

THE UNIVERSITY OF WESTERN ONTARIO  
BIOLOGICAL AGENTS REGISTRY FORM  
Approved Biohazards Subcommittee: April 9, 2010  
Biosafety Website: [www.uwo.ca/humanresources/biosafety/](http://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 biohazardous 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 biohazards 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 Biohazard Subcommittee. For questions regarding this form, please contact the Biosafety Officer at extension 81135 or [biosafety@uwo.ca](mailto: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/](http://www.uwo.ca/humanresources/biosafety/)

PRINCIPAL INVESTIGATOR

DEPARTMENT

ADDRESS

PHONE NUMBER

EMERGENCY PHONE NUMBER(S)

EMAIL

Dr David O'Gorman  
BIOCHEMISTRY  
268 GROSVENOR ST, LONDON, ON Rm EA-137  
x 64397  
519-471-6457  
[dogorman@uwo.ca](mailto:dogorman@uwo.ca)

Location of experimental work to be carried out: Building(s) LAWSON HEALTH RESEARCH Room(s) F1-104  
INSTITUTE

\*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 12.0, Approvals).

FUNDING AGENCY/AGENCIES: CIMR / Plastic Surgery Education Foundation  
GRANT TITLE(S): Molecular mechanisms of Dupuytren's Disease  
Molecular mechanisms of abnormal scarring

PLEASE ATTACH A BRIEF DESCRIPTION OF YOUR WORK THAT EXPLAINS THE BIOHAZARDS USED AND HOW THEY WILL BE STORED, USED AND DISPOSED OF. PROJECTS SUBMITTED WITHOUT A SUMMARY WILL NOT BE REVIEWED.

Names of all personnel working under Principal Investigators supervision in this location:

BRETT THURLOW ANDREW GOULD  
JUSTIN CRAWFORD  
CHRISTINA RAYKHA

\* DESCRIPTION MUST BE ATTACHED TO THIS FORM OR PROJECT WILL NOT BE REVIEWED\*

**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)? Invitrogen

Please describe the risk (if any) of escape and how this will be mitigated:  
No risk (non-pathogenic e. coli)

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

**New Info**

1.2 Please complete the table below:

Name of Biological agent(s)*	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
E. coli (DH5α)	<input type="radio"/> Yes <input checked="" type="radio"/> No	<input type="radio"/> Yes <input checked="" type="radio"/> No	<input type="radio"/> Yes <input checked="" type="radio"/> No	0.1 L	Invitrogen	<input checked="" type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input checked="" type="radio"/> 3
Lentivirus (viral vector, replication deficient)	<input type="radio"/> Yes <input checked="" type="radio"/> No	<input type="radio"/> Yes <input checked="" type="radio"/> No	<input type="radio"/> Yes <input checked="" type="radio"/> No	0.03 L	Invitrogen	<input type="radio"/> 1 <input checked="" type="radio"/> 2 <input type="radio"/> 2+ <input checked="" 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 checked="" 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 checked="" 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?  YES  NO  
 If no, please proceed to Section 3.0

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 checked="" type="radio"/> Yes <input type="radio"/> No	Surgically resected tissue	Not applicable
Rodent	<input type="radio"/> Yes <input checked="" type="radio"/> No		
Non-human primate	<input type="radio"/> Yes <input checked="" type="radio"/> No		
Other (specify)	<input type="radio"/> Yes <input checked="" type="radio"/> No		

\* DESCRIPTION MUST BE ATTACHED TO THIS FORM OR PROJECT WILL NOT BE REVIEWED\*  
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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)*	Supplier / Source
Human	<input checked="" type="radio"/> Yes <input type="radio"/> No	HaCaT	CLS - Germany
Rodent	<input checked="" type="radio"/> Yes <input type="radio"/> No	NIH 3T3	ATCC - USA
Non-human primate	<input type="radio"/> Yes <input checked="" type="radio"/> No		
Other (specify)	<input type="radio"/> Yes <input checked="" 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 types(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/NO	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"/> No <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		<input type="radio"/> Yes <input type="radio"/> No <input type="radio"/> Unknown		<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 2+ <input type="radio"/> 3
Human Organs or Tissues (unpreserved)	Surgically resected tissue (in-house surgeries)	<input type="radio"/> Yes <input checked="" type="radio"/> No <input type="radio"/> Unknown	---	<input type="radio"/> 1 <input checked="" 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

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

\* DESCRIPTION MUST BE ATTACHED TO THIS FORM OR PROJECT WILL NOT BE REVIEWED\*

4.3 Will genetic modification(s) 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
Lentivirus	pLent:Ubc	Invitrogen	POSTN	<del>constitutive</del> constitutive expression of periostin, a cell adhesion molecule <del>expressed</del> in the ECM found

\* Please attach a Material Safety Data Sheet or equivalent.

4.4 Will genetic sequences from the following be involved?

- HIV  YES, please specify gag/pol/Rev  NO
- HTLV 1 or 2 or genes from any Level 1 or Level 2 pathogens  YES, specify VSV-G  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 \_\_\_\_\_

6.3 AUS protocol # \_\_\_\_\_

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:  
\_\_\_\_\_  
\_\_\_\_\_

\* DESCRIPTION MUST BE ATTACHED TO THIS FORM OR PROJECT WILL NOT BE REVIEWED\*

**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 Please specify the animal(s) used:

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

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

8.3 What is the LD<sub>50</sub> (specify species) of the toxin \_\_\_\_\_

8.4 How much of the toxin is handled at one time\*? \_\_\_\_\_

8.5 How much of the toxin is stored\*? \_\_\_\_\_

8.6 Will any biological toxins be used in live animals?  YES, Please provide details: \_\_\_\_\_  NO

\*For information on biosecurity requirements, please see:

[http://www.uwo.ca/humanresources/docandform/docs/healthandsafety/biosafety/Biosecurity\\_Requirements.pdf](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 biohazardous 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. O 1 O 2  2+ O 3

13.2 Has the facility been certified by OHS for this level of containment?  
 YES, permit # if on-campus BIO-LHRI-0052 *re-certification 17/08/2010*  
 NO, please certify  
 NOT REQUIRED for Level 1 containment

**14.0 Procedures to be Followed**

14.1 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 *Don Dezan* Date: 20 July 2010

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

14.3 Please outline what will be done if there is an exposure to the biohazards listed, such as a needlestick injury:  
Individual will report to OH+S, SJHC for care + follow-up.

**15.0 Approvals**

1) UWO Biohazard 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: August 17/2010

Approval Number: \_\_\_\_\_ Expiry Date (3 years from Approval): \_\_\_\_\_

Special Conditions of Approval:

## Description of Work

In brief, our work revolves around the culture of primary tissue explants for the purpose of generating human cells capable of acting as proxy models for a disease state of interest (specifically Dupuytren's Contracture and / or abnormal scarring phenotypes).

