose and exposure time.

### TOXICOLOGY

#### Acute Toxicity:
Lethality after single oral doses was seen in female rats at 903 mg/kg (5826 mg/m²) and male rats at 1703 mg/kg (10967 mg/m²). The minimum lethal dose in mice, rabbits, and dogs was 4000 mg/kg (10909 mg/m²).

#### Chronic Toxicity:
In a 1-year daily repeat dose toxicity study in dogs, the limiting toxicity of risedronate was observed at 8 mg/kg/day (160 mg/m²) and consisted of liver, testicular, renal, and gastrointestinal changes. Gastrointestinal effects at 16 mg/kg (320 mg/m²) and consisted of liver, testicular, and renal toxicities. Gastric lesions were observed at 16 mg/kg (320 mg/m²). These doses are equivalent to approximately 3.5 and 7 times the human 150 mg dose based on surface area, mg/m².

#### Long-term Oral Administration:
In a 104-week study in dogs, the limiting toxicity of risedronate was observed at 8 mg/kg/day (160 mg/m²) and consisted of liver, testicular, renal, and gastrointestinal changes. Gastrointestinal effects at 16 mg/kg (320 mg/m²) and consisted of liver, testicular, and renal toxicities. Gastric lesions were observed at 16 mg/kg (320 mg/m²). These doses are equivalent to approximately 3.5 and 7 times the human 150 mg dose based on surface area, mg/m².

#### Radiographs:
Radiographs taken at baseline and after 6 months from patients treated with ACTONEL 30 mg daily demonstrate that ACTONEL is highly effective in decreasing the extent of osteolysis across all anatomical sites including the appendicular and axial skeleton. Importantly, osteolytic lesions in the lower extremities improved or were unchanged in 15/16 (94%) of assessed patients; 9/15 (60%) patients showed clear improvement in osteolytic lesions. No evidence of new fractures was observed.

### DETAILED PHARMACOLOGY

#### There are extensive preclinical data to support that bone produced during ACTONEL (risedronate sodium) treatment at therapeutic doses is of normal quality, consistent with clinical experience. Risedronate demonstrated potent anti-osteoclast, antiresorptive activity in ovariectomized animals, increasing bone mass and biomechanical strength dose-dependently.

#### Chronic Toxicity:
In a 1-year daily repeat dose toxicity study in dogs, the limiting toxicity of risedronate was observed at 8 mg/kg/day (160 mg/m²) and consisted of liver, testicular, renal, and gastrointestinal changes. Gastrointestinal effects at 16 mg/kg (320 mg/m²) and consisted of liver, testicular, and renal toxicities. Gastric lesions were observed at 16 mg/kg (320 mg/m²). These doses are equivalent to approximately 3.5 and 7 times the human 150 mg dose based on surface area, mg/m².

#### A 13-week oral dog study was performed to evaluate the gastric and lower gastrointestinal toxicity and toxicokinetics of risedronate (8 and 16 mg/kg) when dosed with or without EDTA (2.5 and 12.5 mg/kg) following 14 once-weekly oral doses. No additional GI toxicity was observed with the addition of either dose of EDTA to either dose of risedronate. No new organs of toxicity were identified when dogs were treated with risedronate in combination with EDTA (vs risedronate alone). Treatment with EDTA alone was not associated with any treatment-related changes.

Co-administration of EDTA with 8 and/or 16 mg/kg risedronate was associated with potentiation of risedronate-mediated histologic alterations in the liver, kidneys, and testes (incidence and/or severity). Potentiation of toxicity was more evident at 12.5 mg/kg EDTA when compared with 2.5 mg/kg EDTA. With respect to expected pharmacological effects (e.g. increased bone), 12.5 mg/kg EDTA potentiated the severity of rib hypertrophy and the incidence of increased bone in nasal turbinates when administered in combination with 8 and 16 mg/kg risedronate (vs risedronate alone). These findings may be related to the observed increase in exposure noted when risedronate was administered in combination with EDTA.

#### Carcinogenicity:
Three carcinogenicity studies in two species (mouse and rat) have been completed. All studies clearly showed dose-dependent bone pharmacologic effects. Risedronate was not carcinogenic in male or female rats dosed daily by gavage for 104 weeks at doses up to 24 mg/kg/day (12 times the human 30 mg dose based on surface area, mg/m²). Similarly, there was no evidence of a carcinogenic potential in male or female mice dosed daily by gavage for 80 weeks at doses up to 32 mg/kg/day (5 times the human 30 mg dose based on surface area, mg/m²).

#### Mutagenesis:
In a series of seven in vitro and in vivo mutagenicity assays, risedronate was not genotoxic. An in vitro chromosomal aberration assay in Chinese hamster ovary cells was weakly positive at highly cytotoxic doses (> 675 µg/mL). However, when the assay was repeated at doses exhibiting increased cell survival (300 µg/mL), risedronate was negative.

