

iLite[®] Assay Ready Cells containing cryoprotective medium from Gibco

BM4025, BM4052, BM4075, BM5001, BM5010, BM5011, BM5013, BM5014, BM5015, BM5016, BM5017, BM5018, BM5035, BM5036, BM6069

SECTION 1: IDENTIFICATION OF THE SUBSTANCE/MIXTURE AND OF THE COMPANY/ UNDERTAKING

UNDERTAKING						
1.1 Product identifier						
Product name:	iLite® TLR4 Assay Ready Cells iLite® TLR9 Assay Ready Cells iLite® RANKL Assay Ready Cells iLite® C5a Assay Ready Cells iLite® ADCC Effector (V) Assay Ready Cells iLite® CD20 (+) Target Assay Ready Cells iLite® CD20 (-) Target Assay Ready Cells iLite® HER2 (+) Target Assay Ready Cells iLite® HER2 (-) Target Assay Ready Cells iLite® mTNF-alpha (+) Target Assay Ready Cells iLite® mTNF-alpha (-) Target Assay Ready Cells iLite® EGFR (+) Target Assay Ready Cells iLite® EGFR (-) Target Assay Ready Cells iLite® mVEGF (-) Target Assay Ready Cells iLite® mVEGF (-) Target Assay Ready Cells iLite® mVEGF (-) Target Assay Ready Cells					
Product description	iLite Assay Ready Cells containing cryoprotective medium from Gibco (cat no 12648-010)					
Product code	BM4025, BM4052, BM4075, BM5001, BM5010, BM5011, BM5013, BM5014, BM5015, BM5016, BM5017, BM5018, BM5035, BM5036, BM6069					
1.2 Relevant identified uses of the substance or mixture and uses advised against Use of the product Laboratory chemicals. For research use only.						
1.3 Details of the suppl	ier of the safety data sheet					
Company	Svar Life Science AB					
Address	Lundvägen 151					
Zip code/Place	SE-212 24 Malmö, Sweden					
Telephone	+46 40 53 76 00					
Website	www.svarlifescience.com					
E-mail	info@svarlifescience.com					
1.4 Emergency telepho	ne number					
Emergency telephone	(Sweden) Acute: 112 – Ask for "Giftinformation". If less acute call: +46 010 4566700.					
number	Other countries: Please contact local emergency telephone number.					
SECTION 2: HAZARDS IDENTIFICATION						
2.1 Classification of the	e substance or mixture					
Classification according to the Regulation (EC) No. 1272/2008 (CLP):						
The mixture is not to be class						

and classified as a Class 1 Genetically Modified Microorganism.

The mixture is covered by Directive 2000/54/EC of 18 September 2000 on the protection of workers from risks related to exposure to biological agents at work.

2.2 Label elements

None

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2.3 Other hazards

not result in classification

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Other hazards which do Contain Fetal Bovine serum, which is derived from cattle with US origin. The Certificate of Analysis for FBS show that the substance has been analyzed for Blue tongue Virus. Bovine Adenovirus, Bovine Parvovirus, Rabies Virus, Reovirus, BRSV Fluorescent Antibody, BVDV Fluorescent Antibody, Cytopathogenic agents and Hemadsorbing agents with a negative result.

The products are considered to be biological agents in group 1 (ie. a biological agent that is unlikely to cause human infection). As a precaution, it is recommended that the work is carried out under measures similar to Group 2 in Council Directive 2000/54/EC.

PBT/ vPvB: No

Substance meets the criteria for PBT/ vPvB under Regulation EC No. 1907/2006, appendix XIII

Endocrine disrupting properties

The substances are not identified as having endocrine disrupting properties in accordance with the criteria set out in Regulation 2017/2100 or Regulation 2018/605.

