

- a cycle of worsening systemic hypercalcemia. Reducing excessive bone resorption and maintaining adequate fluid administration are, therefore, essential to the management of hypercalcemia of malignancy. Patients who have hypercalcemia of malignancy can generally be divided into two groups according to the pathophysiologic mechanism involved: humoral hypercalcemia and hypercalcemia due to tumor invasion of bone. In humoral hypercalcemia, osteoclasts are activated and
- bone resorption is stimulated by factors such as parathyroid-hormone-related protein, which are elaborated by the tumor and circulate systemically. Humoral hypercalcemia usually occurs in squamous-cell malignancies of the lung or head and neck or in genitourinary tumors such as renal-cell carcinoma or ovarian cancer. Skeletal metastases may be absent or minimal in these patients

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Head and Neck Genitourinary Malignant Melanoma Hepatobiliary Thyroid Other Sarcoma Neuroendocrine/Carcinoid Mesothelioma

14

Description: Zometa	Rev. Date:	April 2005	LD&C:	Date:		
Material Group No .:	Component No.:	Supersedes Component No.:				
USLFLTH	• 5000349	• 5000227	LD&C:	_ Date:		
•	•	•				
•	•	•	Engineer:	_ Date:		
Dimensions: 17 x 15	T2005-26	T2004-86				
Number of Colors: 1 Black						
Do Not Print Dotted Lines	FPO - For Positio		U NOVARTIS			

Unknown

GI (Other)

Bladder

childbearing potential should be advised to avoid becoming pregnant. PRECAUTIONS General Standard hypercalcemia-related metabolic parameters, such as serum levels of calcium, phosphate, and magnesium, as well as serum creatinine, should be carefully monitored following nitiation of therapy with Zometa® (zoledronic acid) Injection. If hypocalcemia, hypophosphatemia, or hypomagnesemia occur, short-term supplemental therapy may be necessary.

Patients with hypercalcemia of malignancy must be adequately rehydrated prior to administration of Zometa. Loop diuretics should not be used until the patient is adequately rehydrated and should be used with caution in combination with Zometa in order to avoid hypocalcemia. Zometa should be used with caution with other nephrotoxic drugs.

