

Safety Data Sheet

BONIVA(R) Injection (3 mg/3 ml pre-filled syringes)

SECTION 1: Identification of the substance/mixture and of the company/undertaking

1.1. Product identifier

Product name	BONIVA(R) Injection (3 mg/3 ml pre-filled syringes)	
Product code	SAP-10067516	
Synonyms	- BONVIVA Prefilled Syringes 3 mg/3 ml - Boniva	*1

1.2. Relevant identified uses of the substance or mixture and uses advised against

Use	- pharmaceutical active substance (postmenopausal osteoporosis)	*1
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1.3. Details of the supplier of the safety data sheet

Company information	Enquiries: Genentech, Inc. 1 DNA Way South San Francisco USA-CA 94080 United States of America	Local representation:
	Phone 001-(650) 225-1000 E-Mail info.sds@roche.com US Chemtrec phone: (800)-424-9300	

1.4. Emergency telephone number

Emergency telephone number	US Chemtrec phone: (800)-424-9300
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*1 referring to: Ibandronate

SECTION 2: Hazards identification

Classification of the substance or mixture / Label elements

GHS Classification	no classification and labelling according to GHS
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Other hazards

Note	- no information available
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SECTION 3: Composition/information on ingredients

Characterization lbandronate with other inactive ingredients

Ingredients	Concentration	GHS-Classification (pure ingredient)
lbandronate 138926-19-9	0.1 %	<ul style="list-style-type: none">- Combustible dust (No category), USH003- Acute toxicity (Category 4), H302- Skin corrosion/irritation (Category 1B), H314- Specific target organ toxicity - Repeated exposure (Category 2), H373

For the full text of the 'Hazard statements' mentioned in this Section, see Section 16.

*1 referring to: lbandronate

SECTION 4: First aid measures

4.1. Description of first aid measures

Eye contact	- rinse immediately with tap water for at least 20 minutes - open eyelids forcibly
Skin contact	- drench affected skin with water
Inhalation	- in the event of symptoms get medical treatment

4.2. Most important symptoms and effects, both acute and delayed

Note - no information available

4.3. Indication of any immediate medical attention and special treatment needed

Note to physician - treat symptomatically

SECTION 5: Firefighting measures

5.1. Extinguishing media

Suitable extinguishing media	- adapt extinguishing media to surrounding fire conditions
Flash point (liquid)	not applicable

5.2. Special hazards arising from the substance or mixture

Specific hazards - no particular hazards known

5.3. Advice for firefighters

Protection of fire-fighters - precipitate gases/vapours/mists with water spray

SECTION 6: Accidental release measures

6.1. Personal precautions, protective equipment and emergency procedures

Personal precautions - no special precautions required

6.2. Environmental precautions

Environmental protection - no special environmental precautions required

6.3. Methods and material for containment and cleaning up

Methods for cleaning up - mop or flush the contaminated area with water

SECTION 7: Handling and storage

7.1. Precautions for safe handling

Suitable materials - glass

7.2. Conditions for safe storage, including any incompatibilities

Storage conditions - below 30 °C

Validity - after opening the content should be used within a short period, see expiry date on the label

Packaging materials - prefilled syringes

SECTION 8: Exposure controls/personal protection

8.1. Control parameters

Threshold value (Roche) air - IOEL (Internal Occupational Exposure Limit): 0.002 mg/m³ *1

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8.2. Exposure controls

Respiratory protection	<ul style="list-style-type: none">- Respiratory protection is recommended as a precaution to minimize exposure. Effective engineering controls are considered to be the primary means to control worker exposure. Respiratory protection should not substitute for feasible engineering controls.- respiratory protection not necessary during normal operations- in case of intense formation of aerosols: respirator with independent air supply or particle respectively filter mask (depending on the aerosol composition)
Hand protection	<ul style="list-style-type: none">- protective gloves (eg made of neoprene, nitrile or butyl rubber)
Eye protection	<ul style="list-style-type: none">- safety glasses

*1 referring to: lbandronate

SECTION 9: Physical and chemical properties

9.1. Information on basic physical and chemical properties

Color	colorless
Form	clear solution sterile liquid
Density	1.004 g/ml
pH value	3.9 to 4.1

9.2. Other information

Note - no information available

SECTION 10: Stability and reactivity

10.1. Reactivity

Note - no information available

10.2. Chemical stability

Stability - does not contain any antimicrobial preservative; therefore, care must be taken to ensure the sterility of the prepared solution

10.3. Possibility of hazardous reactions

Note - no information available

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10.4. Conditions to avoid

Conditions to avoid - temperatures above 30 °C

10.5. Incompatible materials

Note - no information available

10.6. Hazardous decomposition products

Note - Calcium containing solutions should not be mixed with Bondronat concentrate for solution for infusion

SECTION 11: Toxicological information

11.1. Information on toxicological effects

Acute toxicity	- LD ₅₀ 811 mg/kg (oral, rat)	*1
	- LD ₅₀ 30 mg/kg (i.v., rat)	*1
Local effects	- skin, eyes, mucous membranes: corrosive	*1
Sensitization	- non-sensitizing (guinea pig)	*1
Mutagenicity	- not mutagenic (various in vivo and in vitro test systems)	*1
Carcinogenicity	- not carcinogenic (several species)	*1
Reproductive toxicity	- not teratogenic, not embryotoxic (several species)	*1
	- does not lower parental fertility (several species)	*1
STOT-single exposure	- no information available	
STOT-repeated exposure	- no information available	
Aspiration hazard	- no information available	
Note	- inhibits mechanisms reducing bone mass by long-term binding to bone tissue	*1
	- high doses cause: liver damages, kidney damages	*1
	- decrease in serum calcium level possible	*1
	- dosage (oral): 2.5 to 50 mg/d	*1
	- dosage (i.v.): 0.5 mg/3 months to 2.5 mg/day	*1
Potential Health Effects	- Exposure: Inhalation, Ingestion, Skin contact, Eye contact	
	- Carcinogenicity: formulation not listed by NTP, IARC or OSHA	
Additional Health Information	- Conditions Aggravated: Hypersensitivity to this material and other materials in its chemical class. Uncorrected hypocalcemia. Severe renal impairment.	
*1 referring to:	ibandronate	