We pursue this research through the CL-2 approved culture of tissue explants from surgical resections (done in-house at the Hand and Upper Limb Clinic, St. Joseph's Health Care) and subsequent manipulations of those cultured cells in downstream applications typical of molecular biology research. All samples received from the surgical team have been screened by the referring physician for known infectious diseases and are flagged as such. Regardless, all samples received in the lab are treated as potentially containing infectious materials and are handled with appropriate precautions.

For applications and investigations not suited to the use of primary cell cultures, we also maintain immortalized cell lines (HaCaT keratinocytes and NIH 3T3 fibroblasts) obtained from commercial sources. All cell cultures are stored in approved liquid nitrogen-containing cell storage units when not in active culture

In all cases, cell culture takes place entirely within an approved and inspected CL-2 facility, is carried out only by trained individuals, and is maintained in accordance with published biosafety protocols. Any waste generated is decontaminated with 10% (final volume) bleach prior to disposal in the institutional biohazard waste stream. Virtually all equipment used in cell culture is disposable by nature, and is treated as any other waste as required.

In addition to the use of primary cell cultures, basic molecular investigation in the lab requires the use of standard practices of gene manipulation, including the use of non-pathogenic E. coli bacterial cultures for the purpose of amplification and manipulation of various genetic sequences of interest. Such cultures are commercially obtained and cultured in accordance with standard protocols. Waste or spills generated are decontaminated with 10% (final volume) bleach prior to disposal in the institutional biohazard waste stream. Equipment and glassware used in such cultures is decontaminated with bleach, washed and re-sterilised on-site.

Finally, though not taking place in the lab currently, research requirements dictate the generation of genetically modified organisms through viral transduction of primary cells (at some point in the near future). Such approaches are required when using primary culture due to the demonstrated inefficiency of plasmid transfection.

The current intention is to use a commercially available system for generation of a replication defective lentiviral vector containing our gene(s) of interest. This system (ViraPower from Invitrogen) is already in use in the institute, and our lab technician (Andrew Gould) has experience with the system and with the necessary safety protocols required for operating in a CL-2 environment with CL-3 precautions (previously established, in consultation with the on-site Safety Officer, as the required level of protection for work with this system).

Waste generated from this future objective will be disposed of in accordance with established protocols – bleach decontamination of all waste prior to removal from the culture hood, followed by immediate autoclave sterilization prior to disposal in the institutional biohazard waste stream.

concentration

JF.

MATERIAL SAFETY DATA SHEET

LIBRARY EFFICIENCY DHSALPHA COMPETENT CELLS  
 INVITROGEN CORPORATION  
 MSDS ID: 18263

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 Revised 9/30/03  
 Replaces 9/05/03  
 Printed 9/30/03

**1. PRODUCT AND COMPANY INFORMATION**

INVITROGEN CORPORATION  
 1600 FARADAY AVE.  
 CARLSBAD, CA 92008  
 760/603-7200

GIBCO PRODUCTS  
 INVITROGEN CORPORATION  
 3175 STALEY ROAD P.O. BOX 68  
 GRAND ISLAND, NY 14072  
 716/774-6700

INVITROGEN CORPORATION  
 3 FOUNTAIN DR.  
 INCHINNAN BUSINESS PARK  
 PAISLEY, PA4 9RF  
 SCOTLAND  
 44-141 814-6100

INVITROGEN CORPORATION  
 P.O. BOX 12-502  
 PENROSE  
 AUCKLAND 1135  
 NEW ZEALAND  
 64-9-579-3024

INVITROGEN CORPORATION  
 2270 INDUSTRIAL ST.  
 BURLINGTON, ONT  
 CANADA L7P 1A1  
 905/335-2255

EMERGENCY NUMBER (SPILLS, EXPOSURES): 301/431-8585 (24 HOUR)  
 800/451-8346 (24 HOUR)  
 800/955-6288

NON-EMERGENCY INFORMATION:

Product Name: LIBRARY EFFICIENCY DHSALPHA COMPETENT CELLS  
 Stock Number: 18263012

NOTE: If this product is a kit or is supplied with more than one material, please refer to the MSDS for each component for hazard information.

Product Use:  
 These products are for laboratory research use only and are not intended for human or animal diagnostics, therapeutic, or other clinical uses.

Synonyms:  
 Not available.

**2. COMPOSITION, INFORMATION ON INGREDIENTS**

The following list shows components of this product classified as hazardous based on physical properties and health effects:

Component	CAS No.	Percent
DIMETHYL SULFOXIDE	67-68-5	3 - 7

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**3. HAZARDS IDENTIFICATION**

\*\*\*\*\* EMERGENCY OVERVIEW \*\*\*\*\*  
 Warning!  
 Irritant.  
 Harmful if absorbed.  
 \*\*\*\*\*

Potential Health Effects:

Eye:  
 Can cause moderate irritation, tearing and reddening, but not likely to permanently injure eye tissue.

Skin:  
 Can cause moderate skin irritation, defatting, and dermatitis. Not likely to cause permanent damage.  
 Upon prolonged or repeated exposure, harmful if absorbed through the skin.  
 May cause minor systemic damage.

Inhalation:

Can cause moderate respiratory irritation, dizziness, weakness, fatigue, nausea and headache.  
 No toxicity expected from inhalation.

Ingestion:

Irritating to mouth, throat, and stomach. Can cause abdominal discomfort, nausea, vomiting and diarrhea.

Chronic:

No data on cancer.

**4. FIRST AID MEASURES**

Eye:

Flush eyes with plenty of water for at least 20 minutes retracting eyelids often. Tilt the head to prevent chemical from transferring to the uncontaminated eye. Get immediate medical attention.

Skin:

Wash with soap and water. Get medical attention if irritation develops or persists.

Inhalation:

Remove to fresh air. If breathing is difficult, have a trained individual administer oxygen. If not breathing, give artificial respiration and have a trained individual administer oxygen. Get medical attention immediately.

Ingestion:

Do not induce vomiting and seek medical attention immediately. Drink two

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**4. FIRST AID MEASURES (CONT.)**

glasses of water or milk to dilute. Provide medical care provider with this MSDS.

Note To Physician:  
Treat symptomatically.

**5. FIRE FIGHTING MEASURES**

Flashpoint Deg C: Not available.  
Upper Flammable Limit %: Not available.  
Lower Flammable Limit %: Not available.  
Autoignition Temperature Deg C: Not available.

Extinguishing Media:  
Use alcohol resistant foam, carbon dioxide, dry chemical, or water spray when fighting fires. Water or foam may cause frothing if liquid is burning but it still may be a useful extinguishing agent if carefully applied to the fire. Do not direct a water stream directly into the hot burning liquid. DMSO undergoes a violent exothermic reaction on mixing with copper wool and trichloroacetic acid. On mixing with potassium permanganate it will flash instantaneously. It reacts violently with: acid halides, cyanuric chloride, silicon tetrachloride, phosphorus trichloride and trioxide, thionyl chloride, magnesium perchlorate, silver fluoride, methyl bromide, iodine pentafluoride, nitrogen periodate, diborane, sodium hydride, perchloric and periodic acids. When heated above its boiling point, DMSO degrades giving off formaldehyde, methyl mercaptan, and sulfur dioxide.