Renal Insufficiency Limited clinical data are available regarding use of Zometa in patients with renal impairment. teta is excreted intact primarily via the kidney, and the risk of adverse reactions, in particular renal adverse reactions, may be greater in patients with impaired renal function. Serum creatinine should be monitored in all patients treated with Zometa prior to each dose. Studies of Zometa in the treatment of hypercalcemia of malignancy excluded patients with serum creatinine \geq 400 µmol/L or \geq 4.5 mg/dL. Bone metastasis trials excluded patients with serum creatinine >265 µmol/L or >3.0 mg/dL and there were only eight of 564 patients treated with Zometa 4 mg by 15-minute infusion with a baseline serum creatinine >2 mg/dL. No clinical or pharmacokinetics data are available to guide dose selection or to provide guidance on how to safely use Zometa in patients with severe renal impairment. For multiple myeloma and bone metastases of solid tumors, the use of Zometa in patients with severe renal impairment is not recommended. For hypercalcemia of malignancy. Zometa should be used in patients with severe renal impairment only if the expected clinical benefits outweigh the risk of renal failure and after considering other available treatment options. (See WARNINGS.) Dose adjustments of Zometa are not necessary in treating patients for hypercalcemia presenting with mild-to-moderate rena impairment prior to initiation of therapy (serum creatinine <400 µmol/L or <4.5 mg/dL). II. Analysis of Time to the First SRE Patients receiving Zometa for hypercalcemia of malignancy with evidence of deterioration in renal function should be appropriately evaluated and consideration should be given as to whether the potential benefit of continued treatment with Zometa outweighs the possible risk. Upon initiation of treatment in patients with multiple myeloma or metastatic bone lesions from solid tumors, with mild-to-moderate renal impairment, lower doses of Zometa are recommended. In patients who show evidence of renal deterioration during treatment, Zometa should only be 0.011 0.67 (0.49, 0.91) resumed when serum creatinine returns to within 10% of baseline. (See WARNINGS and DOSAGE AND ADMINISTRATION.) Hepatic Insufficiency Only limited clinical data are available for use of Zometa to treat hypercalcemia of malignancy in patients with hepatic insufficiency, and these data are not adequate to provide guidance on dosage selection or how to safely use Zometa in these patients. 0.023 Patients with Asthma (0.55, 0.96) While not observed in clinical trials with Zometa, administration of other bisphosphonates has been associated with bronchoconstriction in aspirin-sensitive asthmatic patients. Zometa should be used with caution in patients with aspirin-sensitive asthma Osteonecrosis of the Jaw Osteonecrosis of the jaw (ONJ) has been reported in patients with cancer receiving treatment regimens including bisphosphonates. Many of these patients were also receiving chemotherapy and corticosteroids. The majority of reported cases have been associated with dental procedures such as tooth extraction. Many had signs of local infection including osteomyelitis. A dental examination with appropriate preventive dentistry should be considered prior to treatment with bisphosphonates in patients with concomitant risk factors (e.g., cancer, chemotherapy, corticosteroids, poor oral hygiene). While on treatment, these patients should avoid invasive dental procedures if possible. For patients who develop ONJ while on bisphosphonate therapy, dental surgery may exacerbate the condition. For patients requiring dental procedures, there are no data avail able to suggest whether discontinuation of bisphosphonate treatment reduces the risk of ONJ. Clinical judgment of the treating physician should guide the management plan of each patient based on individual II. Analysis of Time to the First SRE benefit/risk assessmen Musculoskeletal Pain In post-marketing experience, severe and occasionally incapacitating bone, joint, and/or muscle pain has been reported in patients taking bisphosphonates. However, such reports have been P-value infrequent. This category of drugs includes Zometa (zoledronic acid for injection). 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. A subset had recurrence of symptoms when rechallenged 0.32 (0.77, 1.09) with the same drug or another bisphosphonate. Laboratory Tests Serum creatinine should be monitored prior to each dose of Zometa. Serum calcium, electrolytes, phosphate, magnesium, and hematocrit/hemoglobin should also be monitored regularly. (See WARNINGS, PRECAUTIONS, DOSAGE AND ADMINISTRATION, and ADVERSE REACTIONS.) **Drug Interactions** In vitro studies indicate that zoledronic acid is approximately 22% bound to plasma proteins. In vitro studies also indicate that zoledronic acid does not inhibit microsomal CYP450 enzymes. In vivo studies showed that zoledronic acid is not metabolized, and is excreted into the urine as the intact drug. However, no in vivo drug interaction studies have been performed. Caution is advised when bisphosphonates are administered with aminoglycosides, since these agents may have an additive effect to lower serum calcium level for prolonged periods. This has not been reported in Zometa clinical trials. Caution should also be exercised when Zometa is used in combination with loop diuretics due to an increased risk of hypocalcemia. Caution is indicated when Zometa is used with other potentially nephrotoxic drugs. In multiple myeloma patients, the risk of renal dysfunction may be increased when Zometa is used in combination with thalidomide. Carcinogenesis, Mutagenesis, Impairment of Fertility Carcinogenesis: Standard lifetime carcinogenicity bioassays were conducted in mice and rats. Mice were given oral doses of zoledronic acid of 0.1, 0.5, or 2.0 mg/kg/day. There was an increased incidence of Harderian gland adenomas in males and females in all treatment groups (at doses ≥0.002 times a human intravenous dose of 4 mg, based on a comparison of relative body surface areas). Rats were given oral doses of zoledronic acid of 0.1, 0.5, or 2.0 mg/kg/day. No increased incidence of tumors was observed (at doses ≤0.2 times the human intravenous dose of 4 mg, based on a comparison of relative body surface areas). Mutagenesis: Zoledronic acid was not genotoxic in the Ames bacterial mutagenicity assay, in the Chinese hamster ovary cell assay, or in the Chinese hamster gene mutation assay, with or without metabolic activation. Zoledronic acid was not genotoxic in the in vivo rat micronucleus assay. Impairment of Fertility: Female rats were given subcutaneous doses of zoledronic acid of 0.01, 0.03, or 0.1 mg/kg/day beginning 15 days before mating and continuing through gestation. Effects observed in the high-dose group (with systemic exposure of 1.2 times the human systemic exposure following an intravenous dose of 4 mg, based on AUC comparison) included inhibition of ovulation and a decrease in the number of pregnant rats. Effects observed in both the mid-dose group (with systemic exposure of 0.2 times the human systemic exposure following an intravenous dose of 4 mg, based on an AUC comparison) and high-dose group included an increase in preimplantation losses and a decrease in the number of implantations and live fetuses. Pregnancy Category D (See WARNINGS.) osphonates are incorporated into the bone matrix, from where they are gradually released over periods of weeks to years. The extent of bisphosphonate incorporation into adult bone, and hence, the amount available for release back into the systemic circulation, is directly related to the total dose and duration of bisphosphonate use. Although there are no data on fetal risk in humans, bisphosphonates do cause fetal harm in animals, and animal data suggest that uptake _ _ _ _ _ Zometa® (zoledronic acid) Injection Zometa[®] taking this drug, the patient should be apprised of the potential harm to the fetus. Women of (zoledronic acid) Injection

Hazard

Hazard

0.92

Median Ratio

373

363

0.46

Median Ratio

NR

321

230

163

0.02

0.13

Zometa[®] (zoledronic acid) Injection

Table 7 provides adverse events that were reported by 10% or more of the 189 patients treated with Zometa 4 mg or pamidronate 90 mg from the two controlled multicenter HCM trials. Adverse events are listed regardless of presumed causality to study drug.

