

**THE UNIVERSITY OF WESTERN ONTARIO
BIOLOGICAL AGENTS REGISTRY FORM**
Approved Biohazards Subcommittee: October 14, 2010
Biosafety Website: www.uwo.ca/humanresources/biosafety/

This form must be completed by each Principal Investigator holding a grant administered by the University of Western Ontario (UWO) or in charge of a laboratory/facility where the use of Level 1, 2 or 3 biological agents is described in the laboratory or animal work proposed. The form must also be completed if any work is proposed involving animals carrying zoonotic agents infectious to humans or involving plants, fungi, or insects that require Public Health Agency of Canada (PHAC) or Canadian Food Inspection Agency (CFIA) permits.

This form must be updated at least every 3 years or when there are changes to the biological agents being used.

Containment Levels will be established in accordance with Laboratory Biosafety Guidelines, 3rd edition, 2004, Public Health Agency of Canada (PHAC) or Containment Standards for Veterinary Facilities, 1st edition 1996, Canadian Food Inspection Agency (CFIA).

Completed forms are to be returned to Occupational Health and Safety, (OHS), (Support Services Building, Room 4190) for distribution to the Biohazards Subcommittee. For questions regarding this form, please contact the Biosafety Officer at extension 81135 or biosafety@uwo.ca. If there are changes to the information on this form (excluding grant title and funding agencies), contact Occupational Health and Safety for a modification form. See website: www.uwo.ca/humanresources/biosafety/

PRINCIPAL INVESTIGATOR	<u>James Lewis, Ruud Veldhuizen, and Cory Yamashita</u>
DEPARTMENT	<u>Medicine</u>
ADDRESS	<u>SJHC, Grosvener St.</u>
PHONE NUMBER	<u>646-6100, Ext 66288</u>
EMERGENCY PHONE NUMBER(S)	<u></u>
EMAIL	<u>jflewis@uwo.ca, rveldhui@uwo.ca, cyamash@uwo.ca</u>

Location of experimental work to be carried out: Building(s) LHRI_ Room(s)_F4-117 for LPS, F4-124 for cell culture

*For work being performed at Institutions affiliated with the University of Western Ontario, the Safety Officer for the Institution where experiments will take place must sign the form prior to its being sent to the University of Western Ontario Biosafety Officer (See Section 15.0, Approvals).

FUNDING AGENCY/AGENCIES: CIHR
GRANT TITLE(S): The Role of Pulmonary Surfactant in Acute Lung Injury and Progression to Multi-Organ Failure

List all personnel working under Principal Investigators supervision in this location:

<u>Name</u>	<u>UWO E-mail Address</u>	<u>Date of Biosafety Training</u>
<u>Lynda McCaig</u>	<u>lmccaig@uwo.ca</u>	<u>2004</u>
<u>Lijuan Yao</u>	<u>jyao@uwo.ca</u>	<u>2004</u>
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Please explain the biological agents and/or biohazardous substances used and how they will be stored, used and disposed of. Projects without this description will not be reviewed.

LPS (lipopolysaccharide):

LPS powder is stored at 4 C fridge, after dissolve in saline, make aliquots and stored in -20C

We handle LPS powder in the fume hood with gloves, mask, lab coat and safety glasses on. When we use this saline dissolved LPS in animals we also wear gloves, lab coat and safety glasses.

The animals that have been used with LPS are disposed of in a biohazard waste box for incineration within the ACF.

Cell line:

MLE-12 (Mouse lung epithelial cells)

Cells are aliquoted and stored in the liquid nitrogen tank for storage

We perform all cell experiment in the cell culture room within F4-124 (Dr. David Hill's laboratory)

All wastes are collected in a biohazard box for incineration

Please include a one page research summary or teaching protocol.

The overall objective of research in our lab is to understand the mechanisms leading to Acute Lung Injury and to develop logical treatment strategy to improve the outcome of patients afflicted with this syndrome.

ALI is defined by the inability of the lung facilitating the diffusion of oxygen from the atmosphere into the blood. This type of lung dysfunction can occur in people of all ages due to one or more insult to the lung. Some of the more common initiating event leading to ALI are systemic infection, pneumonia, aspiration of gastric content and trauma. The reported incidence of ALI ranges from 15 to 79 cases per 100,000 person-years depending on the specific population studied, and has a mortality of approximately 40%. In addition, patient surviving ALI, have often required a prolonged stay in the Intensive Care Unit. As such, ALI represents a significant burden on our healthcare system.

