KADCYLA(R) Lyophilized Powder Safety Data Sheet in Vials (100 mg) SECTION 1: Identification of the substance/mixture and of the company/undertaking 1.1. Product identifier Product name KADCYLA(R) Lyophilized Powder in Vials (100 mg) Product code SAP-10138138 Synonyms - T-DM1 with excipients lyophilized - KADCYLA Lyophilized Vials 1.2. Relevant identified uses of the substance or mixture and uses advised against Use - pharmaceutical active substance (antineoplastic) *1 1.3. Details of the supplier of the safety data sheet Company information Enquiries: Local representation: Genentech, Inc. 1 DNA Way South San Francisco USA-CA 94080 United States of America 001-(650) 225-1000 Phone info.sds@roche.com E-Mail US Chemtrec phone: (800)-424-9300 1.4. Emergency telephone number Emergency telephone number US Chemtrec phone: (800)-424-9300 *1 referring to: ado-trastuzumab emtansine **SECTION 2: Hazards identification Emergency Overview** Form sterile, lyophilized powder Color white or practically white

Classification of the substance or mixture / Label elements					
GHS Classification	H332 Har 3.1 Acute toxic H301 Tox 3.5 Germ cell r H340 May 3.7 Reproduct	 3.1 Acute toxicity (Category 4) H332 Harmful if inhaled. 3.1 Acute toxicity (Category 3) H301 Toxic if swallowed. 3.5 Germ cell mutagenicity (Category 1B) H340 May cause genetic defects. 3.7 Reproductive toxicity (Category 1B) H360FD May damage fertility. May damage the unborn 			
	Signalword: Dange	r			
	Label:				
	 P280 Wear protection. P301 + P310 IF S CENTER or doct P302 + P352 IF G P304 + P312 IF I doctor/physician P309 + P310 IF G 	cial instructions before use. ective gloves/ protective clothing / eye protection / SWALLOWED: Immediately call a POISON			
Other hazards					
Note	- no further information	ation available			
SECTION 3: Composition/information on ingredients					
Characterization	Ado-trastuzumab	emtansine with other inactive ingredients			
Ingredients	Concentration	GHS-Classification (pure ingredient)			
ado-trastuzumab emtansine 1018448-65-1	~ 24 %	 Combustible dust (No category), USH003 Acute toxicity (Category 2), H330 Acute toxicity (Category 2), H300 Skin corrosion/irritation (Category 2), H315 Germ cell mutagenicity (Category 1B), H340 Carcinogenicity (Category 2), H351 Reproductive toxicity (Category 1B), H360FD Specific target organ toxicity - Single exposure (Category 2), H371 			
For the full text of the H-phras	ses mentioned in this	Section, see Section 16.			

SECTION 4: First aid measures				
4.1. Description of first aid mea	sures			
Eye contact	 rinse immediately with tap water for 10 minutes - open eyelids forcibly 			
Skin contact	 remove immediately contaminated clothes, wash affected skin with water and soap - do not use any solvents 			
Inhalation	 remove the casualty to fresh air and keep him/her calm 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	e medical attention and special treatment needed			
Note to physician	- treat symptomatically			
SECTION 5: Firefighting m	neasures			
5.1. Extinguishing media				
Suitable extinguishing media	 adapt extinguishing media to surrounding fire conditions, water spray jet, dry powder, foam, carbon dioxide 			
Flash point (liquid)	not applicable			
5.2. Special hazards arising from the substance or mixture				
Specific hazards	- consider dust explosion hazard			
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, prote	ective equipment and emergency procedures			
Personal precautions	- prevent any exposure			
6.2. Environmental precautions				
Environmental protection	 if the substance reaches waters or the sewer system, inform the competent authority 			