**Firefighting Techniques/Equipment:**

Do not enter fire area without proper protection including self-contained breathing apparatus and full protective equipment. Fight fire from a safe distance and a protected location due to the potential of hazardous vapors and decomposition products.

Hazardous Combustion Products:  
Carbon dioxide Carbon monoxide Sulfur containing gases

**6. ACCIDENTAL RELEASE MEASURES**

Accidental releases may be subject to special reporting requirements and other regulatory mandates. Refer to Section 8 for personal protection equipment recommendations.

**6. ACCIDENTAL RELEASE MEASURES (CONT.)**

Spill Cleanup:  
 Exposure to the spilled material may be irritating or harmful. Follow personal protective equipment recommendations found in Section VIII of this MSDS. Additional precautions may be necessary based on special circumstances created by the spill including; the material spilled, the quantity of the spill, the area in which the spill occurred. Also consider the expertise of employees in the area responding to the spill. Ventilate the contaminated area. Absorb spill. Common absorbent materials should be effective. Deposit in appropriate containers for removal and disposal.

**7. HANDLING AND STORAGE**

Storage of some materials is regulated by federal, state, and/or local laws.

Storage Pressure:  
 Ambient

Handling Procedures:

Harmful or irritating material. Avoid contacting and avoid breathing the material. Use only in a well ventilated area. Keep closed or covered when not in use.

Storage Procedures:

Store in a cool dry ventilated location. Isolate from incompatible materials and conditions. Keep container(s) closed. Suitable for most general chemical storage areas.

**8. EXPOSURE CONTROLS, PERSONAL PROTECTION**

Exposure Limits:

Component DIMETHYL SULFOXIDE	OSHA PEL (ppm) Not established.	AGCIH TWA (ppm) Not established.
---------------------------------	---------------------------------------	--

Engineering Controls:

Local exhaust ventilation or other engineering controls are normally required when handling or using this product to avoid overexposure.

Personal Protective Equipment:

Eye:  
 Safety glasses should be the minimum eye protection.  
 Wear chemically resistant safety glasses with side shields when handling this product. Wear additional eye protection such as chemical splash

MATERIAL SAFETY DATA SHEET

LIBRARY EFFICIENCY DHSALPHA COMPETENT CELLS  
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### 8. EXPOSURE CONTROLS, PERSONAL PROTECTION (CONT.)

goggles and/or face shield when the possibility exists for eye contact with splashing or spraying liquid, or airborne material. Do not wear contact lenses. Have an eye wash station available.

**Skin:**  
 Avoid skin contact by wearing chemically resistant gloves, an apron and other protective equipment depending upon conditions of use. Inspect gloves for chemical break-through and replace at regular intervals. Clean protective equipment regularly. Wash hands and other exposed areas with mild soap and water before eating, drinking, and when leaving work. Gloves should be used as minimum hand protection.

**Respiratory:**  
 Use supplied-air respiratory equipment as required.

### 9. PHYSICAL AND CHEMICAL PROPERTIES

**Appearance/physical state:** Liquid solution / suspension  
**Odor:** No odor.

Not established.  
 Not established.

**Specific Gravity/Density:** Not established.  
**Octanol/water Partition Coeff:** Not established.  
**Volatiles:** Not established.  
**Evaporation Rate:** Not established.  
**Viscosity:** Not established.

### 10. STABILITY AND REACTIVITY

**Stability:**  
 Stable under normal conditions.

**Conditions to Avoid:**  
 Strong oxidizing agents. Temperatures above the high flash point of this combustible material in combination with sparks, open flames, or other sources of ignition. Strong alkalies. DMSO undergoes a violent exothermic reaction on mixing with copper wool and trichloroacetic acid. On mixing with potassium permanganate it will flash instantaneously. It reacts violently with: acid halides, cyanuric chloride, silicon tetrachloride, phosphorus trichloride and trioxide, thionyl chloride, magnesium perchlorate, silver fluoride, methyl bromide, iodine pentafluoride, nitrogen peroxide, diborane, sodium hydride, perchloric and periodic acids. When heated above its boiling point, DMSO

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10. STABILITY AND REACTIVITY (CONT.)

degrades giving off formaldehyde, methyl mercaptan, and sulfur dioxide.

Hazardous Decomposition Products:  
 Carbon monoxide. Carbon dioxide. Sulfur containing gases.

Hazardous Polymerization:  
 Hazardous polymerization will not occur.

11. TOXICOLOGICAL INFORMATION

Acute Toxicity:

Dermal/Skin:  
 DIMETHYL SULFOXIDE: 40 GM/KG

Inhalation/Respiratory:  
 Not determined.

Oral/Ingestion:  
 DIMETHYL SULFOXIDE: 14,500 MG/KG

Target Organs: Blood. Eyes. Skin.

Carcinogenicity:  
 NTP:  
 Not tested.

IARC:  
 Not listed.

OSHA:  
 Not regulated.

Other Toxicological Information

12. Ecological Information

Ecotoxicological Information: No ecological information available.

Environmental Fate (Degradation, Transformation, and Persistence):  
 Bioconcentration is not expected to occur.  
 Biodegrades slowly.

MATERIAL SAFETY DATA SHEET

LIBRARY EFFICIENCY DHSALPHA COMPETENT CELLS  
 INVITROGEN CORPORATION  
 MSDS ID: 18263

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13. DISPOSAL CONSIDERATIONS

Regulatory Information:  
 Not applicable.

Disposal Method:  
 Clean up and dispose of waste in accordance with all federal, state, and local environmental regulations.  
 Dispose of by incineration following Federal, State, Local, or Provincial regulations.

14. TRANSPORT INFORMATION

Proper Shipping Name: Not Determined.  
 Subsidiary Hazards:

15. REGULATORY INFORMATION

UNITED STATES:

TSCA:  
 This product is solely for research and development purposes only and may not be used, processed or distributed for a commercial purpose. It may only be handled by technically qualified individuals.

Prop 65 Listed Chemicals: PROP 65 PERCENT  
 No Prop 65 Chemicals.

NO 313 Chemicals

CANADA:

DSL/NDSL:  
 Not determined.

COMPONENT WHMIS Classification  
 DIMETHYL SULFOXIDE D2B

EUROPEAN UNION:

PRODUCT RISK PHRASES: None assigned.  
 PRODUCT SAFETY PHRASES: Not applicable.  
 PRODUCT CLASSIFICATION:

MATERIAL SAFETY DATA SHEET

LIBRARY EFFICIENCY DH5ALPHA COMPETENT CELLS  
 INVITROGEN CORPORATION  
 MSDS ID: 18263

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**15. REGULATORY INFORMATION (CONT.)**

Not classified

Component  
 DIMETHYL SULFOXIDE

EINECS  
 Number  
 200-664-3

**16. OTHER INFORMATION**

HMS Rating 0-4:  
 FIRE: Not determined.  
 HEALTH: Not determined.  
 REACTIVITY: Not determined.