SECTION 12: Ecological information

12.1. Toxicity

Ecotoxicity	<ul style="list-style-type: none"> - strongly toxic for algae (<i>Selenastrum capricornutum</i>) EbC₅₀ (72 h) 1.4 mg/l ErC₅₀ (72 h) 4.7 mg/l NOEC (72 h) 0.22 mg/l (OECD No. 201) *1 - highly toxic for algae (<i>Scenedesmus (=Desmodesmus) subspicatus</i>) EbC₅₀ (72 h) 0.218 mg/l (nominal concentration) ErC₅₀ (72 h) 0.390 mg/l (nominal concentration) NOEC (72 h) < 0.1 mg/l (nominal concentration) (OECD No. 201) *1 - barely toxic for planktonic crustaceans (<i>Daphnia magna</i>) NOEC (48 h) 100 mg/l EC₅₀ (48 h) > 180 mg/l (OECD No. 202) *1 - no adverse influence on substrate biodegradation (activated sludge) concentration (28 d) 41.5 mg/l (OECD No. 301B, Modified Sturm Test) *1 - barely inhibitory on aerobic bacterial reproduction (activated sludge) NOEC (5 h) 1300 mg/l (growth test) *1 - barely toxic for fish (carp) LC₅₀ (96 h) 200 mg/l LC₀ (96 h) 86 mg/l (OECD No. 203) *1 - highly toxic for algae (<i>Scenedesmus (=Desmodesmus) subspicatus</i>) EC₅₀ (14 d) 0.5 mg/l (nominal concentration) NOEC (14 d) 0.1 mg/l (nominal concentration) (OECD No. 201) *1 - no adverse influence on substrate biodegradation concentration (28 d) 100 mg/l (Manometric Respirometry Test, OECD No. 301 F) *1
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12.2. Persistence and degradability

Ready biodegradability	<ul style="list-style-type: none"> - not readily biodegradable ≤ 3 %, 28 d (CO₂ Evolution Test, Modified Sturm Test, OECD No. 301B) *1 - not readily biodegradable 0 %, 28 d (Manometric Respirometry Test, OECD No. 301 F) *1
Inherent biodegradability	<ul style="list-style-type: none"> - not inherently biodegradable < 10 %, 1 d < 10 %, 15 d < 10 %, 28 d (Zahn-Wellens test, OECD No. 302 B) *1 - not inherently biodegradable < 10 %, 28 d (Zahn-Wellens test, OECD No. 302 B) *1

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Abiotic degradation - stable in water, no photodegradation 200 mg/l, water
< 2 %, 14 d, ~ 22 °C, under illumination *1

12.3. Bioaccumulative potential

Note - no information available

12.4. Mobility in soil

Note - no information available

Mobility - no significant adsorption (, 28 d, ~22 °C)
 $K_d = 1210 \text{ l/kg}$ (activated sludge)
(Adsorption to activated sludge in biodegradability test) *1

12.5. Results of PBT and vPvB assessment

Note - no information available

12.6. Other adverse effects

Note - no information available

Note - after the regular 28 days in the Zahn-Wellens test, without significant degradation and still 400 mg DOC/l, 200 mg DOC/l benzoate was added as a well degradable substrate; after 5 days, only 150 mg DOC/l was left, showing some cometabolic degradation *1

- biphosphonates form complexes with bivalent cations, in the relatively high concentrations in the algal test they deplete the medium as scavengers; hence, the effect on algae is not toxic in the strict sense *1

*1 referring to: lbandronate

SECTION 13: Disposal considerations

13.1. Waste treatment methods

Waste from residues - observe local/national regulations regarding waste disposal

SECTION 14: Transport information

Note - not classified as Dangerous Good according to the Dangerous Goods Regulations, proper shipping name non-regulated

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SECTION 15: Regulatory information

15.1. Safety, health and environmental regulations/legislation specific for the substance or mixture

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| TSCA Status | - FDA Exemption - not on inventory |
| Reporting Requirements | <ul style="list-style-type: none">- The United States Environmental Protection Agency (USEPA) has not established a Reportable Quantity (RQ) for releases of this material.- In New Jersey, report all releases, which are likely to endanger the public health, harm the environment or cause a complaint, to the NJDEPE Hotline and to local officials.- State and local regulations vary and may impose additional reporting requirements. |

SECTION 16: Other information

Full text of H-Statements referred to under section 3

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| H302 | Harmful if swallowed. |
| H314 | Causes severe skin burns and eye damage. |
| H373 | May cause damage to organs through prolonged or repeated exposure. |
| USH003 | May form combustible dust concentrations in the air |

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| Note | - Please note this Safety Data Sheet for the bulk product does not apply for the finished, packaged medicinal product intended for the final user. |
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| Edition documentation | - changes from previous version in sections 2, 15 |
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The information in this safety data sheet is based on current scientific knowledge. It should not be taken as expressing or implying any warranty concerning product characteristics.