#### Reproduction:
In female rats, ovulation was inhibited at an oral dose of 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²). Decreased implantation was noted in female rats treated with doses ≥ 7 mg/kg/day (approximately 2.3 times the 30 mg/day human dose based on surface area, mg/m²). In male rats, testicular and epididymal atrophy and inflammation were noted at 40 mg/kg/day (approximately 13 times the 30 mg/day human dose based on surface area, mg/m²). Testicular atrophy was also noted in male rats after 13 weeks of treatment at oral doses of 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²). There was moderate-to-severe spermatid maturation block after 13 weeks in male dogs at an oral dose of 8 mg/kg/day (approximately 8 times the 30 mg/day human dose based on surface area, mg/m²). These findings tended to increase in severity with increased dose and exposure time.
Survival of neonates was decreased in rats treated during gestation with oral doses ≥ 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²). Body weight was decreased in neonates from dams treated with 80 mg/kg (approximately 26 times the 30 mg/day human dose based on surface area, mg/m²). In rats treated during gestation, the number of fetuses exhibiting incomplete ossification of sternebrae or skull was statistically significantly increased at 7.1 mg/kg/day (approximately 2.3 times the 30 mg/day human dose based on surface area, mg/m²). Both incomplete ossification and un ossified sternebrae were increased in rats treated with oral doses ≥ 16 mg/kg/day (approximately 5.2 times the 30 mg/day human dose based on surface area, mg/m²). A low incidence of cleft palate was observed in fetuses from female rats treated with oral doses ≥ 3.2 mg/kg/day (approximately 1 time the 30 mg/day human dose based on surface area, mg/m²). The relevance of this finding to human use of ACTONEL is unclear. No significant fetal ossification effects were seen in rabbits treated with oral doses up to 10 mg/kg/day during gestation (approximately 6.7 times the 30 mg/day human dose based on surface area, mg/m²). However, in rabbits treated with 10 mg/kg/day, 1 of 14 litters were aborted and 1 of 14 litters were delivered prematurely.

Similar to other bisphosphonates, treatment during mating and gestation with doses as low as 3.2 mg/kg/day (approximately 1 time the 30 mg/day human dose based on surface area, mg/m²) has resulted in parturient hypocalcemia and mortality in pregnant rats allowed to deliver.

Bisphosphonates are incorporated into the bone matrix, from which they are gradually released over periods of weeks to years. The amount of bisphosphonate incorporation into adult bone, and hence, the amount available for release back into the systemic circulation, is directly related to the dose and duration of bisphosphonate use. There are no data on fetal risk in humans. However, there is a theoretical risk of fetal harm, predominantly skeletal, if a woman becomes pregnant after completing a course of bisphosphonate therapy. The impact of variables such as time between cessation of bisphosphonate therapy to conception, the particular bisphosphonate used, and the route of administration (intravenous versus oral) on this risk has not been studied.

REFERENCES

PART III: CONSUMER INFORMATION

ACTONEL®
Risedronate Sodium Tablets

and

ACTONEL DR™
Risedronate Sodium Delayed-Release Tablets

This leaflet is Part III of a three-part “Product Monograph” published for ACTONEL and ACTONEL DR. It is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about ACTONEL or ACTONEL DR. Contact your doctor or pharmacist if you have any questions about the drug.

ABOUT THIS MEDICATION

What the medication is used for:
• Treatment and prevention of osteoporosis in postmenopausal women
• Treatment of osteoporosis in men, to improve bone mineral density
• Treatment and prevention of osteoporosis caused by treatment with steroid medication such as prednisone
• Treatment of Paget’s disease of bone

What it does:
ACTONEL and ACTONEL DR is a bisphosphonate drug that helps to slow bone loss. In many people, ACTONEL and ACTONEL DR help to increase bone density. In osteoporosis, the body removes more bone than it replaces. This causes bones to get weaker and more likely to break or fracture (usually at the spine, wrist or hip). Spine fractures may result in a curved back, height loss or back pain. ACTONEL and ACTONEL DR correct this imbalance by decreasing the elevated rate of bone removal. ACTONEL and ACTONEL DR can therefore help reduce the risk of spine and non-spine fractures.

Your doctor may measure the thickness (i.e., density) of your bone through a bone mineral density (BMD) test or x-ray to check your progress against further bone loss or fracture.

ACTONEL and ACTONEL DR are not pain relievers. Your doctor may prescribe or recommend another medicine specifically for pain relief.

When it should not be used:
• If you have low blood calcium levels (hypocalcemia).
• If you are allergic to risedronate sodium or any other ingredients in ACTONEL or ACTONEL DR.

What the medicinal ingredient is:
Risedronate sodium

What the nonmedicinal ingredients are:
ACTONEL (film-coated) tablets:
Crospovidone, ferric oxide (5, 35 & 75 mg), hydroxypropyl cellulose, hypromellose, indigo carmine (150 mg), lactose monohydrate (5, 30 & 35 mg), magnesium stearate, microcrystalline cellulose, polyethylene glycol, silicon dioxide and titanium dioxide.

ACTONEL DR (delayed-release) tablets
Edetate disodium, ferric oxide yellow, magnesium stearate, methacrylic acid copolymer dispersion, polysorbate 80, silicified microcrystalline cellulose, simethicone, sodium starch glycolate, stearic acid, talc and triethyl citrate.

What dosage form it comes in:
ACTONEL film-coated tablets:
Each ACTONEL tablet contains risedronate sodium 5 mg (yellow), 30 mg (white), 35 mg (orange), 75 mg (pink) or 150 mg (blue).

ACTONEL DR tablets:
Each yellow, enteric coated ACTONEL DR tablet contains risedronate sodium 35 mg.

The ACTONEL DR tablet has an enteric coating, which delays the release of risedronate until the small intestine.

WARNINGS AND PRECAUTIONS

Before you use ACTONEL or ACTONEL DR, talk to your doctor or pharmacist if:
• You have had problems or disease in your kidneys, esophagus (the tube connecting the mouth and the stomach), stomach or intestines.
• You cannot carry out dosing instructions (see PROPER USE OF THIS MEDICATION).
• You are pregnant or nursing.
• You have one of the following risk factors: cancer, chemotherapy, radiotherapy of the head or neck, treatment with corticosteroids or dental problems or dental infections. If so, a dental examination and any necessary dental procedures should be considered before you start treatment with ACTONEL or ACTONEL DR.