From a research perspective, one of the important aspects of this disease that has limited our ability to develop appropriate therapies is the complexity of the disease. The multiple events involved in the development of the disease have been shown to contribute to a very complex, multi-factorial process which may vary among individual patients. To address this issue our lab has focused on specific common pathological processes such as the contribution of mechanical ventilation and the role of pulmonary surfactant.

Experimentally, our lab tries to use several complementary approaches to study the ALI from a physiological to a molecular level. At the physiological level we utilize animal models of ALI that reflect the human disease. Specifically we study the physiological effects of a small amount of acid instilled into the lung of mice and rats as models of gastric acid aspiration that occurs in humans. We also study the effect of mechanical ventilation, the main supportive therapy for ALI, on the lungs of mice and rats. Material from the animal experiments is used for advanced biophysical analysis such as atomic force microscopy and captive bubble surfactometry, as well as inflammatory measurements such as the determination of cytokines, adhesion molecules and NFkB activation. To

study more specific cellular aspects of lung injury we utilize various cell lines (MLE12 and MLEC) to determine the effects of inflammation and cell stretch on cellular responses.

Lipopolysaccharide(LPS) is used in the experiments as a lung insult to make lung injury at a very low dose (1mg/kg). LPS is dissolved in saline and instilled into animal lungs. At the end of the experiment, animal bodies are sent to a biohazard waste box for incineration.

See E-mail

Question 1.1 → yes

1.0 Microorganisms

1.1 Does your work involve the use of biological agents? YES NO
 (non-pathogenic and pathogenic biological agents including but not limited to bacteria and other microorganisms, viruses, prions, parasites or pathogens of plant or animal origin)? **If no, please proceed to Section 2.0**

Do you use microorganisms that require a permit from the CFIA? YES NO
 If YES, please give the name of the species. _____

What is the origin of the microorganism(s)? _____
 Please describe the risk (if any) of escape and how this will be mitigated:

Please attach the CFIA permit.
 Please describe any CFIA permit conditions:

1.2 Please complete the table below:

Name of Biological Agent(s)* (Be specific)	Is it known to be a human pathogen? YES/NO	Is it known to be an animal pathogen? YES/NO	Is it known to be a zoonotic agent? YES/NO	Maximum quantity to be cultured at one time? (in Litres)	Source/ Supplier	PHAC or CFIA Containment Level
	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No			<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No			<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No			<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No	<input type="radio"/> Yes <input type="radio"/> No			<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3

*Please attach a Material Safety Data Sheet or equivalent from the supplier.

2.0 Cell Culture

2.1 Does your work involve the use of cell cultures?
If no, please proceed to Section 3.0

YES

NO

2.2 Please indicate the type of primary cells (i.e. derived from fresh tissue) that will be grown in culture:

Cell Type	Is this cell type used in your work?	Source of Primary Cell Culture Tissue	AUS Protocol Number
Human	<input type="radio"/> Yes <input type="radio"/> No		Not applicable
Rodent	<input type="radio"/> Yes <input type="radio"/> No		
Non-human primate	<input type="radio"/> Yes <input type="radio"/> No		
Other (specify)	<input type="radio"/> Yes <input type="radio"/> No		

2.3 Please indicate the type of established cells that will be grown in culture in:

Cell Type	Is this cell type used in your work?	Specific cell line(s)*	Containment Level of each cell line	Supplier / Source of cell line(s)
Human	<input type="radio"/> Yes <input type="radio"/> No			
Rodent	<input checked="" type="radio"/> Yes <input type="radio"/> No	Mouse MLE-12	2	ATCC
Non-human primate	<input type="radio"/> Yes <input type="radio"/> No			
Other (specify)	<input type="radio"/> Yes <input type="radio"/> No			

*Please attach a Material Safety Data Sheet or equivalent from the supplier. (For more information, see www.atcc.org)

2.4 For above named cell type(s) indicate PHAC or CFIA containment level required 1 2 2+ 3

3.0 Use of Human Source Materials

3.1 Does your work involve the use of human source materials? YES NO

If no, please proceed to Section 4.0

3.2 Indicate in the table below the Human Source Material to be used.

Human Source Material	Source/Supplier /Company Name	Is Human Source Material Infected With An Infectious Agent? YES/UNKNOWN	Name of Infectious Agent (If applicable)	PHAC or CFIA Containment Level (Select one)
Human Blood (whole) or other Body Fluid		<input type="radio"/> Yes <input type="radio"/> Unknown		<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
Human Blood (fraction) or other Body Fluid	Lung lavage from Proteinosis Patients	<input type="radio"/> Yes <input checked="" type="radio"/> Unknown		1 <input checked="" type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
Human Organs or Tissues (unpreserved)		<input type="radio"/> Yes <input type="radio"/> Unknown		<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
Human Organs or Tissues (preserved)		Not Applicable		Not Applicable