Pamidronate

90 mg

n (%)

103 (100)

95 (92.2)

34 (33.0)

21 (20.4)

28 (27.2)

15 (14.6)

Table 7: Percentage of Patients with Adverse Events ≥10% Reported in Hypercalcemia of Malignancy Clinical Trials by Body System

Zometa®

4 mg

n (%)

86 (100)

81 (94.2)

38 (44.2)

14 (16.3)

25 (29.1)

Patients Studied Total No. of Patients Studied

Body as a Whole

Fever

Digestive

Nausea

Total No. of Patients with any AE

Progression of Cancer

Psychiatric Depression 146 (14) 112 (11) Anxiety Confusior 74 (7) Respiratory 282 (27) Dvspnea 224 (22) Cough Skin Alopecia 125 (12) Dermatitis 114 (11)

of bisphosphonates into fetal bone is greater than into maternal bone. Therefore, there is a theoretical risk of fetal harm (e.g., skeletal and other abnormalities) 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 established. In female rats given subcutaneous doses of zoledronic acid of 0.01, 0.03, or 0.1 mg/kg/day

beginning 15 days before mating and continuing through gestation, the number of stillbirths was increased and survival of neonates was decreased in the mid- and high-dose groups (≥0.2 times he human systemic exposure following an intravenous dose of 4 mg, based on an AUC compari son). Adverse maternal effects were observed in all dose groups (with a systemic exposure of \geq 0.07 times the human systemic exposure following an intravenous dose of 4 mg, based on an AUC comparison) and included dystocia and periparturient mortality in pregnant rats allowed to deliver. Maternal mortality may have been related to drug-induced inhibition of skeletal calcium

mobilization, resulting in periparturient hypocalcemia. This appears to be a bisphosphonate-class effect. In pregnant rats given a subcutaneous dose of zoledronic acid of 0.1, 0.2, or 0.4 mg/kg/day during gestation, adverse fetal effects were observed in the mid- and high-dose groups (with sys-

temic exposures of 2.4 and 4.8 times, respectively, the human systemic exposure following an intravenous dose of 4 mg, based on an AUC comparison). These adverse effects included increases in pre- and post-implantation losses, decreases in viable fetuses, and fetal skeletal, vis-ceral, and external malformations. Fetal skeletal effects observed in the high-dose group included

unossified or incompletely ossified bones, thickened, curved or shortened bones, wavy ribs, and shortened jaw. Other adverse fetal effects observed in the high-dose group included reduced lens udimentary cerebellum, reduction or absence of liver lobes, reduction of lung lobes, vessel dilation cleft palate and edema. Skeletal variations were also observed in the low-dose group (with systemic exposure of 1.2 times the human systemic exposure following an intravenous dose of 4 mg, based on an AUC comparison). Signs of maternal toxicity were observed in the high-dose group and included reduced body weights and food consumption, indicating that maximal exposure

levels were achieved in this study. In pregnant rabbits given subcutaneous doses of zoledronic acid of 0.01, 0.03, or 0.1 mg/kg/day during gestation (≤0.5 times the human intravenous dose of 4 mg, based on a comparison of relative body surface areas), no adverse fetal effects were observed. Maternal mortality and abortion

occurred in all treatment groups (at doses ≥0.05 times the human intravenous dose of 4 mg, based on a comparison of relative body surface areas). Adverse maternal effects were associated with, and may have been caused by, drug-induced hypocalcemia. Nursing Mothers

It is not known whether Zometa is excreted in human milk. Because many drugs are excreted in human milk, and because Zometa binds to bone long term, Zometa should not be administered to a nursing woman.

Pediatric Use The safety and effectiveness of Zometa in pediatric patients have not been established. Because of long-term retention in bone, Zometa should only be used in children if the potential benefit outweighs the potential risk.