Calcium and vitamin D are also important for strong bones. Your doctor may ask you to take calcium and vitamin D while you are on ACTONEL or ACTONEL DR therapy (see INTERACTIONS WITH THIS MEDICATION section).
INTERACTIONS WITH THIS MEDICATION

If taken with some other medicines, the effects of ACTONEL, ACTONEL DR or the effects of other medicines may be changed. It is important to tell your health care providers, including doctors and dentists, about all medications you are taking, even if the medicine does not require a prescription (including vitamin and herbal supplements).

ACTONEL

You should not take ACTONEL with food, as it may prevent your body from absorbing or using ACTONEL. You should take ACTONEL on an empty stomach. (See PROPER USE OF THIS MEDICATION for instruction).

Vitamins, mineral supplements, antacids and other medications may contain substances (e.g., calcium, magnesium, aluminum, and iron) which can stop your body from absorbing or using ACTONEL. These medications should be taken at a different time of day than ACTONEL.

If you are taking ACTONEL for Paget's disease, talk to your doctor before taking ASA or other non-steroidal anti-inflammatory drugs used for pain management because the risk of stomach upset may be increased.

ACTONEL DR

ACTONEL DR should be taken with food. Talk to your doctor to ensure that you fully understand which dosage form you have been prescribed.

The following medications may reduce the absorption of ACTONEL DR and should be taken at a different time of the day than ACTONEL DR:

- Calcium supplements, or antacids, or medicines containing magnesium, aluminum, or iron
- Medicines that reduce the stomach acid

PROPER USE OF THIS MEDICATION

As with all medications, it is important to take as directed by your doctor.

Usual Dose:

**Treatment of postmenopausal osteoporosis:**
- 1 ACTONEL tablet (5 mg) per day or
- 1 ACTONEL tablet (35 mg) per week or
- 1 ACTONEL DR tablet (35 mg) per week or
- 1 ACTONEL tablet (75 mg) per day for two consecutive days each month, on the same two calendar days each month or
- 1 ACTONEL tablet (150 mg) once per month, on the same calendar day each month

**Prevention of postmenopausal osteoporosis:**
- 1 ACTONEL tablet (5 mg) per day or
- 1 ACTONEL tablet (35 mg) per week

**Treatment of osteoporosis in men, to improve bone mineral density:**
- 1 ACTONEL tablet (35 mg) per week

**Treatment and prevention of glucocorticoid-induced osteoporosis:**
- 1 ACTONEL tablet (5 mg) per day

**Paget’s disease of bone:**
- 1 ACTONEL tablet (30 mg) per day

ACTONEL DOSING INSTRUCTIONS (For ACTONEL DR dosing instructions see below)

- ACTONEL should be taken first thing in the morning with plain water before you have anything to eat or drink.
- Choose a day of the week to take your tablet.
- On your chosen day, take 1 ACTONEL tablet first thing in the morning with plain water before you have anything to eat or drink.

ACTONEL is not the same as ACTONEL DR. Clinical benefits may be compromised by failure to take ACTONEL on an empty stomach. Only ACTONEL DR can be taken with food. Talk to your health professional to ensure that you are clear on the dosage form you have been prescribed.

Once daily dosing (5 mg or 30 mg per day):
- Take 1 ACTONEL tablet first thing in the morning with plain water before you have anything to eat or drink.

Once weekly dosing (35 mg per week – see below for ACTONEL DR dosing instructions):
- Choose a day of the week to take your tablet.
- On your chosen day, take 1 ACTONEL tablet first thing in the morning with plain water before you have anything to eat or drink.

Monthly duet dosing (75 mg on two consecutive days per month):
- Choose 2 days in a row each month that you will remember and that best fits your schedule to take your ACTONEL doses.
- Take 1 ACTONEL tablet first thing in the morning with plain water before you have anything to eat or drink on your first chosen day.
- Take the second ACTONEL tablet in the morning of the following day with plain water before you have anything to eat or drink.
- ACTONEL should be taken on the same two consecutive calendar days each month.

ACTONEL should be taken in the morning on an empty stomach at least 30 minutes before consuming the first food, drink (other than plain water) and/or any other medication of the day. Food, medication or drink other than plain water can interfere with the absorption of ACTONEL.
- Each ACTONEL tablet should be swallowed whole while you are in an upright position and with sufficient plain water (≥120 mL or ½ cup) to facilitate delivery to the stomach.
- Aside from plain water, do not eat or drink for at least 30 minutes after taking ACTONEL.
- You should not lie down for at least 30 minutes after taking the medication. You may sit, stand or do normal activities like read the newspaper, take a walk, etc.
- ACTONEL tablets should not be chewed, cut, or crushed.

These recommendations help ACTONEL work correctly and help you avoid possible irritation of the esophagus (the tube connecting the mouth and the stomach).
Choose a day of the month that you will remember and that best fits your schedule to take your ACTONEL dose (1 tablet of 150 mg) once per month. On your chosen day each month, take 1 ACTONEL 150 mg tablet first thing in the morning with plain water before you have anything to eat or drink.