4.0 Genetically Modified Organisms and Cell lines

4.1 Will genetic modifications be made to the microorganisms, biological agents, or cells described in Sections 1.0 and 2.0? YES NO If no, please proceed to Section 5.0

4.2 Will genetic modification(s) involving plasmids be done? YES, complete table below NO

Bacteria Used for Cloning *	Plasmid(s) **	Source of Plasmid	Gene Transfected	Describe the change that results from transformation or tranfection

* Please attach a Material Data Sheet or equivalent if available.

** Please attach a plasmid map.

4.3 Will genetic modification(s) of bacteria and/or cells involving viral vectors be made?

YES, complete table below NO

Virus Used for Vector Construction	Vector(s) *	Source of Vector	Gene(s) Transduced	Describe the change that results from transduction

MLE12 cells contain Papovavirus

* Please attach a Material Safety Data Sheet or equivalent.

4.4 Will genetic sequences from the following be involved? No

- ◆ HIV YES, please specify _____ NO
- ◆ HTLV 1 or 2 or genes from any Level 1 or Level 2 pathogens YES, specify _____ NO
- ◆ SV 40 Large T antigen YES NO
- ◆ E1A oncogene YES NO
- ◆ Known oncogenes YES, please specify _____ NO
- ◆ Other human or animal pathogen and or their toxins YES, please specify _____ NO

4.5 Will virus be replication defective? YES NO

4.6 Will virus be infectious to humans or animals? YES NO

4.7 Will this be expected to increase the containment level required? YES NO

5.0 Human Gene Therapy Trials

5.1 Will human clinical trials be conducted involving a biological agent? YES NO
(including but not limited to microorganisms, viruses, prions, parasites or pathogens of plant or animal origin)
If no, please proceed to Section 6.0

5.2 If YES, please specify which biological agent will be used: _____
Please attach a full description of the biological agent.

5.2 Will the biological agent be able to replicate in the host? YES NO

5.3 How will the biological agent be administered? _____

5.4 Please give the Health Care Facility where the clinical trial will be conducted: _____

5.5 Has human ethics approval been obtained? YES, number: _____ NO PENDING

6.0 Animal Experiments

6.1 Will live animals be used? YES NO If no, please proceed to section 7.0

6.2 Name of animal species to be used mice_

6.3 AUS protocol # 2010-272_

6.4 Will any of the agents listed in section 4.0 be used in live animals YES, specify: _____ NO

6.5 Will the agent(s) be shed by the animal: YES NO, please justify:
__LPS is instilled into the lungs of mice. 4 hours later mice are euthanized, this is to short a time for LPS to enter the systemic circulation and be excreted in urine or feces.

7.0 Use of Animal species with Zoonotic Hazards

7.1 Will any animals with zoonotic hazards or their organs, tissues, lavages or other body fluids including blood be used (see list below)? YES No If no, please proceed to section 8.0

7.2 Will live animals be used? YES No

7.3 If yes, please specify the animal(s) used:

- ◆ Pound source dogs YES NO
- ◆ Pound source cats YES NO
- ◆ Cattle, sheep or goats YES, please specify species _____ NO
- ◆ Non-human primates YES, please specify species _____ NO
- ◆ Wild caught animals YES, please specify species & colony # _____ NO
- ◆ Birds YES, please specify species _____ NO
- ◆ Others (wild or domestic) YES, please specify _mouse_____ NO

7.4 If no live animals are used, please specify the source of the specimens:

8.0 Biological Toxins

8.1 Will toxins of biological origin be used? YES NO If no, please proceed to Section 9.0

8.2 If YES, please name the toxin(s) LPS

Please attach information, such as a Material Safety Data Sheet, for the toxin(s) used.

What is the LD₅₀ (specify species) of the toxin 7670 ug/kg IV in mice

8.4 How much of the toxin is handled at one time*? _We purchase 500mg at a time and we aliquot 2 mg which is then suspended in saline and aliquoted into 100uL aliquots. 100uL aliquots are frozen so that personnel can use in animal work.