Geriatric Use Clinical studies of Zometa in hypercalcemia of malignancy included 34 patients who were 65

years of age or older. No significant differences in response rate or adverse reactions were seen in geriatric patients receiving Zometa as compared to younger patients. Controlled clinical studies of Zometa in the treatment of multiple myeloma and bone metastases of solid tumors in patients over age 65 revealed similar efficacy and safety in older and younger patients. Because decreased

renal function occurs more commonly in the elderly, special care should be taken to monitor renal function.

ADVERSE REACTIONS Hypercalcemia of Malignancy

Adverse reactions to Zometa® (zoledronic acid) Injection are usually mild and transient and similar to those reported for other bisphosphonates. Intravenous administration has been most com monly associated with fever. Occasionally, patients experience a flu-like syndrome consisting of fever, chills, bone pain and/or arthralgias, and myalgias. Gastrointestinal reactions such as nausea and vomiting have been reported following intravenous infusion of Zometa. Local reactions at the infusion site, such as redness or swelling, were observed infrequently. In most cases, no spe-

cific treatment is required and the symptoms subside after 24-48 hours. Rare cases of rash, pruritus, and chest pain have been reported following treatment with Zometa.

As with other bisphosphonates, cases of conjunctivitis and hypomagnesemia have been

reported following treatment with Zometa. Grade 3 and Grade 4 laboratory abnormalities for serum creatinine, serum calcium, serum phosphorus, and serum magnesium observed in two clinical trials of Zometa in patients with HCM are shown in Table 6.

Table 6: Grade 3-4 Laboratory Abnormalities for Serum Creatinine, Serum Calcium. Serum Phosphorus, and Serum Magnesium in Two Clinical Trials in Patients

with HCM Grade 3 Grade 4

1	Laboratory Parameter		meta® ma	Pamidronate 90 mg		Zometa [®] 4 mg		Pamidronate 90 mg	
1		n/N	ັ (%)	n/N	ັ(%)	n/N	(%)	n/N	(%)
I.	Serum Creatinine ¹	2/86	(2.3%)	3/100	(3.0%)	0/86	_	1/100	(1.0%)
	Hypocalcemia ²	1/86	(1.2%)	2/100	(2.0%)	0/86	_	0/100	_
I.	Hypophosphatemia ³	36/70	(51.4%)	27/81	(33.3%)	1/70	(1.4%)	4/81	(4.9%)
	Hypomagnesemia ⁴	0/71		0/84		0/71		1/84	(1.2%)

¹ Grade 3 (>3x Upper Limit of Normal); Grade 4 (>6x Upper Limit of Normal) ² Grade 3 (<7 mg/dL); Grade 4 (<6 mg/dL)

³ Grade 3 (<2 mg/dL); Grade 4 (<1 mg/dL) ⁴ Grade 3 (<0.8 mEg/L); Grade 4 (<0.5 mEg/L)

_ _ _ _ _

Constipation	23	(26.7)	13	(12.6)	
Diarrhea	15	(17.4)	17	(16.5)	
Abdominal Pain	14	(16.3)	13	(12.6)	
Vomiting	12	(14.0)	17	(16.5)	
Anorexia	8	(9.3)	14	(13.6)	
Cardiovascular					
Hypotension	9	(10.5)	2	(1.9)	
Hemic and Lymphatic System					
Anemia	19	(22.1)	18	(17.5)	
Infections					
Moniliasis	10	(11.6)	4	(3.9)	
Laboratory Abnormalities					
Hypophosphatemia	11	(12.8)	2	(1.9)	
Hypokalemia	10	(11.6)	16	(15.5)	
Hypomagnesemia	9	(10.5)	5	(4.9)	
Musculoskeletal					
Skeletal Pain	10	(11.6)	10	(9.7)	
Nervous					
Insomnia	13	(15.1)	10	(9.7)	
Anxiety	12	(14.0)	8	(7.8)	
Confusion	11	(12.8)	13	(12.6)	
Agitation	11	(12.8)	8	(7.8)	
Respiratory					
Dyspnea	19	(22.1)	20	(19.4)	
Coughing	10	(11.6)	12	(11.7)	
Urogenital					

The following adverse events from the two controlled multicenter HCM trials (n=189) were reported by a greater percentage of patients treated with Zometa 4 mg than with pamidronate 90 mg and occurred with a frequency of greater than or equal to 5% but less than 10%. Adverse events are listed regardless of presumed causality to study drug.