**ACTONEL DR DOSSING INSTRUCTIONS**

- ACTONEL DR should be taken in the morning with breakfast (including high fat foods, coffee, tea, milk, orange juice, etc.). Do not take ACTONEL DR before breakfast or on an empty stomach. This may cause upper abdominal pain.
- Each ACTONEL DR tablet should be swallowed whole while you are in an upright position and with sufficient plain water (≥ 120 mL or ½ cup) to facilitate delivery to the stomach.
- You should not lie down for at least 30 minutes after taking the medication. You may sit, stand or do normal activities like read the newspaper, take a walk, etc.
- ACTONEL DR tablets should not be chewed, cut, or crushed. Care should be taken not to break the outer coating which is designed to remain intact until the tablet reaches the small intestine where the tablet coating dissolves and releases the active ingredient.

These recommendations help ACTONEL DR work correctly and help you avoid possible irritation of the esophagus (i.e., the tube connecting the mouth and the stomach).

**ACTONEL DR Once monthly dosing (35 mg per week):**

- Choose a day of the week to take your tablet.
- On your chosen day, take 1 ACTONEL DR tablet in the morning with plain water.

You should take ACTONEL or ACTONEL DR for as long as your doctor recommends, to continue to prevent bone loss and protect your bones from fractures.

**Missed Dose:**

**Daily dose (ACTONEL 5 mg or 30 mg tablet):** If you forget to take your dose, do not double your next dose (i.e., do not take 2 tablets on the same day). Simply take 1 tablet at your next scheduled time.

**Weekly dose (35 mg ACTONEL tablet or 35 mg ACTONEL DR tablet):** If you forget to take your dose on the regularly scheduled day, simply take 1 tablet on the day you first remember having missed your dose. Then resume your schedule by taking 1 tablet on the originally chosen day of the week. If you’ve missed your dose by exactly one week, do not take 2 tablets on the same day. Simply take 1 tablet as you normally would have on this day and resume your usual weekly schedule.

**Monthly duet dose (ACTONEL 75 mg tablet):** If you miss one or both tablets of your dose of ACTONEL 75 mg in the morning, do not take it later in the day. If the next month’s scheduled doses are more than 7 days away do the following:

- If both tablets were missed, take the first ACTONEL 75 mg tablet on the morning after the day it is remembered and the second tablet on the next consecutive morning.
- If only one tablet is missed, take the missed tablet on the morning after the day it is remembered.

You should then continue your usual schedule of ACTONEL 75 mg on two consecutive days each month. Do not take more than two 75 mg tablets within 7 days.

If the next month’s scheduled doses are 1 to 7 days away, you should wait until next month’s scheduled doses and then resume taking ACTONEL 75 mg on two consecutive days each month as originally scheduled.

**Once monthly dose (ACTONEL 150 mg tablet):** If you miss the 150 mg dose of ACTONEL (1 tablet of 150 mg) in the morning, do not take it later in the day. If the next month’s scheduled dose is more than 7 days away, take the missed tablet on the morning after the day it is remembered. You should then return to your usual schedule of ACTONEL 150 mg.

If the next month’s scheduled dose is 1 to 7 days away, you should wait until next month’s scheduled dose and then resume taking ACTONEL 150 mg as originally scheduled. Do not take more than 150 mg of ACTONEL within 7 days.

**Overdose:**

If you take too many tablets on any given day, contact your doctor, or a Poison Control Centre, or an emergency room of the nearest hospital immediately.

For ACTONEL overdose, drink a full glass of milk. Do not induce vomiting.

**SIDE EFFECTS AND WHAT TO DO ABOUT THEM**

Drugs like ACTONEL and ACTONEL DR may cause problems in your esophagus (the tube connecting the mouth and the stomach), stomach and intestines, including ulcers. If you have trouble or pain upon swallowing, heartburn, chest pain and black or bloody stools, stop taking ACTONEL or ACTONEL DR and tell your doctor right away. Remember to take ACTONEL or ACTONEL DR as directed.

**ACTONEL side effects:** In clinical studies of osteoporosis with ACTONEL, the most commonly reported side effects were abdominal pain, heartburn and nausea. In studies of Paget’s disease, diarrhea and headache were also commonly reported.

ACTONEL may cause pain in bones, joints or muscles, rarely severe. Pain may start as soon as one day or up to several months after starting ACTONEL.

ACTONEL at monthly doses may cause short-lasting, mild flu-like symptoms. These symptoms usually decrease after subsequent doses.

Very rarely patients have reported non-healing jaw wounds while receiving ACTONEL or other drugs in this class. Consult your doctor if you experience persistent pain in your mouth, teeth or jaw, or if your gums or mouth heal poorly.

Very rarely patients have reported unusual fractures in their thigh bone while receiving drugs in this class. Consult your doctor if you experience new or unusual pain in your hip, groin, or thigh.

**ACTONEL DR side effects:** Other side effects of ACTONEL DR include:

- Most common side effects:
  - Abdominal pain, heartburn, nausea

- Other common side effects:
  - Pain in bones, joints, or muscles, rarely severe. The pain may start as soon as one day or up to several months after starting ACTONEL DR.
  - Diarrhea, constipation, inflammation of the pharynx and/or nose, upper respiratory tract infection.
Very Rare Side Effects:

- Non-healing jaw. Talk to your doctor if you experience persistent pain in your mouth, teeth or jaw, or if your gums or mouth poorly.
- Unusual fractures in the thigh bone. Talk to your doctor if you experience new or unusual pain in your hip, groin or thigh.

### IMPORTANT SIDE EFFECTS, HOW OFTEN THEY HAPPEN AND WHAT TO DO ABOUT THEM

<table>
<thead>
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<td></td>
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<tr>
<td>Pain in bones, joints, or muscles</td>
<td>✓</td>
<td></td>
</tr>
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<td>Abdominal pain</td>
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</tr>
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</tr>
<tr>
<td>Painful tongue</td>
<td>✓</td>
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<td></td>
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This is not a complete list of side effects. For any unexpected effects while taking ACTONEL or ACTONEL DR, contact your doctor or pharmacist.