SECTION 3: COMPOSITION/INFORMATION ON INGREDIENTS

3.2 Mixtures

Assay	Ready Cells suspended in	cryoprotective r	nedium from	Gipco containing the follo	wing sub	ostances:
No	Product/ingredient name	EC-number	CAS- number	REACH registration number	Conc. (%w/w)	Classification Regulation (EC) No. 1272/2008 [CLP]
	Fetal Bovine Serum (Heat inactivated FBS)				10	None
	Dimethyl Sulfoxid (DMSO)	200-664-3	67-68-5		10	None

SECTION 4: FIRST-AID MEASURES

4.1 Description of first aid measures

On suspicion of possible infection from biological agents – seek medical advice! Inhalation: Move to fresh air. Keep at rest and under surveillance. If needed: seek medical advice. Remove contaminated clothing at once. Flush skin and wash thoroughly with soap and Skin contact: water. Eye contact: Keep eyelids well apart. Rinse with water for a couple of minutes, remember to remove contact lenses if any. If irritation persists: Seek medical advice. Rinse mouth and drink plenty of water. If needed or if larger amounts has been Ingestion swallowed: Seek medical advice.

4.2 Most important symptoms and effects, both acute and delayed

Skin contact:	May cause irritation of skin.
Eye contact:	May cause irritation of eyes.
Inhalation	Prolonged or frequent exposure to vapours of volatile organic compounds may result in
	damage on liver, kidneys, blood or central nervous system (including brain damage).

4.3 Indication of any immediate medical attention and special treatment needed

Show this safety data sheet to a physician or emergency ward

SECTION 5: FIREFIGHTING MEASURES

5.1 Extinguishing media

Use water spray, carbon dioxide, dry chemical or foam. Suitable extinguishing media Unsuitable Waterjet extinguishing media

5.2 Special hazards arising from the substance or mixture

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Hazards from the substance or mixture Hazardous thermal decomposition products	None Decomposition products may include the following materials: oxides of carbon and sulphur.
5.3 Advice for firefighters Special protective actions for fire-fighters	Promptly isolate the scene by removing all persons from the vicinity of the incident if there is a fire. No action shall be taken involving any personal risk or without suitable training.
Special protective equipment for fire- fighters	Fire-fighters should wear appropriate protective equipment and self-contained breathing apparatus (SCBA) with a full face-piece operated in positive pressure mode. Clothing for fire-fighters (including helmets, protective boots and gloves) conforming to European standard EN 469 will provide a basic level of protection for chemical incidents.
Further information	Not applicable

SECTION 6: ACCIDENTAL RELEASE MEASURES

6.1 Personal precautions, protective equipment and emergency procedures

For non-emergency personnel	Use personal protective equipment – see section 8. No action shall be taken involving any personal risk or without suitable training. Evacuate surrounding areas. Keep unnecessary and unprotected personnel from entering. Do not touch or walk through
For emergency responders	spilt material. The employees or the company's occupational health and safety organization must be informed immediately of any accident or incident that may have resulted in the release of biological agents, which may cause disease in humans. If specialized clothing is required to deal with the spillage, take note of any information in Section 8 on suitable and unsuitable materials. See also Section 8 for additional information on hygiene measures.

6.2 Environmental precautions

Do not empty into drains - see section 12. Inform appropriate authorities in accordance with local regulations.

6.3 Methods and material for containment and cleaning up

er
nt

6.4 Reference to other sections

See Section 8 for information on appropriate personal protective equipment. See Section 13 for additional waste treatment information.

SECTION 7: HANDLING AND STORAGE

7.1 Precautions for sat	fe handling
Protective measures	Before use, a workplace assessment of the safety and health conditions at the working place must be carried out according to the general specifications mentioned in the EU Directive 2000/54/EC on biological agents and work. Work must be planned, organized and carried out so that the influence of biological agents is avoided as much as possible. Use laboratory facilities, which generally qualify for handling of biological agents. No tool or used material should after end use be placed on tables or similar but collected immediately in special sealed containers. Recycling of tools should only take place after proper disinfecting and purification. If appliances are contaminated, washing must be made with appropriate disinfectant before further use.
Advice on general occupational hygiene	Eating, drinking and smoking should be prohibited in areas where this material is handled. Avoid contact with skin, eyes and clothing. Always wash hands with soap and
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water after completing work, and when leaving the laboratory (e.g. before going to the toilet and at the end of the workday). Do not pipette by mouth pipetting. See also Section 8 for additional information on hygiene measures.