- Abbreviations
- N/A - Data is not applicable or not available
  - SARA - Superfund and Reauthorization Act
  - HMSIS - Hazard Material Information System
  - WHMIS - Workplace Hazard Materials Information System
  - NTP - National Toxicology Program
  - OSHA - Occupational Health and Safety Administration
  - IARC - International Agency for Research on Cancer
  - PROP 65 - California Safe Drinking Water and Toxic Enforcement Act of 1986
  - EINECS - European Inventory of Existing Commercial Chemical Substances

The above information was acquired by diligent search and/or investigation and the recommendations are based on prudent application of professional judgment. The information shall not be taken as being all inclusive and is to be used only as a guide. All materials and mixtures may present unknown hazards and should be used with caution. Since Invitrogen Corporation cannot control the actual methods, volumes, or conditions of use, the Company shall not be held liable for any damages or losses resulting from the handling or from contact with the product as described herein. THE INFORMATION IN THIS MSDS DOES NOT CONSTITUTE A WARRANTY, EXPRESS OR IMPLIED, INCLUDING ANY IMPLIED WARRANTY OF MERCHANTABILITY OR FITNESS FOR ANY PARTICULAR PURPOSE.

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## HaCaT

Cell name	Description	Order no.	Units	Price, Euro
HaCaT	Human keratinocyte cell line	300493	cryovial	430,00
HaCaT	Human keratinocyte cell line	330493	vital	490,00

### P. Boukamp



**Boukamp P, et al.** (3.7 MB)

Normal keratinization in a spontaneously immortalized aneuploid human keratinocyte cell line.

Designation:	HaCaT
Depositor:	DKFZ, Heidelberg
Organism:	Homo sapiens (human)
Ethnicity:	Caucasian
Age/Stage:	62 years
Gender:	male
Tissue:	Skin
Celltype:	keratinocyte
Growth	monolayer
Properties:	
Description:	in vitro spontaneously transformed keratinocytes from histologically normal skin.
Culture	DMEM medium (high glucose) supplemented with 2 mM L-glutamine and 10% fetal calf serum.
Medium:	
Subculturing:	Remove medium, rinse with 0.05% EDTA, add 0.05% EDTA solution and incubate for 10 min at 37°C. Take off EDTA, add fresh 0.05% trypsin/0.025% EDTA solution (final concentrations) and let culture sit at 37°C until the cells detach (approx. 5 minutes). Add fresh medium, aspirate and dispense into new flasks.
Split Ratio:	A ratio of 1:5 to 1:10 is recommended
Fluid	2 times weekly
Renewal:	
Freeze	CM-1 (CLS - Cell Lines Service)
Medium:	
Sterility:	Tests for mycoplasma, bacteria and fungi were negative
Biosafety	1
Level:	
Tumorigenic:	no
Karyotype:	Aneuploid (hypotetraploid)

### References:

Boukamp P, Dzarlieva-Petrusevska RT, Breitkreuz D, Hornung J, Markham A, Fusenig NE. Normal keratinization in a spontaneously immortalized aneuploid human keratinocyte cell line. *J. Cell Biol.* 106: 761-771, 1988.

Boukamp P, Popp S, Altmeyer S, Hülsen A, Fasching C, Cremer T, Fusenig NE. Sustained nontumorigenic phenotype correlates with a largely stable chromosome content during long-term culture of the human keratinocyte line HaCat. *Genes, Chromosomes and Cancer* 19: 201-214, 1997.

**Cell Line Designation: NIH/ 3T3****ATCC Catalog No. CRL-1658™****Table of Contents:**

- Cell Line Description
- Biosafety Level
- Use Restrictions
- Handling Procedure for Frozen Cells
- Handling Procedure for Flask Cultures
- Subculturing Procedure
- Medium Renewal Procedure
- Complete Growth Medium
- Cryoprotectant Medium
- References
- Replacement Policy
- Specific Batch Information

**Cell Line Description****Organism:** *Mus musculus* (mouse)**Strain:** NIH/Swiss**Tissue:** embryo**Morphology:** fibroblast**Growth properties:** adherent**VirusSuscept:** murine sarcoma viruses; murine leukemia viruses**Depositors:** S.A. Aaronson

**Comments:** The NIH/3T3, a continuous cell line of highly contact-inhibited cells was established from NIH Swiss mouse embryo cultures in the same manner as the original random bred 3T3 (ATCC CCL-92™) and the inbred BALB/c 3T3 (ATCC CCL-163™). The established NIH/3T3 line was subjected to more than 5 serial cycles of subcloning in order to develop a subclone with morphologic characteristics best suited for transformation assays. These cells are useful for DNA transfection and transformation studies.

Tested and found negative for ectromelia virus (mousepox).

**Biosafety Level: 1**

Appropriate safety procedures should always be used with this material. Laboratory safety is discussed in the following publication: *Biosafety in Microbiological and Biomedical Laboratories*, 4th ed. HHS Publication No. (CDC) 93-8395. U.S. Department of Health and Human Services, Centers for Disease Control and Prevention. Washington DC: U.S. Government Printing Office; 1999. The entire text is available online at [www.cdc.gov/od/ohs/biosfty/bmbl4/bmbl4toc.htm](http://www.cdc.gov/od/ohs/biosfty/bmbl4/bmbl4toc.htm).

**Use Restrictions**

These cells are distributed for research purposes only. ATCC recommends that individuals contemplating commercial use of any cell line first contact the originating investigator to negotiate an agreement. Third party distribution of this cell line is discouraged, since this practice has resulted in the unintentional spreading of cell lines contaminated with inappropriate animal cells or microbes.

**Handling Procedure for Frozen Cells**

To insure the highest level of viability, thaw the vial and initiate the culture as soon as possible upon receipt. If upon arrival, continued storage of the frozen culture is necessary, it should be stored in liquid nitrogen vapor phase and not at -70°C. Storage at -70°C will result in loss of viability.

**SAFETY PRECAUTION: ATCC highly recommends that protective gloves and clothing always be used and a full face mask always be worn when handling frozen vials. It is important to note that some vials leak when submersed in liquid nitrogen and will slowly fill with liquid nitrogen. Upon thawing, the conversion of the liquid nitrogen back to its gas phase may result in the vessel exploding or blowing off its cap with dangerous force creating flying debris.**

1. Thaw the vial by gentle agitation in a 37°C water bath. To reduce the possibility of contamination, keep the O-ring and cap out of the water. Thawing should be rapid (approximately 2 minutes).
2. Remove the vial from the water bath as soon as the contents are thawed, and decontaminate by dipping in or spraying with 70% ethanol. *All of the operations from this point on should be carried out under strict aseptic conditions.*
3. Transfer the vial contents to a centrifuge tube containing 9.0 ml complete growth medium and spin at approximately 125 xg for 5 to 7 minutes.
4. Resuspend cell pellet with the recommended complete growth medium (see the specific batch information for the culture recommended dilution ratio) and dispense into a 25 cm<sup>2</sup> or a 75 cm<sup>2</sup> culture flask. *It is important to avoid excessive alkalinity of the medium during recovery of the cells. It is suggested that, prior to the addition of the vial contents, the culture vessel containing the complete growth medium be placed into the incubator for at least 15 minutes to allow the medium to reach its normal pH (7.0 to 7.6).*
5. Incubate the culture at 37°C in a suitable incubator. A 5% CO<sub>2</sub> in air atmosphere is recommended if using the medium described on this product.