8.5 How much of the toxin is stored*? _500 mg

8.6 Will any biological toxins be used in live animals? YES, Please provide details: _50 uL (containing 30ug LPS) is instilled intratracheally in mice. Mice are euthanized 4 hours later. NO

*For information on biosecurity requirements, please see:

http://www.uwo.ca/humanresources/docandform/docs/healthandsafety/biosafety/Biosecurity_Requirements.pdf

9.0 Insects

9.1 Do you use insects? YES NO If no, please proceed to Section 10.0

9.2 If YES, please give the name of the species. _____

9.3 What is the origin of the insect? _____

9.4 What is the life stage of the insect? _____

9.5 What is your intention? Initiate and maintain colony, give location: _____
 "One-time" use, give location: _____

9.6 Please describe the risk (if any) of escape and how this will be mitigated:

9.7 Do you use insects that require a permit from the CFIA permit? YES NO
If YES, Please attach the CFIA permit & describe any CFIA permit conditions:

10.0 Plants

10.1 Do you use plants? YES NO If no, please proceed to Section 11.0

10.2 If YES, please give the name of the species. _____

10.3 What is the origin of the plant? _____

10.4 What is the form of the plant (seed, seedling, plant, tree...)? _____

10.5 What is your intention? Grow and maintain a crop "One-time" use

10.6 Do you do any modifications to the plant? YES NO
If yes, please describe: _____

10.7 Please describe the risk (if any) of loss of the material from the lab and how this will be mitigated:

10.8 Is the CFIA permit attached? YES NO
If YES, Please attach the CFIA permit & describe any CFIA permit conditions:

11.0 Import Requirements

11.1 Will any of the above agents be imported? YES, please give country of origin _____ NO
If no, please proceed to Section 12.0

11.2 Has an Import Permit been obtained from HC for human pathogens? YES NO

11.3 Has an import permit been obtained from CFIA for animal or plant pathogens? YES NO

11.4 Has the import permit been sent to OHS? YES, please provide permit # _____ NO

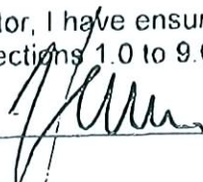
12.0 Training Requirements for Personnel Named on Form

All personnel named on the above form who will be using any of the above named agents are required to attend the following training courses given by OHS:

- ◆ Biosafety
- ◆ Laboratory and Environmental/Waste Management Safety
- ◆ WHMIS (Western or equivalent)
- ◆ Employee Health and Safety Orientation

As the Principal Investigator, I have ensured that all of the personnel named on the form who will be using any of the biological agents in Sections 1.0 to 9.0 have been trained.

SIGNATURE _____



13.0 Containment Levels

13.1 For the work described in sections 1.0 to 9.0, please indicate the highest HC or CFIA Containment Level required. 1 2 2+ 3

13.2 Has the facility been certified by OHS for this level of containment?
X YES, date of most recent biosafety inspection: 2011/02/02 *[Signature]*
 NO, please certify
 NOT REQUIRED for Level 1 containment

13.3 Please indicate permit number (not applicable for first time applicants): _____

14.0 Procedures to be Followed

14.1 Please describe additional risk reduction measures will be taken beyond containment level 1, 2, 2+ or 3 measures, that are unique to this agent.

14.2 Please outline what will be done if there is an exposure to the biological agents listed, such as a needlestick injury or an accidental splash:
Personnel will go immediately to Occupational Health at SJHC

14.3 As the Principal Investigator, I will ensure that this project will follow the Western Biosafety Guidelines and Procedures Manual for Containment Level 1 & 2 Laboratories (and the Level 3 Facilities Manual for Level 3 projects). I will ensure that UWO faculty, staff and students working in my laboratory have an up-to-date Hazard Communication Form, found at <http://www.wph.uwo.ca/>

SIGNATURE *[Signature]*
Date: *Feb 2/11*

15.0 Approvals

1) UWO Biohazards Subcommittee: SIGNATURE: _____
Date: _____

2) Safety Officer for the University of Western Ontario
SIGNATURE: _____
Date: _____

3) Safety Officer for Institution where experiments will take place (if not UWO):
SIGNATURE: *[Signature]*
Date: 2011/02/02

Approval Number: _____ Expiry Date (3 years from Approval): _____

Special Conditions of Approval:

T-11

Subject: Re: Biological Agents Registry Form (Lewis et al)

From: Lynda McCaig <lmccaig@uwo.ca>

Date: Fri, 26 Aug 2011 11:10:28 -0400

To: Jennifer Stanley <jstanle2@uwo.ca>

Hi Jennifer,

The cell line is MLE-12. It is a mouse lung epithelial cell (MLEC). In the summary it states, 'various cell lines (MLE-12 and MLEC) '. This is incorrect as MLE-12 is a specific MLEC.