12 (14.0)

Body as a Whole: asthenia, chest pain, leg edema, mucositis, and metastases

Digestive System: dysphagia Hemic and Lymphatic System: granulocytopenia, thrombocytopenia, and pancytopenia

Infection: non-specific infection

Laboratory Abnormalities: hypocalcemia Metabolic and Nutritional: dehydration

Musculoskeletal: arthralgias Nervous System: headache, somnolence

Urinary Tract Infection

Respiratory System: pleural effusion

NOTE: In the HCM clinical trials, pamidronate 90 mg was given as a 2-hour intravenous infusion. The relative safety of pamidronate 90 mg given as a 2-hour intravenous infusion compared to the same dose given as a 24-hour intravenous infusion has not been adequately studied in controlled clinical trials.

Multiple Myeloma and Bone Metastases of Solid Tumors

The safety analysis includes patients treated in the core and extension phases of the trials. The analysis includes the 2,042 patients treated with Zometa 4 mg, pamidronate 90 mg or placebo in the three controlled multicenter bone metastases trials, including 969 patients completing the efficacy phase of the trial, and 619 patients that continued in the safety extension phase. Only 347 patients completed the extension phases and were followed for two years (or 21 months for the other solid tumor patients). The median duration of exposure for safety analysis for Zometa 4 mg (core plus extension phases) was 12.8 months for breast cancer and multiple myeloma, 10.8 months for prostate cancer, and 4.0 months for other solid tumors.

Table 8 describes adverse events that were reported by $\geq 10\%$ of patients. Adverse events are listed regardless of presumed causality to study drug.

Table 8: Percentage of Patients with Adverse Events ≥10% Reported in Three Bone

Metastases Clinical Trials by Body System							
	Zom 4 n n (*	ng	Pamidro 90 m n (%	ng	Place n (ebo (%)	
Patients Studied							
Total No. of Patients	1031	(100)	556	(100)	455	(100)	
Total No. of Patients with any AE	1015	(98)	548	(99)	445	(98)	
Blood and Lymphatic		``'		. ,		```	
Anemia	344	(33)	175	(32)	128	(28)	
Neutropenia	124	(12)	83	(15)	35	(8)	
Thrombocytopenia	102	(10)	53	(10)	20	(4)	
Gastrointestinal		``'		. ,		. ,	
Nausea	476	(46)	266	(48)	171	(38)	
Vomiting	333	(32)	183	(33)	122	(27)	
Constipation	320	(31)	162	(29)	174	(38)	
Diarrhea	249	(24)	162	(29)	83	(18)	
Abdominal Pain	143	(14)	81	(15)	48	(11)	
Dyspepsia	105	(10)	74	(13)	31	(7)	
Stomatitis	86	(8)	65	(12)	14	(3)	
Sore Throat	82	(8)	61	(11)	17	(4)	
General Disorders and Administration	Site	(-)		()		()	
Fatigue	398	(39)	240	(43)	130	(29)	
Pyrexia	328	(32)	172	(31)	89	(20)	
Weakness	252	(24)	108	(19)	114	(25)	
Edema Lower Limb	215	(21)	126	(23)	84	(19)	
Rigors	112	(11)	62	(11)	28	(6)	
Infections		()		```		(-)	
Urinary Tract Infection	124	(12)	50	(9)	41	(9)	
Upper Respiratory Tract Infection	101	(10)	82	(15)	30	(7)	
Metabolism		``'		. ,		. ,	
Anorexia	231	(22)	81	(15)	105	(23)	
Weight Decreased	164	(16)	50	(9)	61	(13)	
Dehydration	145	(14)	60	(11)	59	(13)	
Appetite Decreased	130	(13)	48	(9)	45	(10)	
Musculoskeletal		``'		. ,		```	
Bone Pain	569	(55)	316	(57)	284	(62)	
Myalgia	239	(23)	143	(26)	74	(16)	
Arthralgia	216	(21)	131	(24)	73	(16)	
Back Pain	156	(15)	106	(19)	40	(9)	
Pain in Limb	143	(14)	84	(15)	52	(11)	
Neoplasms		``'		. ,		```	
Malignant Neoplasm Aggravated	205	(20)	97	(17)	89	(20)	
Nervous		. ,		. ,		```	
Headache	191	(19)	149	(27)	50	(11)	
Dizziness (excluding vertigo)	180	(18)	91	(16)	58	(13)	
Insomnia	166	(16)	111	(20)	73	(16)	
Paresthesia	149	(15)	85	(15)	35	(8)	
Hypoesthesia	127	(12)	65	(12)	43	(10)	
21 1 1 1 1 1 1		, .,		• •	-	(-)	