**Handling Procedure For Flask Cultures**

The flask was seeded with cells (see specific batch information) grown and completely filled with medium at ATCC to prevent loss of cells during shipping.

1. Upon receipt visually examine the culture for macroscopic evidence of any microbial contamination. Using an inverted microscope (preferably equipped with phase-contrast optics), carefully check for any evidence of microbial contamination. Also check to determine if the majority of cells are still attached to the bottom of the flask; during shipping the cultures are sometimes

handled roughly and many of the cells often detach and become suspended in the culture medium (but are still viable).

- If the cells are still attached, aseptically remove all but 5 to 10 ml of the shipping medium. The shipping medium can be saved for reuse. Incubate the cells at 37°C in a 5% CO<sub>2</sub> in air atmosphere until they are ready to be subcultured.
- If the cells are not attached, aseptically remove the entire contents of the flask and centrifuge at 125 xg for 5 to 10 minutes. Remove shipping medium and save. Resuspend the pelleted cells in 10 ml of this medium and add to 25 cm<sup>2</sup> flask. Incubate at 37°C in a 5% CO<sub>2</sub> in air atmosphere until cells are ready to be subcultured.

### Subculturing Procedure

**Never allow the culture to become completely confluent. Subculture at 80% confluency or less.**

Volumes used in this protocol are for 75 cm<sup>2</sup> flask; proportionally reduce or increase amount of dissociation medium for culture vessels of other sizes.

- Remove and discard culture medium.
- Briefly rinse the cell layer with 0.25% (w/v) Trypsin-0.53mM EDTA solution to remove all traces of serum which contains trypsin inhibitor.
- Add 2.0 to 3.0 ml of Trypsin-EDTA solution to flask and observe cells under an inverted microscope until cell layer is dispersed (usually within 5 to 10 minutes).

**Note:** To avoid clumping do not agitate the cells by hitting or shaking the flask while waiting for the cells to detach. Cells that are difficult to detach may be placed at 37°C to facilitate dispersal.

- Add 6.0 to 8.0 ml of complete growth medium and aspirate cells by gently pipetting.
- Add appropriate aliquots of the cell suspension to new culture vessels. Use 3-5 x 10<sup>3</sup> cells/cm<sup>2</sup> and subculture about every 3 days.

**Note:** In order to maintain this property of **high contact inhibition** it is necessary to transfer routinely at only high dilutions, otherwise variants tend to be selected having reduced contact inhibition. Such low density make culture vessels appear sparse and cell growth sensitive to sub-optimal temperature and media conditions.

- Incubate cultures at 37°C.

**Note:** For more information on enzymatic dissociation and subculturing of cell lines consult Chapter 10 in *Culture of Animal Cells, a manual of Basic Technique* by R. Ian Freshney, 3rd edition, published by Alan R. Liss, N.Y., 1994.

### Medium Renewal

Two times per week.

### Complete Growth Medium

The base medium for this cell line is ATCC-formulated Dulbecco's Modified Eagle's Medium, Catalog No. 30-2002. To make the complete growth medium, add the following components to the base medium:

- bovine calf serum to a final concentration of 10%

This medium is formulated for use with a 5% CO<sub>2</sub> in air atmosphere. (Standard DMEM formulations contain 3.7 g/L sodium bicarbonate and a 10% CO<sub>2</sub> in air atmosphere is then recommended).

The calf serum initially employed and found to be satisfactory was from the Colorado Serum Co. Denver.

### Cryoprotectant Medium

Complete growth medium described above supplemented with 5% (v/v) DMSO.

Cell culture tested DMSO is available as ATCC Catalog No. 4-X.

### Additional Information

Additional product and technical information can be obtained from the catalog references and the ATCC Web site at [www.atcc.org](http://www.atcc.org), or by e-mail at [tech@atcc.org](mailto:tech@atcc.org).

### References

(additional references are available in the catalog at [www.atcc.org](http://www.atcc.org))

Copeland NG and Cooper GM. **Transfection by exogenous and endogenous murine retrovirus DNAs.**

Cell 16: 347-356, 1979 PubMed: 79211204

Loffler S et al. **CD9, a tetraspan transmembrane protein, renders cells susceptible to canine distemper virus.** J. Virol. 71: 42-49, 1997 PubMed: 97138295

Berson JF et al. **A seven-transmembrane domain receptor involved in fusion and entry of T-cell-tropic human immunodeficiency virus type 1 strains.** J. Virol. 70: 6288-6295, 1996 PubMed: 96323150

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Siess DC et al. **Exceptional fusogenicity of chinese hamster ovary cells with murine retrovirus suggests roles for cellular factor(s) and receptor clusters in the membrane fusion process.** J. Virol. 70: 3432-439, 1996 PubMed: 96211474

Jang SI et al. **Activator protein 1 activity is involved in the regulation of the cell type-specific expression from the proximal promoter of the human profilaggrin gene.** J. Biol. Chem. 271: 24105-24114, 1996 PubMed: 96394543

Medin JA et al. **Correction in trans for Fabry disease: expression, secretion, and uptake of alpha-**

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galactosidase A in patient-derived cells driven by a high-titer recombinant retroviral vector. Proc. Natl. Acad. Sci. USA 93: 7917-7922, 1996 PubMed: 96353919

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Shisler J et al. **Induction of susceptibility to tumor necrosis factor by E1A is dependent on binding to either p300 or p105-Rb and induction of DNA synthesis.** J. Virol. 70: 68-77, 1996 PubMed: 96099415

Cavanaugh VJ et al. **Murine cytomegalovirus with a deletion of genes spanning HindIII-J and -I displays altered cell and tissue tropism.** J. Virol. 70: 1365-1374, 1996 PubMed: 96190530

Westerman KA and Leboulch P. **Reversible immortalization of mammalian cells mediated by retroviral transfer and site-specific recombination.** Proc. Natl. Acad. Sci. USA 93: 8971-8976, 1996 PubMed: 96392350

Jainchill J.L. et al. (1969), **Murine sarcoma and leukemia viruses: assay using clonal lines of contact-inhibited mouse cells.** J. Virol. 4:549-553. PubMed:70064316.

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Caputo, J. L., **Biosafety procedures in cell culture.** J. Tissue Culture Methods 11:223-227, 1988.

Fleming, D.O., Richardson, J. H., Tulis, J.J. and Vesley, D., (1995) **Laboratory Safety: Principles and Practice.** Second edition, ASM press, Washington, DC.

### ATCC Warranty

The viability of ATCC products is warranted for 30 days from the date of shipment. If you feel there is a problem with this product, contact Technical Services by phone at 800-638-6597 (U.S., Canada, and Puerto Rico) or 703-365-2700 (elsewhere) or by e-mail at [tech@atcc.org](mailto:tech@atcc.org).

### Disclaimers

This product is intended for laboratory research purposes only. It is not intended for use in humans.