### HOW TO STORE IT

- Keep ACTONEL or ACTONEL DR and all other medications out of the reach of children.
- Keep the tablets in their original package and store at controlled room temperature (20°-25°C).
- Do not keep medicine that is out of date or that you no longer need.

### REPORTING SUSPECTED SIDE EFFECTS

You can report any suspected adverse reactions associated with the use of health products to the Canada Vigilance Program by one of the following 3 ways:

- Report online at www.healthcanada.gc.ca/medeffect
- Call toll-free at 1-866-234-2345
- Complete a Canada Vigilance Reporting Form and:
  - Fax toll-free to 1-866-678-6789, or
  - Mail to: Canada Vigilance Program
    Health Canada
    Postal Locator 0701D
    Ottawa, Ontario K1A 0K9

Postage paid labels, Canada Vigilance Reporting Form and the adverse reaction reporting guidelines are available on the MedEffect™ Canada Web site at www.healthcanada.gc.ca/medeffect.

NOTE: Should you require information related to the management of side effects, contact your health professional. The Canada Vigilance Program does not provide medical advice.

### MORE INFORMATION

For more information, please visit [http://www.actonel.ca](http://www.actonel.ca).

This document plus the full product monograph, prepared for health professionals is available by contacting Warner Chilcott Canada Co. at: 1-800-565-0814.

This leaflet was prepared by Warner Chilcott Canada Co.

Last Revised: July 15, 2011
IMPORTANT: PLEASE READ

PART III: CONSUMER INFORMATION

ACTONEL DR™

Risedronate Sodium Delayed-Release Tablets

This leaflet is Part III of a three-part “Product Monograph” for ACTONEL DR. It is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about ACTONEL DR. Contact your doctor or pharmacist if you have any questions about the drug.

ABOUT THIS MEDICATION

What the medication is used for:
• Treatment of osteoporosis in postmenopausal women

What it does:
ACTONEL DR is a bisphosphonate drug that helps to slow bone loss. In many people, ACTONEL DR helps to increase bone density. In osteoporosis, the body removes more bone than it replaces. This causes bones to get weaker and more likely to break or fracture (usually at the spine, wrist or hip). Spine fractures may result in a curved back, height loss or back pain. ACTONEL DR corrects this imbalance by decreasing the elevated rate of bone removal. ACTONEL DR can therefore help reduce the risk of spine and non-spine fractures. Your doctor may measure the thickness (i.e., density) of your bone through a bone mineral density (BMD) test or x-ray to check your progress against further bone loss or fracture.

ACTONEL DR is not a pain reliever. Your doctor may prescribe or recommend another medicine specifically for pain relief.

When it should not be used:
• If you have low blood calcium levels (hypocalcemia).
• If you are allergic to risedronate sodium or any other ingredients in ACTONEL DR.

What the medicinal ingredient is:
Risedronate sodium

What the nonmedicinal ingredients are:
Edetate disodium, ferric oxide yellow, magnesium stearate, methacrylic acid copolymer dispersion, polysorbate 80, silicified microcrystalline cellulose, simethicone, sodium starch glycolate, stearic acid, talc and triethyl citrate.

What dosage form it comes in:
Each yellow, enteric coated ACTONEL DR tablet contains risedronate sodium 35 mg.

The ACTONEL DR tablet has an enteric coating, which delays the release of risedronate until the small intestine.

WARNINGS AND PRECAUTIONS

Before you use ACTONEL DR, talk to your doctor or pharmacist if:
• You have had problems or disease in your kidneys, esophagus (the tube connecting the mouth and the stomach), stomach, or intestines.
• You cannot carry out dosing instructions (see PROPER USE OF THIS MEDICATION).
• You are pregnant or nursing.
• You have one of the following risk factors: cancer, chemotherapy, radiotherapy of the head or neck, treatment with corticosteroids, or dental problems or dental infections. If so, a dental examination and any necessary dental procedures should be considered before you start treatment with ACTONEL DR.

Calcium and vitamin D are also important for strong bones. Your doctor may ask you to take calcium and vitamin D while you are on ACTONEL DR therapy (see INTERACTIONS WITH THIS MEDICATION section).

INTERACTIONS WITH THIS MEDICATION

If taken with some other medicines, the effects of ACTONEL DR or the effects of other medicines may be changed. It is important to tell your health care providers, including doctors and dentists, about all medications you are taking, even if the medicine does not require a prescription (including vitamin and herbal supplements).

The following medications may reduce the absorption of ACTONEL DR and should be taken at a different time of the day than ACTONEL DR:
• Calcium supplements, or antacids, or medicines containing magnesium, aluminum, or iron
• Medicines that reduce the stomach acid

PROPER USE OF THIS MEDICATION

As with all medications, it is important to take as directed by your doctor.

Usual Dose:
• 1 ACTONEL DR tablet (35 mg) per week
• Choose a day of the week to take your tablet.
• On your chosen day, take 1 ACTONEL DR tablet in the morning with plain water.

Instructions for Use
• ACTONEL DR should be taken in the morning with breakfast (including high fat foods, coffee, tea, milk, orange juice, etc.). Do not take ACTONEL DR before breakfast or on an empty stomach. This may cause upper abdominal pain.
Each ACTONEL DR tablet should be swallowed whole while in an upright position and with sufficient plain water (≥ 120 mL or ½ cup) to facilitate delivery to the stomach. You should not lie down for at least 30 minutes after taking the medication. You may sit, stand or do normal activities like read the newspaper, take a walk, etc. ACTONEL DR tablets should not be chewed, cut, or crushed. Care should be taken not to break the outer coating which is designed to remain intact until the tablet reaches the small intestine where the tablet coating dissolves and releases the active ingredient. These recommendations help ACTONEL DR work correctly and help you avoid possible irritation of the esophagus (i.e., the tube connecting the mouth and the stomach).