74 (1 Grade 3 and Grade 4 laboratory abnormalities for serum creatinin phosphorus, and serum magnesium observed in three clinical trials of bone metastases are shown in Tables 9 and 10. Table 9: Grade 3 Laboratory Abnormalities for Serum Creatinin Phosphorus, and Serum Magnesium in Three Clinical 1 Grade 3 Laboratory Parameter Zometa 90 mg 4 mg n/N (%) n/N (%) Serum Creatinine 7/529 (1.3%) 4/268 (1.5%)

Hypophosphatemia³ 115/973 (11.8%) 38/537 (7.1%) 19/971 (2.0%) 2/535 (0.4%) Hypermagnesemia⁴ Hypomagnesemia⁵ 1/971 (0.1%) 0/535 — ¹ Grade 3 (>3x Upper Limit of Normal); Grade 4 (>6x Upper Limit of Normal) Serum creatinine data for all patients randomized after the 15-minute infu ² Grade 3 (<7 mg/dL); Grade 4 (<6 mg/dL) ³ Grade 3 (<2 mg/dL); Grade 4 (<1 mg/dL)</p> ⁴ Grade 3 (>3 mEq/L); Grade 4 (>8 mEq/L) ⁵ Grade 3 (<0.9 mEq/L); Grade 4 (<0.7 mEq/L)

6/973 (0.6%)

4/536 (0.7%)

Hypocalcemia

Table 10: Grade 4 Laboratory Abnormalities for Serum Creatini Phosphorus, and Serum Magnesium in Three Clinical Bone Metastases

Laboratory Parameter	Zometa [®] 4 mg			ronate mg
	n/N	(%)	n/N	(%)
Serum Creatinine ^{1*}	2/529	(0.4%)	1/268	(0.4%)
Hypocalcemia ²	7/973	(0.7%)	3/536	(0.6%)
Hypophosphatemia ³	5/973	(0.5%)	0/537	_
Hypermagnesemia ⁴	0/971	_	0/535	_
Hypomagnesemia ⁵	2/971	(0.2%)	1/535	(0.2%)
¹ Grade 3 (>3x Upper Lin [*] Serum creatinine data fr ² Grade 3 (<7 mg/dL); Gr	or all pat ade 4 (<	ients rando 6 mg/dL)		

³ Grade 3 (<2 mg/dL); Grade 4 (<1 mg/dL)</p> ⁴ Grade 3 (>3 mEq/L); Grade 4 (>8 mEq/L) ⁵ Grade 3 (<0.9 mEq/L); Grade 4 (<0.7 mEq/L)

Among the less frequently occurring adverse events (<15% of pat influenza-like illness, and hypocalcemia showed a trend for more eve administration (Zometa 4 mg and pamidronate groups) compared to Less common adverse events reported more often with Zometa 4 ed decreased weight, which was reported in 16% of patients in the Zo

with 9% in the pamidronate group. Decreased appetite was reported the Zometa 4-mg group (13%) compared with the pamidronate (9%) a but the clinical significance of these small differences is not clear. Renal Toxicity

In the bone metastases trials, renal deterioration was defined as an ir patients with normal baseline creatinine (<1.4 mg/dL) or an increase an abnormal baseline creatinine (≥1.4 mg/dL). The following are data deterioration in patients receiving Zometa 4 mg over 15 minutes in the

Table 11: Percentage of Patients with Renal Function Deteriora Randomized Following the 15-Minute Infusion Amend Patient Population/Baseline Creatinine

Multiple Myeloma	

Zome	ta [®] 4 mg	Pamidro
<u>n/N</u>	(<u>%)</u>	<u>n/N</u>
2/26	(7.7%)	23/246 2/22 25/268
Zome	ta® 4 mg	Pla
<u>n/N</u>	<u>(%)</u>	<u>n/N</u>
1/11	`(9.1%)	10/143 1/20 11/163
Zome	ta® 4 mg	Pla
<u>n/N</u>	<u>(%)</u>	<u>n/N</u>
12/82	(14.6%)	8/68
4/10	(40%)	2/10
16/92	(17.4%)	10/78
	n/N 27/246 2/26 29/272 Zome n/N 17/154 1/11 18/165 Zome n/N 12/82 4/10	27/246 (11%) 2/26 (7.7%) 29/272 (10.7%) Zometa® 4 mg n/N 17/154 (11%) 1/11 (9.1%) 18/165 (10.9%) Zometa® 4 mg n/N 18/165 (10.9%) 12/82 (14.6%) 4/10 (40%)

The risk of deterioration in renal function appeared to be related to patients were receiving Zometa (4 mg over 15 minutes), placebo, or Evaluation of serum creatinine is recommended prior to each cycle patients receiving Zometa for multiple myeloma and bone metastases vidence of deter in renal function, Zometa trea tinine returns to within 10% of baseline.