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Please see the enclosed Material Transfer Agreement (MTA) for further details regarding the use of this product. The MTA is also available on our Web site at [www.atcc.org](http://www.atcc.org).

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### American Type Culture Collection

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Fax: 703-365-2750

E-mail: [tech@atcc.org](mailto:tech@atcc.org)

Or contact your local distributor.

**1. IDENTIFICATION OF THE SUBSTANCE/PREPARATION AND THE COMPANY/UNDERTAKING**

**Product code** 351275  
**Product name** VIRAPOWER PKG. MIX 195 UG, LYOPHILIZED

**Contact manufacturer**  
 INVITROGEN CORPORATON  
 1600 FARADAY AVENUE  
 PO BOX 6482  
 CARLSBAD, CA 92008  
 760-603-7200

INVITROGEN CORPORATION  
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 BURLINGTON, ONT  
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 800-263-6236

GIBCO PRODUCTS  
 INVITROGEN CORPORATION  
 3175 STALEY ROAD P.O. BOX 68  
 GRAND ISLAND, NY 14072  
 716-774-6700

**2. COMPOSITION/INFORMATION ON INGREDIENTS**

**Hazardous/Non-hazardous Components**

The product contains no substances which at their given concentration, are considered to be hazardous to health

**3. HAZARDS IDENTIFICATION**

**Emergency Overview**

The product contains no substances which at their given concentration, are considered to be hazardous to health.

Form  
Solid

**Principle Routes of Exposure/**

**Potential Health effects**

<b>Eyes</b>	No information available
<b>Skin</b>	No information available
<b>Inhalation</b>	No information available
<b>Ingestion</b>	No information available

**Specific effects**

<b>Carcinogenic effects</b>	No information available
<b>Mutagenic effects</b>	No information available
<b>Reproductive toxicity</b>	No information available

Sensitization No information available

Target Organ Effects No information available

#### 4. FIRST AID MEASURES

Skin contact	Wash off immediately with plenty of water
Eye contact	Rinse thoroughly with plenty of water, also under the eyelids.
Ingestion	Never give anything by mouth to an unconscious person
Inhalation	Move to fresh air
Notes to physician	Treat symptomatically

#### 5. FIRE-FIGHTING MEASURES

Suitable extinguishing media	Dry chemical
Special protective equipment for firefighters	Wear self-contained breathing apparatus and protective suit

#### 6. ACCIDENTAL RELEASE MEASURES

Personal precautions	Use personal protective equipment
Methods for cleaning up	Soak up with inert absorbent material

#### 7. HANDLING AND STORAGE

Handling	No special handling advice required
Storage	Keep in properly labelled containers

#### 8. EXPOSURE CONTROLS / PERSONAL PROTECTION

##### Occupational exposure controls

##### Exposure limits

Engineering measures Ensure adequate ventilation, especially in confined areas

##### Personal protective equipment

Respiratory protection	In case of insufficient ventilation wear suitable respiratory equipment
Hand protection	Protective gloves
Eye protection	Safety glasses with side-shields
Skin and body protection	Lightweight protective clothing
Hygiene measures	Handle in accordance with good industrial hygiene and safety practice
Environmental exposure controls	Prevent product from entering drains

#### 9. PHYSICAL AND CHEMICAL PROPERTIES

##### General Information

Form Solid

##### Important Health Safety and Environmental Information

Boiling point/range	°C No data available	°F No data available
Melting point/range	°C No data available	°F No data available
Flash point	°C No data available	°F No data available
Autoignition temperature	°C No data available	°F No data available
Oxidizing properties	No information available	

Water solubility

No data available

## 10. STABILITY AND REACTIVITY

Stability	Stable.
Materials to avoid	No information available
Hazardous decomposition products	No information available
Polymerization	Hazardous polymerisation does not occur

## 11. TOXICOLOGICAL INFORMATION

### Acute toxicity

### Principle Routes of Exposure/

### Potential Health effects

Eyes	No information available
Skin	No information available
Inhalation	No information available
Ingestion	No information available

### Specific effects

Carcinogenic effects	No information available
Mutagenic effects	No information available
Reproductive toxicity	No information available
Sensitization	No information available

### Target Organ Effects

No information available

## 12. ECOLOGICAL INFORMATION

Ecotoxicity effects	No information available.
Mobility	No information available.
Biodegradation	Inherently biodegradable.
Bioaccumulation	Does not bioaccumulate.

## 13. DISPOSAL CONSIDERATIONS

Dispose of in accordance with local regulations

## 14. TRANSPORT INFORMATION

### IATA

Proper shipping name	Not classified as dangerous in the meaning of transport regulations
Hazard Class	No information available
Subsidiary Class	No information available
Packing group	No information available
UN-No	No information available

## 15. REGULATORY INFORMATION

### International Inventories

### U.S. Federal Regulations

#### **SARA 313**

Not regulated

#### **Clean Air Act, Section 112 Hazardous Air Pollutants (HAPs) (see 40 CFR 61)**

This product contains the following HAPs:

### U.S. State Regulations

#### **California Proposition 65**

This product contains the following Proposition 65 chemicals:

#### **WHMIS hazard class:**

Non-controlled

This product has been classified according to the hazard criteria of the CPR and the MSDS contains all of the information required by the CPR

## 16. OTHER INFORMATION

This material is sold for research and development purposes only. It is not for any human or animal therapeutic or clinical diagnostic use. It is not intended for food, drug, household, agricultural, or cosmetic use. An individual technically qualified to handle potentially hazardous chemicals must supervise the use of this material.

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End of Safety Data Sheet

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 AUSTRALIA  
 1-800-331-627

EMERGENCY NUMBER (SPILLS, EXPOSURES): 301/431-8585 (24 HOUR)  
 800/451-8346 (24 HOUR)  
 800/955-6288

NON-EMERGENCY INFORMATION:

Product Name: pLenti4/Ubc/V5-DESTGW Vector, 6UG Lyophilized  
 Stock Number: 350855

NOTE: If this product is a kit or is supplied with more than one material, please refer to the MSDS for each component for hazard information.

Product Use:  
 These products are for laboratory research use only and are not intended for human or animal diagnostics, therapeutic, or other clinical uses.

Synonyms:  
 Not available.

**2. COMPOSITION, INFORMATION ON INGREDIENTS**

The following list shows components of this product classified as hazardous based on physical properties and health effects:

Component	CAS No.	Percent
EDTA	60-00-4	1 - 5
TRIZMA BASE	MIXTURE	60 - 100

**3. HAZARDS IDENTIFICATION**

\*\*\*\*\* EMERGENCY OVERVIEW \*\*\*\*\*  
 Warning!  
 Irritant.  
 Harmful if swallowed.  
 Harmful if absorbed.  
 Harmful by inhalation.  
 May cause allergic skin reaction.  
 Possible reproductive system hazard based on animal data.  
 \*\*\*\*\*

Potential Health Effects:

Eye:  
 Can cause moderate irritation, tearing and reddening, but not likely to permanently injure eye tissue.

