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MATERIAL SAFETY DATA SHEET NOVARTIS PHARMACEUTICALS CORPORATION One Health Plaza East Hanover, NJ 07936

24-Hour Emergency Telephone Number: 1-862-778-7000 Customer Interaction Center (MSDS requests): 1-888-669-6682 For Technical Information: 1-862-778-3680 (9:00 AM – 5:00 PM E.S.T.)

SECTION 1. PRODUCT IDENTIFICATION

PRODUCT NAME: SYNONYMS: THERAPEUTIC CATEGORY: GENERIC NAME: CHEMICAL NAME:	Gleevec [™] 100mg STI571 Antitumor agent (tyrosine protein kinase inhibitor) Imatinib mesylate 4-[(4-Methyl-1-piperazinyl)methyl]-N-[4-methyl-3-[[4-(3-pyridinyl)-2-pyrimidinyl]amino]- phenyl]benzamide methanesulfonate
CHEMICAL FORMULA:	$C_{29}H_{31}N_7O \cdot CH_4SO_3$
MOLECULAR WEIGHT:	589.7

SECTION 2. COMPOSITION/INFORMATION ON INGREDIENTS

<u>COMPOSITION</u>	CAS#	CONCENTRATION (% by wt.)
Active Ingredients		
Gleevec Active Ingredient	220127-57-1	51.95
Inactive Ingredients		
Microcrystalline cellulose	9004-34-6	40.00
Crospovidone	9003-39-8	6.52
Colloidal silicon dioxide	7631-86-9	0.87
Magnesium stearate	557-04-0	0.65

SECTION 3. HAZARDS IDENTIFICATION

EMERGENCY OVERVIEW

FINISHED PHARMACEUTICAL PRODUCT REFER TO PHYSICIANS' DESK REFERENCE OR PACKAGE INSERT MAY CAUSE NAUSEA, VOMITING AND DIARRHEA

GleevecTM Capsules 100 mg Approval Date: 15 Jan 03

MAY CAUSE FLUID RETENTION MAY ADVERSELY AFFECT THE DEVELOPING FETUS

PRIMARY ROUTE(S) OF ENTRY: Oral

EFFECTS OF OVEREXPOSURE:	Finished pharmaceutical product. Potential for exposure is reduced in this form.
Skin:	No hazard is expected from normal clinical use.
Eye:	No hazard is expected from normal clinical use.
Inhalation:	No hazard is expected from normal clinical use.
Ingestion:	No hazard is expected from normal clinical use.
THERAPEUTIC SIDE EFFECTS:	Nausea, vomiting, diarrhea, fluid retention, muscle cramps, skin rash, headache, fatigue, arthralgia, and abdominal pain.
TARGET ORGAN EFFECTS:	Prolonged or repeated exposure may cause liver and kidney toxicity, and immunosuppresion.
REPRODUCTIVE HAZARDS:	FDA Pregnancy Category D (see section 11).
CARCINOGENICITY:	See section 11.
MUTAGENICITY:	Imatinib mesylate was clastogenic in one in vitro assay, and non-mutagenic in three assays (see Section 11).

MEDICAL CONDITIONS AGGRAVATED BY EXPOSURE: Pregnancy; known hypersensitivity to imatinib or any other components of the formulation; pre-existing liver impairment.

Skin Contact:	Wash contaminated area with soap and water.
Eye Contact:	Flush with running water for 15 minutes holding eyelids open.
Inhalation:	No specific treatment is necessary since this product is not likely to be
	hazardous by inhalation if capsule is left intact.
Ingestion:	Get medical attention immediately; induce vomiting if victim is conscious.

SECTION 5. FIRE FIGHTING MEASURES

Flash Point:Not applicableMethoFlammable Limits (% in air)	od Used: Not applicable
	plicable
Autoignition Temperature:	Not available
Extinguishing Media:	Use media suitable for fire in surrounding area.
Special Fire Fighting Procedures and Precautions	Evacuate area and fight fire from safe distance.
Fire and Explosion Hazards:	Not available
Fire-Fighting Equipment:	Wear full protective clothing and positive pressure self-
	contained breathing apparatus.
Hazardous Products of Combustion:	COx, NOx, SOx

SECTION 6. ACCIDENTAL RELEASE MEASURES

Steps to be taken if Material is Released or Spilled: Using appropriate protective equipment, sweep up and containerize spilled material. Avoid contamination of sewers and waterways.