You should take ACTONEL DR for as long as your doctor recommends, to continue to prevent bone loss and protect your bones from fractures.

**Missed Dose:**
If you forget to take your dose on the regularly scheduled day, simply take 1 tablet on the day you first remember having missed your dose. Then resume your schedule by taking 1 tablet on the originally chosen day of the week. If you’ve missed your dose by exactly one week, do not take 2 tablets on the same day. Simply take 1 tablet as you normally would have on this day and resume your usual weekly schedule.

**Overdose:**
If you take too many tablets on any given day, contact your doctor, or a Poison Control Centre, or an emergency room of the nearest hospital immediately.

**SIDE EFFECTS AND WHAT TO DO ABOUT THEM**

ACTONEL DR may cause problems in your esophagus (the tube connecting the mouth and the stomach), stomach and intestines, including ulcers. If you have trouble or pain upon swallowing, heartburn, chest pain and black or bloody stools, stop taking ACTONEL DR and tell your doctor right away. Other side effects of ACTONEL DR include:

Most common side effects:
- Abdominal pain, heartburn, nausea

Other common side effects:
- Pain in bones, joints, or muscles, rarely severe. The pain may start as soon as one day or up to several months after starting ACTONEL DR.
- Diarrhea, constipation, inflammation of the pharynx and/or nose, upper respiratory tract infection.

Very Rare Side Effects:
- Non-healing jaw. Talk to your doctor if you experience persistent pain in your mouth, teeth or jaw, or if your gums or mouth heal poorly.
- Unusual fractures in the thigh bone. Talk to your doctor if you experience new or unusual pain in your hip, groin or thigh.

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**HOW TO STORE IT**

- Keep ACTONEL DR and all other medications out of the reach of children.
- Keep the tablets in their original package and store at controlled room temperature (20°-25°C).
- Do not keep medicine that is out of date or that you no longer need.
REPORTING SUSPECTED SIDE EFFECTS
You can report any suspected adverse reactions associated with the use of health products to the Canada Vigilance Program by one of the following 3 ways:

- Report online at www.healthcanada.gc.ca/medeffect
- Call toll-free at 1-866-234-2345
- Complete a Canada Vigilance Reporting Form and:
  - Fax toll-free to 1-866-678-6789, or
  - Mail to: Canada Vigilance Program
    Health Canada
    Postal Locator 0701D
    Ottawa, Ontario K1A 0K9

Postage paid labels, Canada Vigilance Reporting Form and the adverse reaction reporting guidelines are available on the MedEffect™ Canada Web site at www.healthcanada.gc.ca/medeffect.

NOTE: Should you require information related to the management of side effects, contact your health professional. The Canada Vigilance Program does not provide medical advice.

MORE INFORMATION
For more information, please visit [http://www.actonel.ca](http://www.actonel.ca).

This document plus the full product monograph, prepared for health professionals is available by contacting Warner Chilcott Canada Co. at: 1-800-565-0814.

This leaflet was prepared by Warner Chilcott Canada Co.

Last Revised: July 15, 2011
PART III: CONSUMER INFORMATION

ACTONEL® 75 mg

Risedronate Sodium

This leaflet is Part III of a three-part “Product Monograph” for ACTONEL. It is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about ACTONEL. Contact your doctor or pharmacist if you have any questions about the drug.

ABOUT THIS MEDICATION

What the medication is used for:

Treatment of osteoporosis in postmenopausal women.

What it does:

ACTONEL is a bisphosphonate drug that helps to slow bone loss. In many people, ACTONEL helps to increase bone density. In osteoporosis, the body removes more bone than it replaces. This causes bones to get weaker and more likely to break or fracture (usually at the spine, wrist or hip). Spine fractures may result in a curved back, height loss or back pain. ACTONEL corrects this imbalance by decreasing the elevated rate of bone removal. ACTONEL can therefore help reduce the risk of spine and non-spine fractures.

Your doctor may measure the thickness (i.e., density) of your bone through a bone mineral density (BMD) test or x-ray to check your progress against further bone loss or fracture. ACTONEL is not a pain reliever. Your doctor may prescribe or recommend another medicine specifically for pain relief.

When it should not be used:

• If you have low blood calcium levels (hypocalcemia).
• If you are allergic to risedronate sodium or any other ingredients in ACTONEL.

What the medicinal ingredient is:

Risedronate sodium

What the nonmedicinal ingredients are:

Crospovidone, ferric oxide, hydroxypropyl cellulose, hypromellose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, silicon dioxide and titanium dioxide.

What dosage form it comes in:

Each ACTONEL 75 mg pink tablet contains risedronate sodium 75 mg.

WARNINGS AND PRECAUTIONS

Before you use ACTONEL, talk to your doctor or pharmacist if:

• You have had problems or disease in your kidneys, esophagus (the tube connecting the mouth and the stomach), stomach or intestines.
• You cannot carry out dosing instructions (see PROPER USE OF THIS MEDICATION).
• You are pregnant or nursing.
• You have one of the following risk factors: cancer, chemotherapy, radiotherapy of the head or neck, treatment with corticosteroids or dental problems or dental infections. If so, a dental examination and any necessary dental procedures should be considered before you start treatment with ACTONEL.