Skin:  
 Can cause moderate skin irritation, defatting, and dermatitis. Not likely to cause permanent damage.  
 May cause allergic skin reaction.  
 Upon prolonged or repeated exposure, harmful if absorbed through the skin.  
 May cause minor systemic damage.

Inhalation:  
 Can cause moderate respiratory irritation, dizziness, weakness, fatigue, nausea and headache.  
 Harmful! Can cause systemic damage (see "Target Organs").

Ingestion:  
 Mildly irritating to mouth, throat, and stomach. Can cause abdominal discomfort.  
 Harmful if swallowed. May cause systemic poisoning.

Chronic:  
 No data on cancer.  
 Contains a substance that is a possible reproductive system hazard based on animal studies at doses that could be encountered in the workplace.

**4. FIRST AID MEASURES**

Eye:  
 Immediately flush eyes with plenty of water for at least 20 minutes retracting eyelids often. Tilt the head to prevent chemical from transferring to the uncontaminated eye. Get immediate medical attention and monitor the eye daily as advised by your physician.

Skin:  
 Wash with soap and water. Remove contaminated clothing, launder

MATERIAL SAFETY DATA SHEET

PLENTI6/UBC/V5-DEBTGW VECTOR, 6UG LYOPHILIZED INVITROGEN CORPORATION MSDS ID: 350855	Page 3 of 8 Revised 12/18/03 Replaces 11/17/03 Printed 12/18/03
--	--

**4. FIRST AID MEASURES (CONT.)**

immediately, and discard contaminated leather goods. Get medical attention immediately.

**Inhalation:**

Remove to fresh air. If breathing is difficult, have a trained individual administer oxygen. If not breathing, give artificial respiration and have a trained individual administer oxygen. Get medical attention immediately.

**Ingestion:**

Severely irritating. Do not induce vomiting. Seek medical attention immediately. Drink 2 glasses of water or milk to dilute.

Note To Physician:  
Treat symptomatically.

**5. FIRE FIGHTING MEASURES**

Flashpoint Deg C:

Not available.

Upper Flammable Limit %:

Not available.

Lower Flammable Limit %:

Not available.

Autoignition Temperature Deg C:

Not available.

**Extinguishing Media:**

Can cause moderate irritation, tearing and reddening, but not likely to permanently injure eye tissue.  
Use water spray/fog for cooling.

**Firefighting Techniques/Equipment:**

Do not enter fire area without proper protection including self-contained breathing apparatus and full protective equipment. Fight fire from a safe distance and a protected location due to the potential of hazardous vapors and decomposition products.

**Hazardous Combustion Products:**

Includes carbon dioxide, carbon monoxide, dense smoke.

**6. ACCIDENTAL RELEASE MEASURES**

Accidental releases may be subject to special reporting requirements and other regulatory mandates. Refer to Section 8 for personal protection equipment recommendations.

MATERIAL SAFETY DATA SHEET

PLENTI6/UBC/V5-DESTGW VECTOR, 6UG LYOPHILIZED  
 INVITROGEN CORPORATION  
 MSDS ID: 350855

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**6. ACCIDENTAL RELEASE MEASURES (CONT.)**

Spill Cleanup:  
 Exposure to the spilled material may be irritating or harmful. Follow personal protective equipment recommendations found in Section VIII of this MSDS. Additional precautions may be necessary based on special circumstances created by the spill including; the material spilled, the quantity of the spill, the area in which the spill occurred. Also consider the expertise of employees in the area responding to the spill. Ventilate the contaminated area. Prevent the spread of any spill to minimize harm to human health and the environment if safe to do so. Wear complete and proper personal protective equipment following the recommendation of Section VIII at a minimum. Dike with suitable absorbent material like granulated clay. Gather and store in a sealed container pending a waste disposal evaluation.

**7. HANDLING AND STORAGE**

Storage of some materials is regulated by federal, state, and/or local laws.

Storage Pressure:  
 Ambient

Handling Procedures:  
 Harmful or irritating material. Avoid contacting and avoid breathing the material. Use only in a well ventilated area.  
 Keep closed or covered when not in use.

Storage Procedures:  
 Store in a cool dry ventilated location. Isolate from incompatible materials and conditions. Keep container(s) closed.  
 Suitable for most general chemical storage areas.

**8. EXPOSURE CONTROLS, PERSONAL PROTECTION**

Exposure Limits:

Component	OSHA PEL (ppm)	AGCIH TWA (ppm)
EDTA	Not established.	Not established.
TRIZMA BASE	Not established.	Not established.

Engineering Controls:  
 Local exhaust ventilation or other engineering controls are normally required when handling or using this product to avoid overexposure.

Personal Protective Equipment:

MATERIAL SAFETY DATA SHEET

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PLENTI6/UBC/V5-DESGW VECTOR, 6UG LYOPHILIZED  
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#### 8. EXPOSURE CONTROLS, PERSONAL PROTECTION (CONT.)

**Eye:**  
 An eye wash station must be available where this product is used. Wear chemically resistant safety glasses with side shields when handling this product. Wear additional eye protection such as chemical splash goggles and/or face shield when the possibility exists for eye contact with splashing or spraying liquid, or airborne material. Do not wear contact lenses. Have an eye wash station available.

**Skin:**  
 Avoid skin contact by wearing chemically resistant gloves, an apron and other protective equipment depending upon conditions of use. Inspect gloves for chemical break-through and replace at regular intervals. Clean protective equipment regularly. Wash hands and other exposed areas with mild soap and water before eating, drinking, and when leaving work. Have a safety shower available.

**Respiratory:**  
 NIOSH approved air purifying respirator with dust/mist filter. A respiratory protection program that meets OSHA's 29 CFR 1910.134 and ANSI Z88.2 requirements must be followed whenever workplace conditions warrant a respirator's use.

#### 9. PHYSICAL AND CHEMICAL PROPERTIES

**Appearance/physical state:** Liquid solution / suspension

**Odor:** No odor.  
 Not established.  
 Not established.

**Specific Gravity/Density:** Not established.  
**Octanol/water Partition Coeff:** Not established.  
**Volatiles:** Not established.  
**Evaporation Rate:** Not established.  
**Viscosity:** Not established.

#### 10. STABILITY AND REACTIVITY

**Stability:**  
 Stable under normal conditions.

**Conditions to Avoid:** Strong oxidizing agents. High temperatures. Strong alkalis. Copper alloys. Aluminum alloys.

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PLENTIC/UBC/V5-DESGW VECTOR, 6UG LYOPHILIZED  
 INVITROGEN CORPORATION  
 MSDS ID: 350855

**15. REGULATORY INFORMATION (CONT.)**

PRODUCT CLASSIFICATION: XI

Component EINECS  
 EDTA Number 200-449-4  
 TRIZMA BASE Not established.

**16. OTHER INFORMATION**

HMS Rating 0-4:  
 FIRE: Not determined.  
 HEALTH: Not determined.  
 REACTIVITY: Not determined.