In the trials and in post-marketing experience, renal deterioration, and dialysis have occurred in patients with normal and abnormal bas patients treated with 4 mg infused over a 15-minute period. There has occurring after the initial Zometa dose.

eting Experier

Cases of osteonecrosis (primarily involving the jaws) have been repo bisphosphonates. The majority of the reported cases are in cancer pa procedure. Osteonecrosis of the jaws has multiple well-documented r nosis of cancer, concomitant therapies (e.g., chemotherapy, radiother morbid conditions (e.g., anemia, coagulopathies, infection, pre-existin causality cannot be determined, it is prudent to avoid dental surgery longed. (See PRECAUTIONS.) Cases of uveitis and episcleritis have been reported during post-

OVERDOSAGE

known.

There is no experience of acute overdose with Zometa® (zoledronic a received Zometa 32 mg over 5 minutes in clinical trials. Neither patient laboratory toxicity. Overdosage may cause clinically significant hypocalce and hypomagnesemia. Clinically relevant reductions in serum levels of magnesium should be corrected by intravenous administration of calc sodium phosphate, and magnesium sulfate, respectively. In an open-label study of zoledronic acid 4 mg in breast cancer pa received a single 48-mg dose of zoledronic acid in error. Two days af experienced a single episode of hyperthermia (38°C), which resolved a evaluations were normal, and the patient was discharged seven days A patient with non-Hodgkin's lymphoma received zoledronic acid 4 mg daily on four successive days for a total dose of 16 mg. The patient developed paresthesia and abnormal liver function tests with increased GGT (nearly 100U/L, each value unknown). The outcome of this case is not