Calcium and vitamin D are also important for strong bones. Your doctor may ask you to take calcium and vitamin D while you are on ACTONEL therapy. (see INTERACTIONS WITH THIS MEDICATION section).

INTERACTIONS WITH THIS MEDICATION

If taken with some other medicines, the effects of ACTONEL or the effects of other medicines may be changed. It is important to tell your health care provider, including doctors and dentists, about all medications you are taking, even if the medicine does not require a prescription (including vitamin and herbal supplements).

You should not take ACTONEL with food, as it may prevent your body from absorbing or using ACTONEL. You should take ACTONEL on an empty stomach. (See PROPER USE OF THIS MEDICATION for instruction).

Vitamin and mineral supplements, as well as antacids, may contain substances (e.g., calcium, magnesium, aluminum, and iron) which can stop your body from absorbing or using ACTONEL. These should be taken at a different time of day.
PROPER USE OF THIS MEDICATION

As with all medications, it is important to take ACTONEL as directed by your doctor.

Usual Dose:

Treatment of postmenopausal osteoporosis:

Monthly Duet Dose

- ACTONEL 75 mg is a “monthly duet” given as 1 tablet per day for two consecutive days each month, on the same two calendar days each month.
- Choose 2 days in a row each month that you will remember and that best fit your schedule to take your ACTONEL 75 mg doses.
- Take the first ACTONEL 75 mg tablet of the monthly duet first thing in the morning with plain water before you have anything to eat or drink on your first chosen day.
- Take the second ACTONEL 75 mg tablet of the monthly duet in the morning of the following day with plain water before you have anything to eat or drink.
- ACTONEL 75 mg should be taken on the same two consecutive calendar days each month.

DOSING INSTRUCTIONS

- ACTONEL should be taken in the morning on an empty stomach at least 30 minutes before consuming the first food, drink (other than plain water) and/or any other medication of the day. Food, medication or drink other than plain water can interfere with the absorption of ACTONEL.
- Each ACTONEL tablet should be swallowed whole while you are in an upright position and with sufficient plain water (≥ 120 mL or ½ cup) to facilitate delivery to the stomach.
- You should not lie down for at least 30 minutes after taking the medication. You may sit, stand or do normal activities like read the newspaper, take a walk, etc.
- ACTONEL tablets should not be chewed, cut, or crushed.

These recommendations help ACTONEL work correctly and help you avoid possible irritation of the esophagus (the tube connecting the mouth and the stomach).

You should take ACTONEL for as long as your doctor recommends, to continue to prevent bone loss and protect your bones from fractures.

Missed Dose:

Monthly duet: If you miss one or both tablets of your dose of ACTONEL 75 mg in the morning, do not take it later in the day. If the next month’s scheduled doses are more than 7 days away do the following:

- If both tablets were missed, take the first ACTONEL 75 mg tablet on the morning after the day it is remembered and the second tablet on the next consecutive morning.
- If only one tablet is missed, take the missed tablet on the morning after the day it is remembered.

You should then continue your usual schedule of ACTONEL 75 mg on two consecutive days each month. Do not take more than two 75 mg tablets within 7 days.

If the next month’s scheduled doses are 1 to 7 days away, you should wait until next month’s scheduled doses and then resume taking ACTONEL 75 mg on two consecutive days each month as originally scheduled.

Overdose:

If you take too many tablets on any given day, contact your doctor, or a Poison Control Centre, or an emergency room of the nearest hospital immediately. Drink a full glass of milk. Do not induce vomiting.

SIDE EFFECTS AND WHAT TO DO ABOUT THEM

Drugs like ACTONEL may cause problems in your esophagus (the tube connecting the mouth and the stomach), stomach and intestines, including ulcers. If you have trouble or pain upon swallowing, heartburn, chest pain and black or bloody stools, stop taking ACTONEL and tell your doctor right away. Remember to take ACTONEL as directed.

In clinical studies of osteoporosis with ACTONEL, the most commonly reported side effects were abdominal pain, heartburn and nausea. In studies of Paget’s disease, diarrhea and headache were also commonly reported.

ACTONEL may cause pain in bones, joints or muscles, rarely severe. Pain may start as soon as one day or up to several months after starting ACTONEL.

ACTONEL at monthly doses may cause short-lasting, mild flu-like symptoms. These symptoms usually decrease after subsequent doses.

Very rarely patients have reported non-healing jaw wounds while receiving ACTONEL or other drugs in this class. Consult your doctor if you experience persistent pain in your mouth, teeth or jaw, or if your gums or mouth heal poorly.

Very rarely patients have reported unusual fractures in their thigh bone while receiving drugs in this class. Consult your doctor if you experience new or unusual pain in your hip, groin, or thigh.
### IMPORTANT SIDE EFFECTS, HOW OFTEN THEY HAPPEN AND WHAT TO DO ABOUT THEM

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<tbody>
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<td>Only if severe</td>
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*This is not a complete list of side effects. For any unexpected effects while taking ACTONEL, contact your doctor or pharmacist.*

### HOW TO STORE IT

- Keep ACTONEL and all other medications out of the reach of children.
- Keep the tablets in their original package and store at controlled room temperature (20 °-25°C).
- Do not keep medicine that is out of date or that you no longer need.