SECTION 7. HANDLING AND STORAGE

Do not store above $86^{\circ}F(30^{\circ}C)$.
See container packaging.
None known.
None known.

SECTION 8. EXPOSURE CONTROLS/PERSONAL PROTECTION

Eye Protection:	Not required under normal conditions of therapeutic administration and use.
Skin Protection:	Not required under normal conditions of therapeutic administration and use. Protective gloves should be worn if contents of capsule are expelled.
Respiratory Protection:	Not required under normal conditions of therapeutic administration and use.
Ventilation Requirements:	Not required under normal conditions of therapeutic administration and use.
Additional Measures:	None

Exposure Limits (Definition of terms):

NPIEL: Novartis Pharma Internal Exposure Limit

Component

Imatinib mesylate

 $\frac{\text{Exposure Limit}}{\text{NPIEL} = 0.013 \text{ mg/m}^3}$

SECTION 9. PHYSICAL AND CHEMICAL PROPERTIES

Appearance: Color: Boiling Point: Melting/Freezing Pt.: pH: Specific Gravity: Soluble In: Capsule Orange to grayish orange opaque Not applicable Not available Not available Water

Odor Threshold: Odor Characteristics: Vapor Pressure (mm Hg): Vapor Density: % Volatile by Wt:

Not available Not available Not applicable Not applicable Not applicable

SECTION 10. STABILITY AND REACTIVITY

Yes Will not occur. Protect from temperatures exceeding 86°F (30°C). None known None known

SECTION 11. TOXICOLOGICAL INFORMATION

No toxicological data on finished product; data is for drug substance.

Eye Irritation:	No data available.
Skin Irritation/Sensitization:	Non irritating to the skin of rabbits; not sensitizing to the skin of guinea pigs.
Oral Toxicity:	MTD Oral (rat): > 600 mg/kg
Parenteral Toxicity:	LD ₅₀ Intravenous (rat): >100 mg/kg
Dermal Toxicity:	No data available.
Inhalation Toxicity:	Respiratory irritant (human)
Chronic/Carcinogenicity:	Carcinogenicity studies have not been performed with imatinib mesylate.
Mutagenicity:	<u>Positive in the following tests</u> : <i>in vitro</i> chromosome aberration test in ovarian cells of the Chinese hamster. <u>Negative in the following tests</u> : <i>in vitro</i> bacterial cell assay (Ames test), <i>in vitro</i> mammalian cell assay (mouse lymphoma) and an <i>in vivo</i> rat micronucleus assay.
Reproductive Effects:	Imatinib mesylate was teratogenic in rats when administered during organogenesis at doses \geq 100 mg/kg, approximately equal to the maximum clinical dose of 800 mg/day, based on body surface area. Teratogenic effects included exencephaly or encephalocele, absent/reduced frontal and absent parietal bones. Female rats administered this dose also experienced significant post-implantation loss in the form of early fetal resorption. At doses higher than 100 mg/kg, total fetal loss was noted in all animals. Women of childbearing potential should be advised to avoid becoming pregnant.
	In a study of fertility, in male rats dosed for 70 days prior to mating, testicular and epididymal weights and percent motile sperm were decreased at 60 mg/kg, approximately equal to the maximum clinical dose of 800 mg/day. When female rats were dosed 14 days prior to mating and through to gestational day 6, there was no effect on mating or on number of pregnant females.
	It is not known whether imatinib or its metabolites are excreted in human milk. However, in lactating female rats administered 100 mg/kg, a dose approximately equal to the maximum clinical dose of 800 mg/day based on body surface area, imatinib and/or its metabolites were extensively excreted in milk. Therefore, women should be advised against breastfeeding while taking Gleevec.

SECTION 12. ECOLOGICAL INFORMATION

No ecological data on finished product; data is for drug substance.