Would you like us to change the wording in the summary or can you cross out the words 'various' and 'MLEC' from the summary.

Sorry for the confusion, I missed that one completely!!

Lynda

On 26/08/2011 11:02 AM, Jennifer Stanley wrote:

Hi there

I noticed that in the research summary you mention using a cell line called MLEC. This cell line is not listed in Table 2.3. Please clarify.

Regards
Jennifer

E-mail

SIGMA-ALDRICH

Material Safety Data Sheet

Version 3.0
 Revision Date 01/03/2009
 Print Date 05/31/2010

1. PRODUCT AND COMPANY IDENTIFICATION

Product name : Lipopolysaccharides, from *Escherichia coli* 026:B6

Product Number : L3755
 Brand : Sigma

Company : Sigma-Aldrich Canada, Ltd
 2149 Winston Park Drive
 OAKVILLE ON L6H 6J8
 CANADA

Telephone : +19058299500
 Fax : +19058299292
 Emergency Phone # : 800-424-9300

2. COMPOSITION/INFORMATION ON INGREDIENTS

CAS-No.	EC-No.	Index-No.	Concentration
Lipopolysaccharides from <i>Escherichia coli</i> 026:B6			
-	-	-	-

3. HAZARDS IDENTIFICATION

Emergency Overview

Other hazards which do not result in classification
 Pyrogen. May cause fever.

WHMIS Classification

Not WHMIS controlled.

Not WHMIS controlled.

HMIS Classification

Health Hazard: 0
 Flammability: 0
 Physical hazards: 0

Potential Health Effects

Inhalation : May be harmful if inhaled. May cause respiratory tract irritation.
Skin : May be harmful if absorbed through skin. May cause skin irritation.
Eyes : May cause eye irritation.
Ingestion : May be harmful if swallowed.

4. FIRST AID MEASURES

If inhaled

If breathed in, move person into fresh air. If not breathing give artificial respiration

In case of skin contact

Wash off with soap and plenty of water.

In case of eye contact

Flush eyes with water as a precaution.

If swallowed

Never give anything by mouth to an unconscious person. Rinse mouth with water.

5. FIRE-FIGHTING MEASURES**Flammable properties**

Flash point no data available

Ignition temperature no data available

Suitable extinguishing media

Use water spray, alcohol-resistant foam, dry chemical or carbon dioxide.

Special protective equipment for fire-fighters

Wear self contained breathing apparatus for fire fighting if necessary.

6. ACCIDENTAL RELEASE MEASURES**Personal precautions**

Avoid dust formation.

Environmental precautions

Do not let product enter drains.

Methods for cleaning up

Sweep up and shovel. Keep in suitable, closed containers for disposal.

7. HANDLING AND STORAGE**Handling**

Provide appropriate exhaust ventilation at places where dust is formed. Normal measures for preventive fire protection.

Storage

Keep container tightly closed in a dry and well-ventilated place.

Recommended storage temperature: 2 - 8 °C

8. EXPOSURE CONTROLS/PERSONAL PROTECTION

Contains no substances with occupational exposure limit values.

Personal protective equipment**Respiratory protection**

Respiratory protection is not required. Where protection from nuisance levels of dusts are desired, use type N95 (US) or type P1 (EN 143) dust masks. Use respirators and components tested and approved under appropriate government standards such as NIOSH (US) or CEN (EU).

Hand protection

For prolonged or repeated contact use protective gloves.

Eye protection

Safety glasses

Hygiene measures

General industrial hygiene practice.

9. PHYSICAL AND CHEMICAL PROPERTIES

Appearance

Form powder, lyophilized

Safety data

pH no data available

Melting point no data available

Boiling point no data available

Flash point no data available

Ignition temperature no data available

Lower explosion limit no data available

Upper explosion limit no data available

Water solubility no data available

10. STABILITY AND REACTIVITY

Storage stability

Stable under recommended storage conditions.

Hazardous decomposition products

Hazardous decomposition products formed under fire conditions. - Nature of decomposition products not known.