7.2 Conditions for safe storage, including any incompatibilities

Storage:

Upon receipt confirm that adequate dry-ice is present and the cells are frozen. Immediately transfer to -80 °C (do not store at any other temperature). Cells should be used within 30 min of thawing and should be diluted immediately after thawing. Not applicable

7.3 Specific end use(s)

Further information:

Laboratory chemicals for research use only.

SECTION 8: EXPOSURE CONTROLS/PERSONAL PROTECTION

	I parameters	limito F	MEO					
European Ur	nal exposure	innits - L	JMISO					
UK: None								
Sweden:	<u>NGV</u> 50 ppm = 150	a/am ³	KGV 150 ppm = {	$EQQ = \alpha/2m^3$	<u>Comments</u> H.V			
	H: Skin perme			e short-term exp				
Denmark: 50 Finland: 50 p Austria: 50 p	AK: 50 ppm = 16) ppm = 160 mg/ ppm (8h) pm = 160 mg/m ² 50 ppm = 160 m	m ³						
			1					
Recomment monitoring	nded g procedure	Not rele	evant					
-								
Derived eff		T	F	Malasa	Demoletien			
Product/in name	grealent	Туре	Exposure	Value	Population	Effects		
Predicted e	ffeet	Notovo	ilabla					
concentrat		Not available						
PNEC Sum	mary	Not available – No CSR						
8.2 Exposure controls								
	e engineering	Sufficient ventilation.						
Hygiene m	easures	Wash hands thoroughly after handling chemical products, before eating, drinking, smoking and using the lavatory and at the end of the working period. Appropriate techniques should be used to remove potentially contaminated clothing. Wash contaminated clothing before reusing. Ensure that eyewash stations and safety showers are close to the workstation location.						
Respirator	y protection	Not rele	vant during norm	nal condition.				
Eye/face pr	rotection	Use safe	ety glasses (acc	ording to EN166	6) when there is risk of	splashes.		
Hand prote	ection	Wear protective gloves (according to EN374) of butyl rubber or nitrile rubber.				⁻ nitrile rubber.		
Body prote	ection	Wear suitable protective clothing.						
Environme controls	ntal exposure	Not app	licable					

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SECTION 9: PHYSICAL AND CHEMICAL PROPERTIES

9.1 Information on basic physical and chemical properties				
Physical state:	Liquid			
Colour:	n.d			
Odour:	n.d			
Melting point/freezing point (°C):	n.d			
Boiling point or initial boiling point and boiling range (°C):	n.d			
Flammability (solid, gas):	n.a			
Lower and upper explosion limit (vol-%):	n.d			
Flash point (°C):	n.d			
Auto-ignition temperature (°C):	n.d			
Decomposition temperature (°C):	n.d			
pH:	n.d			
Kinematic viscosity:	n.d			
Solubility:	n.d			
Partition coefficient n-octanol/water (log value):	n.d			
Vapour pressure:	n.d			
Density and/or relative density:	n.d			
Relative vapour density:	n.d			
Particle characteristics:	n.a			

n.d = not determined n.a = not applicable

9.2 Other information

Not applicable

SECTION 10: STABILITY AND REACTIVITY				
10.1 Reactivity	No available	information		
10.2 Chemical stability	Stabile at rec	commended storage conditions – see section 7.		
10.3 Possibility of hazardous reactions No available information.				
10.4 Condition to avoid	No available	information.		
10.5 Incompatible materials	No available	information.		
10.6 Hazardous decomposition	n products	When heated to high temperatures (decomposition) toxic fumes are emitted: Oxides of carbon and sulphur.		



SECTION 11: TOXICOLOGICAL INFORMATION

In addition to the hazardous properties mentioned below, the risk of infection from the biological agents present in the product must also be taken into account.