· ·				
73 (1	17) 49 (11) 13) 37 (8) (7) 47 (10)	minutes has been shown to incr istered as a 15-minute intraveno	ministration of Zometa 4 mg as an i ease the risk of renal toxicity compa us infusion. In controlled clinical tria increased risk of renal toxicity comp	ared to the same dose admin- als, Zometa 8 mg has been
	28) 107 (24) 23) 65 (14)	patients with hypercalcemia of n	venous infusion, and was not assoc nalignancy. Single doses of Zomet enous infusion should be no less	a should not exceed 4 mg
80 (1	4) 36 (8)	WARNINGS.) In the trials and sion to renal failure and dialys	in post-marketing experience, rer sis, have occurred in patients, inc	nal deterioration, progres- luding those treated with
74 (1 n creatinin	3) 38 (8) ne, serum calcium, serum	the approved dose of 4 mg inf occurring after the initial Zom	used over 15 minutes. There have eta dose.	e been instances of this
	of Zometa in patients with	DOSAGE AND ADMINISTRATI Hypercalcemia of Malignancy		
	e, Serum Calcium, Serum Trials in Patients with Bone	hypercalcemia when considering	o the severity of, as well as the sym use of Zometa® (zoledronic acid) Ir nt to treat mild, asymptomatic hyper	njection. Vigorous saline
-		The maximum recommended serum calcium* ≥12 mg/dL [3.0	dose of Zometa in hypercalcemia of mmol/L]) is 4 mg. The 4-mg dose m	malignancy (albumin-corrected
F	Placebo	intravenous infusion over no les Patients should be adequatel and PRECAUTIONS.)	s than 15 minutes. y rehydrated prior to administration	of Zometa. (See WARNINGS
n/	N (%) 241 (1.7%)	Retreatment with Zometa 4 n	ng, may be considered if serum cald atment. It is recommended that a mi	
) 14/-	415 — 415 (3.4%)	monitored in all patients receiving	ull response to the initial dose. Rena og Zometa and possible deterioration ith Zometa. (See WARNINGS and F	n in renal function must be
1/-	(415 (1.9%) (415 (0.2%)		m (Cca, mg/dL) = Ca + 0.8 (mid-rate)	,
nit of Norma minute infu	ai) Jsion amendment	Multiple Myeloma and Metastat	tic Bone Lesions From Solid Tumo ta in patients with multiple myeloma a	
		from solid tumors for patients with than 15 minutes every three to fo	n creatinine clearance >60 mL/min is ur weeks. The optimal duration of the	4 mg infused over no less erapy is not known.
	ine, Serum Calcium, Serum I Trials in Patients with	tion (mild and moderate renal imp to achieve the same AUC as that	recommended Zometa doses for pati airment) are listed in the following tab achieved in patients with creatinine cl	le. These doses are calculated earance of 75 mL/min.
	2	PHARMACOLOGY, Special Popul	alculated using the Cockcroft-Gault f lations, Renal Insufficiency.)	ormula. (See CLINICAL
n	Placebo /N (%)	Baseline Creatinine Clearance > 60 50 - 60	e (mL/min) Zometa® Recomm 4.0 m 3.5 m	ng
2/4	241 — 415 (0.5%) 415 (0.2%)	40 - 49 30 - 39	3.3 m 3.0 m	ng
2/4	415 (0.2%) 415 (0.5%) 415 —		AUC of 0.66(mg•hr/L) (CrCl=75 mL/min) tinine should be measured before e	
it of Norma ninute infu	al) usion amendment		al deterioration. In the clinical studie	
		 For patients with normal base 	eline creatinine, increase of 0.5 mg/ aseline creatinine, increase of 1.0 m	
		In the clinical studies, Zometa	a treatment was resumed only wher 2. Zometa should be re-initiated at the	the creatinine returned to
more eve	tients), rigors, hypokalemia, ents with bisphosphonate	treatment interruption. Patients should also be admin	istered an oral calcium supplement of	of 500 mg and a multiple vita-
Zometa 4	the placebo group. mg than pamidronate includ- cometa 4-mg group compared	min containing 400 IU of Vitamin Preparation of Solution		Lallanda for the state of
reported	in slightly more patients in and placebo (10%) groups,	5 mL of concentrate (equivalent	ncentrate for infusion contain overfil to 4 mg zoledronic acid). This conce	entrate should immediately be
clear.			 Sodium Chloride, USP, or 5% Dexi syringe, to avoid inadvertent injection or no less than 15 minutes 	
increase	increase of 0.5 mg/dL for of 1.0 mg/dL for with		<i>ith Baseline CrCl</i> ≤60 <i>mL/min:</i> Wi	thdraw an appropriate volum
	a on the incidence of renal se trials. (See Table 11.)	4.4 mL for 3.5 mg dose 4.1 mL for 3.3 mg dose		
Deteriora n Amend	tion Who Were dment		nust be diluted in 100 mL of sterile (The dose must be given as a single	
		less than 15 minutes. If not used immediately after	dilution with infusion media, for mic	robiological integrity, the solu
amidrona <u>n/N</u>	te 90 mg (%)	ibrated to room temperature price	C-8°C (36°F-46°F). The refrigerated or to administration. The total time by ration must not exceed 24 hours.	
23/246	(9.3%)	Zometa must not be mixed Ringer's solution, and should	with calcium-containing infusion be administered as a single intra	
2/22 25/268	(9.1%) (9.3%)	separate from all other drugs. Method of Administration: Due	e to the risk of clinically significa	nt deterioration in renal
Plac <u>n/N</u>	ebo (<u>%)</u>	function, which may progress t and the duration of infusion s	o renal failure, single doses of Zor hould be no less than 15 minutes	meta should not exceed 4 mg s. (See WARNINGS.) In the
0/143	(7%)	and dialysis, have occurred in	patients, including those treated	with the approved dose of
1/20 1/163	(5%) (6.7%)	Zometa dose.	There have been instances of th ce to the intravenous administration	-
Plac <u>n/N</u>	ebo (%)	in order to decrease the risk of o	deterioration in renal function.	
8/68 2/10	(11.8%) (20%)		s should be inspected visually fo ion, whenever solution and conta	
0/78	(12.8%)		g zoledronic acid monohydrate, corre	
cebo, or p	o time on study, whether pamidronate.	citrate, USP.) mg of mannitol, USP, water for inje	-
netastase	e of therapy with Zometa. In s of solid tumors, who show be withheld until serum crea-		permitted to 15-30°C (59-86°F) [se	
	progression to renal failure	Temperature].		
ormal bas	seline renal function, including ve been instances of this	REV: APRIL 2005	Printed in U.S.A.	T2005-26 5000349
	orted in patients treated with			
cancer pa umented r	atients attendant to a dental risk factors including a diag-			
radiothei pre-existir	rapy, corticosteroids) and co- ng oral disease). Although	N. November		
• •	as recovery may be pro-	U novartis		
ng post-a	approval use.	Manufactured by Novartis Pharma Stein AG		
	acid) Injection. Two patients nt experienced any clinical or	Stein, Switzerland for Novartis Pharmaceuticals Corpo	pration	
hypocalce	emia, hypophosphatemia, of calcium, phosphorus, and	East Hanover, NJ 07936		
	cium gluconate, potassium or			
o days afl	atients, a female patient ter the overdose the patient			
even days	after treatment. All other after the overdose.			
onic acid 4	4 mg daily on four successive			