11. TOXICOLOGICAL INFORMATION

Acute toxicity

no data available

Irritation and corrosion

no data available

Sensitisation

no data available

Chronic exposure

IARC: No component of this product present at levels greater than or equal to 0.1% is identified as probable, possible or confirmed human carcinogen by IARC.

Signs and Symptoms of Exposure

To the best of our knowledge, the chemical, physical, and toxicological properties have not been thoroughly investigated.

Potential Health Effects

Inhalation	May be harmful if inhaled. May cause respiratory tract irritation.
Skin	May be harmful if absorbed through skin. May cause skin irritation.
Eyes	May cause eye irritation.
Ingestion	May be harmful if swallowed.

12. ECOLOGICAL INFORMATION

Elimination information (persistence and degradability)

no data available

Ecotoxicity effects

no data available

Further information on ecology

no data available

13. DISPOSAL CONSIDERATIONS

Product

Observe all federal, state, and local environmental regulations.

Contaminated packaging

Dispose of as unused product.

14. TRANSPORT INFORMATION

DOT (US)

Not dangerous goods

IMDG

Not dangerous goods

IATA

Not dangerous goods

15. REGULATORY INFORMATION

DSL Status

This product contains the following components that are not on the Canadian DSL nor NDSL lists.

Lipopolysaccharides from Escherichia coli 026:B6

CAS-No.

-

WHMIS Classification

Not WHMIS controlled.

Not WHMIS controlled.

16. OTHER INFORMATION

Further information

Copyright 2008 Sigma-Aldrich Co. License granted to make unlimited paper copies for internal use only.

The above information is believed to be correct but does not purport to be all inclusive and shall be used only as a guide. The information in this document is based on the present state of our knowledge and is applicable to the product with regard to appropriate safety precautions. It does not represent any guarantee of the properties of the product. Sigma-Aldrich Co., shall not be held liable for any damage resulting from handling or from contact with the above product. See reverse side of invoice or packing slip for additional terms and conditions of sale.



TOXIN USE RISK ASSESSMENT

Name of Toxin:	LPS
Proposed Use Dose:	2000 µg
Proposed Storage Dose:	500000 µg
LD ₅₀ (species):	7670 µg

Calculation:
$7670 \text{ µg/kg} \quad \times \quad 50 \text{ kg/person}$
Dose per person based on LD ₅₀ in µg = 383500
LD ₅₀ per person with safety factor of 10 based on LD ₅₀ in µg = 38350

Comments/Recommendations: The amount of toxin stored exceeds the recommended storage dose.

Info on Cell Line(s)

Cell Biology

ATCC® Number: **CRL-2110™** Price: **\$379.00**

Designations: MLE 12
Depositors: JA Whitsett
Biosafety Level: 2 [CELLS CONTAIN PAPOVAVIRUS]
Shipped: frozen
Medium & Serum: [See Propagation](#)
Growth Properties: adherent
Organism: Mus musculus, transgenic (mouse, transgenic)
Morphology: epithelial

Source: **Organ:** lung
Strain: FVB/N
Cell Type: epithelialSV40 transformed

Cellular Products: lung surfactant proteins B and C (SP-B, SP-C)

Permits/Forms: In addition to the [MTA](#) mentioned above, other [ATCC and/or regulatory permits](#) may be required for the transfer of this ATCC material. Anyone purchasing ATCC material is ultimately responsible for obtaining the permits. Please [click here](#) for information regarding the specific requirements for shipment to your location.

Isolation: **Isolation date:** 1992

Tumorigenic: Yes

Age: 5 month old

Gender: female

Comments: This line was established in 1992 by Kathryn A. Wikenheiser from pulmonary tumors in a mouse transgenic for the SV40 large T antigen under the control of the promoter region of the human surfactant protein C gene.

The cells express the mRNA for large T antigen. Lung surfactant proteins B and C were detected.

The cells secrete phospholipids in response to phorbol esters and ATP but not in response to forskolin.

Propagation:

Related Links ▶

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[Frequently Asked Questions](#)

[Material Transfer Agreement](#)

[Technical Support](#)

[Related Cell](#)

[Culture Products](#)

[BioProducts](#)

[Cell, microbial and molecular genomics products for the life](#)

- [sciences](#)

[BioServices](#)

[Bio-materials management; basic repository to complex partnership-](#)

- [level services](#)

[BioStandards](#)

[Biological Reference Material and Consensus Standards for the life science](#)

- [community](#)