11.1 Information on toxicological effects

Hazard class	Data (DMSO)	Test	Reference
Acute toxicity:			
Inhalation	LD ₅₀ (rat) > 2 mg/l/4h	No information	IUCLID
Dermal	LD _{Lo} (rat) > 40000 mg/kg	No information	IUCLID
Oral	LD ₅₀ (rat) = 14500 mg/kg	No information	IUCLID
Corrosion/irritation	Mild eye and skin irritation, rabbit	OECD 404, EU	ECHA
		Method B.5	
Sensitization	No skin sensitization, guinea pig	Buehler	IUCLID
CMR	No mutagenicity, carcinogenicity, genotoxicity	Several	Merck/IUCLID

Acute toxicity

Assessment for other reagents than DMSO: No data available.

Irritation/Corrosion

Assessment for other reagents than DMSO: No data available.

Sensitization by inhalation/skin contact

Assessment for other reagents than DMSO: No data available.

Germ cell mutagenicity

Assessment for other reagents than DMSO: The chemical structure of the different reagents doesn't indicate any mutagenic effects.

Carcinogenicity

Assessment for other reagents than DMSO: The chemical structure of the different reagents doesn't indicate any carcinogenic effects.

Reproduction toxicity

Assessment for other reagents than DMSO: The chemical structure of the different reagents doesn't indicate any reproduction toxic effects.

Developmental toxicity

Assessment for other reagents than DMSO: The chemical structure of the different reagents doesn't indicate any teratogenic effects.

Specific target organ toxicity (single exposure)

STOT assessment single dose toxicity: No data available.

Repeated dose toxicity and specific organ toxicity (repeated exposure)

Prolonged or frequent exposure to vapours of volatile organic compounds may result in damage on liver, kidneys, blood or central nervous system (including brain damage).

11.2. Information on other hazards:

None known.



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SECTION 12: ECOLOGICAL INFORMATION

12.1 Toxicity

12.1.1 Acute toxicity in the aquatic environment for DMSO

Test	
Fish LC ₅₀	
Daphnia EC50	
Algae EC ₅₀	

Value/unit (mg/l) 32000 7000 12350-25500

Test method Static (FW) No info. (FW) No info. (SW)

 Not
 Exp. time (h)

 96
 96

 V)
 24

 N)
 96

Species Oncorhynchus mykiss Daphnia sp. Skeletonema costatum

12.1.2 Acute toxicity in the aquatic environment other reagents than DMSO No data available.

12.1.3 Ecotoxicity No data available.

12.2 Persistence and degradability

Conclusion/Summary DMSO is not readily degradable (3.1% after 14 days in OECD 301C test).

12.3 Bioaccumulative potential

Conclusion/Summary DMSO: Log K_{ow} -1,35 – No significant bioaccumulation.

12.4 Mobility in soil Soil/water partition

coefficient (KOC)

DMSO: K_{oc} (calculated) < 10 – Very high mobility expected in soil environments.

No available data

12.5 Results of PBT and vPvB assessment

PBTThe substance is not considered PBT according to criteria in Annex XIIII.vPvBThe substance is not considered vPvB according to criteria in Annex XIIII.

12.6. Endocrine disrupting properties

None known.

Mobility

12.7 Other adverse effects

None known.

SECTION 13: DISPOSAL CONSIDERATIONS

13.1 Waste treatment methods

Method of disposal	Biological agents are considered hazardous waste. Disposal should be according to local, state or national legislation.
	Note! Waste containers containing biological material must be labeled with: 🐼 (black
	symbol on yellow background). The generation of waste should be avoided or minimized wherever possible. This material and its container must be disposed of in a safe way by incineration.
Hazardous waste	Within the present knowledge of the supplier, this product is regarded as hazardous waste, as defined by EU Directive 2008/98/EC.

European Waste Catalogue (EWC)

EWC Waste Code	Type of waste
18 01 03	Wastes whose collection and disposal is subject to special requirements in order to prevent infection
15 02 02	Absorbent material containing residues of or contaminated by dangerous substances

Packaging

Method of disposal	Incineration.
Special precautions	None.