6.3. Methods and material for containment and cleaning up						
Methods for cleaning up	 collect solids (avoid dust formation) and hand over to waste removal wash contaminated surfaces with sodium hydroxide solution, c(NaOH)=0.5 mol/l to 1 mol/l, and rinse with water 					
SECTION 7: Handling and	SECTION 7: Handling and storage					
7.1. Precautions for safe handli	ng					
Suitable materials	- glass					
7.2. Conditions for safe storage	, including any incompatibilities					
Storage conditions	 2 - 8 °C do not freeze protected from light 					
Validity	- 36 months, 2 to 8 °C, see expiry date on the label					
SECTION 8: Exposure con	trols/personal protection					
8.1. Control parameters						
Threshold value (Roche) air	- IOEL (Internal Occupational Exposure Limit): 0.0003 mg/m ³ *1					
8.2. Exposure controls						
General protective and hygiene measures	- instruction of employees mandatory					
Respiratory protection	 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 is recommended for dusty operations. respiratory protection not necessary during normal operations 					
Hand protection	- protective gloves (eg made of neoprene, nitrile or butyl rubber)					
Eye protection	- safety glasses					
*1 referring to:	ado-trastuzumab emtansine					
SECTION 9: Physical and chemical properties						
9.1. Information on basic physic						
Color	white or practically white					

KADCYLA(R) Lyophilized Powder in Vials (100 mg)

Form sterile, tyophilized powder Solubility solubie, water 9.2. Other information no information available SECTION 10: Stability and reactivity Second available 10.1. Reactivity no information available 10.2. Chemical stability - no information available 10.3. Possibility of hazardous reactions - Note - no information available 10.4. Conditions to avoid - Conditions to avoid - D.5. Incompatible materials - Note - Note - 0.6. Hazardous decomposition products Note - 10.6. Hazardous decomposition products Note - 2 referring to: Herceptin Vials (2% Trastuzumab solution with excipients) SECTION 11: Toxicological effects Acute toxicity - HNSTD 20 mg/kg (i.v., rai) - HNSTD 0 <t< th=""><th></th><th></th></t<>						
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- HNSTD 20 mg/kg (i.v., rat) *1 - LD ₅₀ 0 to 5 mg/kg (oral, mouse)	11.1. Information on toxicological effects					
	Acute toxicity	- HNSTD 20 mg/kg (i.v., rat) *1 - LD ₅₀ 0 to 5 mg/kg (oral, mouse)				

	- LC ₀ 0.5 μg/l (inhal., rat, 4 h) (OECD No. 403) LC ₁₀₀ 11.5 μg/l (inhal., rat, 4 h) (OECD No. 403) *3
Subacute toxicity	- HNSTD 10 mg/kg/3w(i.v., cynomolgus monkey, 9 weeks) *1
Local effects	- skin, eyes, mucous membranes: corrosive *4
Sensitization	anaphylactic reactions may occur following the intravenous application of proteins; rare cases of hypersensitivity have been described *5
Mutagenicity	 negative (Ames test) *4 OECD No. 474 (Micronucleus Test); positive: evidence of aneugenicity and/or clastogenicity *4
Carcinogenicity	- no information available
Reproductive toxicity	- no information available
STOT-single exposure	- no information available
STOT-repeated exposure	- no information available
Aspiration hazard	- no information available
Note	 HNSTD = Highest Non-Severely Toxic Dose
Potential Health Effects	 Exposure: Inhalation, Ingestion, Skin contact, Eye contact Target Organs: liver, Cardiovascular system, gastrointestinal system, Hematopoietic/blood system, Immune System, respiratory system
	 Acute Effects: May cause allergic reactions., Harmful if swallowed., May cause headache., May cause musculoskeletal effects., May cause general body weakness, fatigue and nausea.
	- Chronic Effects: May cause hepatic (liver) system effects., Signs and symptoms may include elevation of liver enzyme levels and jaundice (yellowing of the skin and eyes)., May cause cardiovascular effects., Signs and symptoms may include increase or decrease in blood pressure, irregular heartbeat, chest pains and cardiac arrest., May cause blood system changes., May cause respiratory effects., Signs and symptoms may include difficulty in breathing, coughing, wheezing, irritation (inflammation) and respiratory arrest.
	- Carcinogenicity: not listed by NTP, IARC or OSHA
Additional Health Information	 Conditions Aggravated: Hypersensitivity to this material and other materials in its chemical class.
 *1 referring to: *3 referring to: *4 referring to: *5 referring to: 	ado-trastuzumab emtansine Ansamitocin P3 DM1 Trastuzumab