Bacteria toxicity (respiration inhibition): activated sludge (3h):

$$\begin{split} & EC_{10}: 65 mg/l \\ & EC_{50}: 232 mg/l \\ & EC_{80}: 605 mg/l \\ & \textbf{Fish toxicity: common carp (cyprinus carpio) (96h):} \\ & LC_{0}: 56 mg/l \\ & LC_{50}: 82 mg/l \\ & \textbf{Daphnia toxicity: } daphnia magna (water flea) (48h): \\ & EC_{50}: 80 mg/l \\ & NoEC: 32 mg/l \\ & \textbf{Algae toxicity: } Selenastrum capricornutum. Green algae. (72h): \\ & EbC_{50}: 2.5 mg/l \\ & EbC_{10}: 1.1 mg/l \\ & NOEC: 0.96 mg/l \\ & \textbf{Biological elimination: } 9 - 12\% (aerobic) (28d) \\ & Inhibitory effects can be excluded. \end{split}$$

Bioaccumulation in water organisms is not likely based on the n-octanol/water partition coefficient (log pOW < 3.0). Avoid release into the environment

SECTION 13. DISPOSAL CONSIDERATIONS

Waste Disposal Method:	All wastes must be disposed of in accordance with local, state and federal laws and regulations. (Contact local or state environmental agency for specific rules).
EPA Hazardous Waste Number:	None

SECTION 14. TRANSPORTATION INFORMATION

Ground Regulations:

Proper Shipping Description:	Drugs, N.O.I. NMFC Item 60000
DOT Proper Shipping Name:	Not Applicable
DOT Hazard Class:	Not Applicable
DOT Identification Number:	Not Applicable
Packing Group:	Not Applicable
Hazard Label:	Not Applicable
Package Weight Limits:	Not Applicable
Special Requirements:	Not Applicable
Exceptions:	Not Applicable
Non-Bulk Requirements:	Not Applicable
Bulk Requirements:	Not Applicable
Reportable Quantity (lbs.):	Not Applicable
Stowage:	Not Applicable

Other Requirements:

Not Applicable

Air Regulations:

Proper Shipping Description:	Drugs, N.O.I. NMFC Item 60000
IATA Proper Shipping Name:	Not Applicable
IATA Hazard Class:	Not Applicable
IATA Identification Number:	Not Applicable
Packing Group:	Not Applicable
Hazard Label:	Not Applicable
Special Requirements:	Not Applicable
Max. wgt/pkg - Passgr. Aircra	ft: Not Applicable
Max. wgt/pkg - Cargo Only Ai	r: Not Applicable

SECTION 15. REGULATORY INFORMATION

OSHA (Occupational Safety & Health Administration	a): This Material Safety Data Sheet contains the information required by the Federal OSHA Hazard Communication Standard (29 CFR 1910.1200).
OSHA PSM (Process Safety Management):	Not listed (29 CFR 1910.119, Appendix A)
NJ TCPA (Toxic Catastrophe Prevention Act):	This product contains NONE of the substances subject to the reporting requirements of Section N.J.A.C. 7:31 of this act.
TSCA (Toxic Substance Control Act):	Not applicable
CERCLA (Comprehensive Response Compensation &	& Liability Act): Not listed
SARA Title III (Superfund Amendments & Reauthori	ization Act):
Section 302 Extremely Hazardous Substances	: Not listed
Section 311/312 Hazard Categories:	None
Section 313 Reportable Ingredients:	Not listed
RCRA (Resource Conservation & Recovery Act):	Not listed
Other State Regulatory Information:	
New Jersey:	NJ RTK Threshold Planning Quantity = 10,000 lbs.
Other USA Regulations:	None
California Proposition 65:	The following statement is made in order to comply with the California Safe Drinking Water and Toxic Enforcement Act of 1986. This product does not contain any ingredient known to the State of California to cause cancer or reproductive toxicity.
Canada:	WHMIS Ingredient Disclosure List Not listed
EEC Classification (European Economic Community):	Warning Symbol: not available.

Risk Phrases: not available. **Safety Phrases:** not available.

SECTION 16. OTHER INFORMATION

Reason for Issue: New

Approved By:J. AffusoDate:15 Jan 03

To the best of our knowledge, the information contained herein is accurate. However, Novartis Pharmaceuticals Corporation does not assume any liability whatsoever for the accuracy or completeness of the information contained herein except for the product's administration/use as intended. Final determination of the suitability of any material is the sole responsibility of the user. All materials may present unknown hazards and should be used with caution. Although certain hazards are described herein, we cannot guarantee that these are the only hazards which exist.