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SECTION 14: TRANSPORT INFORMATION

Product is not classified as dangerous goods.

	ADR/RID	ADN/ADNR	IMDG	IATA
14.1 UN number or ID number				
14.2 UN proper shipping name				
14.3 Transport hazard class(es)				
14.4 Packing Group				
14.5 Environmental hazards				
14.6 Special precautions for user	No	No	No	No
14.7 Maritime transport in bulk	Not applicable	Not applicable	Not applicable	Not applicable
according to IMO instruments				
Additional information	Waste containing used biological agents <u>may</u> be considered as dangerous goods; UN 3291, CLINICAL WASTE, UNSPECIFIED; N.O.S., or (BIO) MEDICAL WASTE N.O.S. or REGULATED MEDICAL WASTE, N.O.S. Class 6.2 Packing Group II			

SECTION 15: REGULATORY INFORMATION

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

Must not be used by persons under 18 years of age (Directive 94/33/EC).

The employer shall assess the working conditions and, if there is any risk to the safety or health and any effects on the pregnancy or breastfeeding of workers, take the necessary measures to adjust the working conditions (Directive 92/85/EEC)

The mixture is covered by:

Directive 2009/41/EC on the contained use of genetically modified micro-organisms

Directive 2000/54/EC - biological agents at work

EU Regulation (EC) No. 1272/2008 (CLP): Not classified.

EU Regulation (EC) No. 1907/2006 (REACH)

Annex XIV – List of substances subject to authorization

Substances of very high concern

None of the components are listed.

Annex XVII – Restrictions on the manufacture, placing on the market and use of certain dangerous substances, mixtures and articles Not applicable

Not applicable

15.2 Chemical Safety Assessment No CSR.

Other information

Tariff Code harmonized	Not applicable
system	
The EU Seveso Directive	Not applicable

International regulations

Chemical Weapons Convention List	Chemical Weapons Convention List	Chemical Weapons Convention List
Schedule I Chemicals	Schedule II Chemicals	Schedule III Chemicals
Not regulated	Not regulated	Not regulated

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SECTION 16: OTHER INFORMATION

Conforms to Regulation (EC) No. 1907/2006 (REACH), Annex II.

LIST OF HAZARD STATEMENTS MENTIONED UNDER SECTION 3: None

Abbreviations:

BRSV = Bovine Respiratory Syncytial Virus BVDV = Bovine Viral Diarrhea Virus CMR = Carcinogenicy, Mutagenicity, and Reproduction toxicity CSR = Chemical Safety Report DNEL = Derived No-Effect Level EC50 - Half maximal effective concentration FW = Fresh Water (Färskvatten) KGV = Korttidsvärde (Swedish for short term exposure limit) LC50 = Lethal Concentration 50 % LD50 = Lethal Dose 50 % MAK = Maximale Arbeitsplatzkonzentrationen (German for maximum workplace concentration) NGV = Nivågränsvärde (Swedish for exposure limit) PBT = Persistent, Bioaccumulative, Toxic PNEC = Predicted No-Effect Concentration vPvB = very Persistent, very Bioaccumulative

Literature:

Merck (Safety Data Sheet) IUCLID = International Uniform ChemicaL Information Database ECHA = European Chemicals Agency

Other information

No special training is required. However, the user should be well instructed in the execution of his/her task, be familiar with this Safety Data Sheet and have normal training in the use of personal protective equipment.

Revisions

Version	Valid from (date)	Changes
1.0	14-Mar-2019	New document
2.0	15-Oct-2019	Addition of BM5017 and BM5018
3.0	18-Mar-2020	Addition of BM4052
4.0	14-May-2020	Addition of BM4075
5.0	24-Sep-2020	Addition of BM4073
6.0	21-Jun-2021	Addition of BM4025. Template updated in compliance with
		Regulation 2020/878
7.0	08-Oct-2021	Addition of BM6069. Removal of BM4073



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