SECTION 12: Ecologic	al information	
12.1. Toxicity		
Ecotoxicity	 barely toxic for algae, growth inhibition possibly due to turbidity caused by test substance (Desmodesmus (=Scenedesmus) subspicatus) ErC₅₀ (72 h) > 100 mg/l (nominal concentration) EyC₅₀ (72 h) ~ 100 mg/l (nominal concentration) (OECD No. 201) barely toxic for planktonic crustaceans (Daphnia magna) EC₅₀ (48 h) > 100 mg/l (nominal concentration) NOEC (48 h) 100 mg/l (nominal concentration) (OECD No. 202) barely toxic for fish (guppy) LC₅₀ (96 h) > 100 mg/l (nominal concentration) NOEC (96 h) < 100 mg/l (nominal concentration) (OECD No. 203, semistatic) barely inhibitory on aerobic bacterial respiration (activated sludge concentration (14 d) 49.5 mg/l (nominal concentration) (Manometric Respirometry Test, OECD No. 301 F) 	*1 *1 e) *1
12.2. Persistence and degr	adability	
Ready biodegradability	 readily biodegradable 84 %, 28 d (Manometric Respirometry Test, OECD No. 301 F) 	*1
12.3. Bioaccumulative pote	ential	
Note	- no information available	
12.4. Mobility in soil		
Note	- no information available	
12.5. Results of PBT and v	PvB assessment	
Note	- no information available	
12.6. Other adverse effects		
Note	- no information available	
*1 referring to:	ado-trastuzumab emtansine	

SECTION 13: Disposal considerations

13.1. Waste treatment methods

Waste from residues

observe local/national regulations regarding waste disposalincinerate in qualified installation with flue gas scrubbing

SECTION 14: Transport information

ΙΑΤΑ	Class	UN/ID	PG		PI	Label	Mark	
	6.1	3249	III		670/677	6.1		
IMDG	Class	UN	PG	EmS	PI	Label	Mark	
	6.1	3249	111	F-A S-A	P002/ -	6.1		
RID/ADR	Class	UN	PG	Haz.no	PI	Label	Mark	Classif. code
	6.1	3249	ш	60	P002/ -	6.1		T2
DOT	Class	UN/ID	PG	PI	RQ	Label	Haz.no	
	6.1	3249	111			6.1		
Proper shipping name MEDICINE, SOLID, TOXIC, N.O.S.								
SECTIO	SECTION 15: Regulatory information							

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

TSCA Status

- FDA Exemption - not on inventory

Reporting Requirements

- 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 (1-609-292-5560) and to local officials.
 - State and local regulations vary and may impose additional reporting requirements.

SECTION 16: Other information				
Full text of H-Statements r	H300 H315 H330 H340 H351 H360FD H371	under section 3 Fatal if swallowed. Causes skin irritation. Fatal if inhaled. May cause genetic defects. Suspected of causing cancer. May damage fertility. May damage the unborn child. May cause damage to organs. May form combustible dust concentrations in the air		
Note		This product may be shipped using De Minimis Quantity Exceptions, if the requirements of US 49 CFR §173.4b and ICAO 5.6/IATA 2.6.10 are met. Please note this Safety Data Sheet for the bulk product does not apply for the finished, packaged medicinal product intended for the final user.		
Edition documentation	-	changes from previous version in sections 2, 3, 11, 16		

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.