### REPORTING SUSPECTED SIDE EFFECTS

You can report any suspected adverse reactions associated with the use of health products to the Canada Vigilance Program by one of the following 3 ways:

- Report online at www.healthcanada.gc.ca/medeffect
- Call toll-free at 1-866-234-2345
- Complete a Canada Vigilance Reporting Form and:
  - Fax toll-free to 1-866-678-6789, or
  - Mail to: Canada Vigilance Program  
    Health Canada  
    Postal Locator 0701D  
    Ottawa, Ontario K1A 0K9

Postage paid labels, Canada Vigilance Reporting Form and the adverse reaction reporting guidelines are available on the MedEffect™ Canada Web site at www.healthcanada.gc.ca/medeffect.

*NOTE: Should you require information related to the management of side effects, contact your health professional. The Canada Vigilance Program does not provide medical advice.*

### MORE INFORMATION

For more information, please visit [http://www.actonel.ca](http://www.actonel.ca)

This document plus the full product monograph, prepared for health professionals is available by contacting Warner Chilcott Canada Co. at: 1-800-565-0814.

This leaflet was prepared by Warner Chilcott Canada Co.

Last Revised: July 15, 2011
PART III: CONSUMER INFORMATION

ACTONEL® 150 mg
Risedronate Sodium

This leaflet is Part III of a three-part “Product Monograph” published for ACTONEL. It is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about ACTONEL. Contact your doctor or pharmacist if you have any questions about the drug.

ABOUT THIS MEDICATION

What the medication is used for:
Treatment of osteoporosis in postmenopausal women.

What it does:
ACTONEL is a bisphosphonate drug that helps to slow bone loss. In many people, ACTONEL helps to increase bone density. In osteoporosis, the body removes more bone than it replaces. This causes bones to get weaker and more likely to break or fracture (usually at the spine, wrist or hip). Spine fractures may result in a curved back, height loss or back pain. ACTONEL corrects this imbalance by decreasing the elevated rate of bone removal. ACTONEL can therefore help reduce the risk of spine and non-spine fractures.

Your doctor may measure the thickness (i.e., density) of your bone through a bone mineral density (BMD) test or x-ray to check your progress against further bone loss or fracture.

ACTONEL is not a pain reliever. Your doctor may prescribe or recommend another medicine specifically for pain relief.

When it should not be used:

• If you have low blood calcium levels (hypocalcemia).
• If you are allergic to risedronate sodium or any other ingredients in ACTONEL.

What the medicinal ingredient is:
Risedronate sodium

What the nonmedicinal ingredients are:
Crospovidone, hydroxypropyl cellulose, hypromellose, indigo carmine, magnesium stearate, microcrystalline cellulose, polyethylene glycol, silicon dioxide and titanium dioxide.

What dosage form it comes in:
ACTONEL 150 mg is available as tablets. Each blue tablet contains risedronate sodium 150 mg.

WARNINGS AND PRECAUTIONS

Before you use ACTONEL, talk to your doctor or pharmacist if:

• You have had problems or disease in your kidneys, esophagus (i.e., the tube connecting the mouth and the stomach), stomach or intestines.
• You cannot carry out dosing instructions (see PROPER USE OF THIS MEDICATION).
• You are pregnant or nursing.
• You have one of the following risk factors: cancer, chemotherapy, radiotherapy of the head or neck, treatment with corticosteroids or dental problems or dental infections. If so, a dental examination and any necessary dental procedures should be considered before you start treatment with ACTONEL.

Calcium and vitamin D are also important for strong bones. Your doctor may ask you to take calcium and vitamin D while you are on ACTONEL therapy (see INTERACTIONS WITH THIS MEDICATION section).

INTERACTIONS WITH THIS MEDICATION

If taken with some other medicines, the effects of ACTONEL or the effects of other medicines may be changed. It is important to tell your health care providers, including doctors and dentists, about all medications you are taking, even if the medicine does not require a prescription (including vitamin and herbal supplements).

You should not take ACTONEL with food, as it may prevent your body from absorbing or using ACTONEL. You should take ACTONEL on an empty stomach. (See PROPER USE OF THIS MEDICATION for instruction).

Vitamin and mineral supplements, as well as antacids, may contain substances (e.g., calcium, magnesium, aluminum, and iron) which can stop your body from absorbing or using ACTONEL. These should be taken at a different time of day.

PROPER USE OF THIS MEDICATION

As with all medications, it is important to take ACTONEL as directed by your doctor.

Usual Dose:

Treatment of postmenopausal osteoporosis:

• Choose a day of the month that you will remember and that best fits your schedule to take your ACTONEL dose (1 tablet of 150 mg) once per month.
• On your chosen day each month, take 1 ACTONEL 150 mg tablet first thing in the morning with plain water before you have anything to eat or drink.
DOsing Instructions

- ACTONEL should be taken in the morning on an empty stomach at least 30 minutes before consuming the first food, drink (other than plain water) and/or any other medication of the day. Food, medication or drink other than plain water can interfere with the absorption of ACTONEL.
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- ACTONEL tablets should not be chewed, cut, or crushed.

These recommendations help ACTONEL work correctly and help you avoid possible irritation of the esophagus (the tube connecting the mouth and the stomach).

You should take ACTONEL for as long as your doctor recommends, to continue to prevent bone loss and protect your bones from fractures.

Missed Dose:
If you miss a dose of ACTONEL 150 mg in the morning, do not take it later in the day. If the next month’s scheduled dose is more than 7 days away, take the missed tablet on the morning after the day it is remembered. You should then return to your usual schedule of ACTONEL 150 mg.

If the next month’s scheduled dose is 1 to 7 days away, you should wait until next month’s scheduled dose and then resume taking ACTONEL 150 mg as originally scheduled. Do not take more than 150 mg of ACTONEL within 7 days.

Overdose:
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SIDE EFFECTS AND WHAT TO DO ABOUT THEM
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