- Abbreviations  
 N/A - Data is not applicable or not available  
 SARA - Superfund and Reauthorization Act  
 HMIS - Hazard Material Information System  
 WHMIS - Workplace Hazard Materials Information System  
 NTP - National Toxicology Program  
 OSHA - Occupational Health and Safety Administration  
 IARC - International Agency for Research on Cancer  
 PROP 65 - California Safe Drinking Water and Toxic Enforcement Act of 1986  
 EINECS - European Inventory of Existing Commercial Chemical Substances

The above information was acquired by diligent search and/or investigation and the recommendations are based on prudent application of professional judgment. The information shall not be taken as being all inclusive and is to be used only as a guide. All materials and mixtures may present unknown hazards and should be used with caution. Since Invitrogen Corporation cannot control the actual methods, volumes, or conditions of use, the Company shall not be held liable for any damages or losses resulting from the handling or from contact with the product as described herein. THE INFORMATION IN THIS MSDS DOES NOT CONSTITUTE A WARRANTY, EXPRESS OR IMPLIED, INCLUDING ANY IMPLIED WARRANTY OF MERCHANTABILITY OR FITNESS FOR ANY PARTICULAR PURPOSE.

**Subject:** Biological Agents Registry Form: O'Gorman  
**From:** Jennifer Stanley <jstanle2@uwo.ca>  
**Date:** Thu, 19 Aug 2010 15:55:47 -0400  
**To:** David O'Gorman <dogorman@uwo.ca>

Hello Dr. O'Gorman -

Thank you for your recent submission.

I was looking for information on the POSTN gene that you transduce using lentivirus....can you verify that one (or both) of these descriptions are accurate? If so, I will include it with your form.

<http://www.genecards.org/cgi-bin/carddisp.pl?gene=POSTN>  
<http://www.wikigenes.org/e/gene/e/50706.html>

Regards  
Jennifer

Show Advanced Search

GeneAlaCart

GeneDecks



keyword(s)

Search

**POSTN Gene**  
 protein-coding **GIFtS: 54**  
 GC13M038136

**periostin, osteoblast specific factor**

Symbol approved by the HUGO Gene Nomenclature Committee (HGNC) database

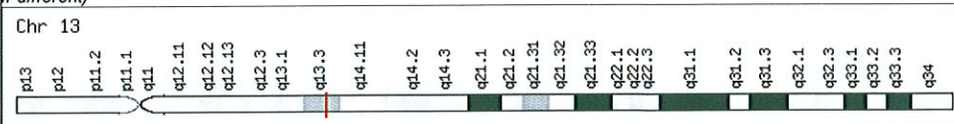
**SIGMA** Pathways  
 Antibodies  
 Proteins / shRNA / esiRNA  
 Small Molecules / siRNA / miRNA

**M** Antibodies/cDNA/RNAi  
 Proteins & Enzymes  
 Assays & Kits/Pathways

**Services**  
 Jump to Section...

**AB** applied Expression  
 biosystems RNAi  
 SNPs / Genotyping / Pathways

**ORIGENE** Proteins  
 Antibodies  
 Assays/Genes/shRNA/Primers

<p><b>Aliases &amp; Descriptions for POSTN gene</b></p> <p>(According to <sup>1</sup>HGNC, <sup>2</sup>Entrez Gene, <sup>3</sup>UniProtKB/Swiss-Prot, <sup>4</sup>UniProtKB/TrEMBL, <sup>5</sup>OMIM, <sup>6</sup>GeneLoc, and/or <sup>7</sup>Ensembl, <sup>8</sup>miRBase)</p> <p><a href="#">About This Section</a></p> <p>Jump to Section...  <a href="#">User Feedback</a></p>	<p><b>Aliases &amp; Descriptions</b></p> <p>periostin, osteoblast specific factor<sup>1 2</sup>    periostin isoform thy<sup>4 2</sup>          PN1<sup>2 3 5</sup>    osteoblast specific factor 2 (fascin I-like)<sup>2</sup>          OSF-2<sup>1 2 3</sup>    RP11-412K4.1<sup>2</sup>          OSF2<sup>2 3 5</sup>    periostin isoform thy<sup>6 2</sup>          periostin<sup>1 2</sup>    PDLPOSTN<sup>2</sup>          Osteoblast-specific factor<sup>2 3</sup>    MGC119510<sup>2</sup>          periostin isoform thy<sup>2 2</sup>    periostin isoform thy<sup>8 2</sup>          MGC119514<sup>2</sup>    periodontal ligament-specific periostin<sup>2</sup></p> <p>External Ids:    HGNC: 16953<sup>1</sup>    Entrez Gene: 10631<sup>2</sup>    Ensembl: ENSG00000133110<sup>7</sup>    UniProtKB: Q15063<sup>3</sup></p> <p>Search outside databases for aliases for POSTN gene</p> <p>Previous GC identifiers: GC13M035934 GC13M037034</p>
<p><b>Summaries for POSTN gene</b></p> <p>(According to Entrez Gene, Tocris Bioscience, Wikipedia's Gene Wiki, UniProtKB/Swiss-Prot, and/or UniProtKB/TrEMBL)</p> <p><a href="#">About This Section</a></p> <p>Jump to Section...  <a href="#">User Feedback</a></p>	<p>UniProtKB/Swiss-Prot: <a href="#">POSTN_HUMAN_Q15063</a></p> <p><b>Function:</b> Binds to heparin. Induces cell attachment and spreading and plays a role in cell adhesion. May play a role in extracellular matrix mineralization</p> <p><b>Gene Wiki entry for POSTN</b></p>
<p><b>Genomic Views for POSTN gene</b></p> <p>(According to GeneLoc and/or HGNC, and/or Entrez Gene (NCBI build 37), and/or miRBase, Genomic Views according to UCSC and Ensembl (release 56), Regulatory elements and Epigenetics data according to SABiosciences)</p> <p><a href="#">About This Section</a></p> <p>Jump to Section...  <a href="#">User Feedback</a></p>	<p><b>Regulatory elements:</b></p> <p> SABiosciences <a href="#">Regulatory transcription factor binding sites</a> in the POSTN gene upstream (promoter) region <b>improved</b>: STAT1alpha STAT1beta STAT1 NF-kappaB1 NF-kappaB STAT5A CUTL1 FOXL1 p300 E4BP4</p> <p><b>Epigenetics:</b></p> <p> Search SABiosciences <a href="#">Methyl-Profiler DNA Methylation qPCR Primer Assays for POSTN</a> <b>NEW</b></p> <p><b>Genomic Location:</b></p> <p>Genomic View: <a href="#">UCSC Golden Path with GeneCards custom track</a></p> <p>Entrez Gene cytogenetic band: <a href="#">13q13.3</a>    Ensembl cytogenetic band: <a href="#">13q13.3</a>    HGNC cytogenetic band: <a href="#">13q13.3</a></p> <p>POSTN Gene in genomic location: bands according to Ensembl, locations according to <a href="#">GeneLoc</a> (and/or Entrez Gene and/or Ensembl if different)</p>  <p>GeneLoc gene densities for chromosome 13    <a href="#">GeneLoc Exon Structure</a></p> <p>GeneLoc location for GC13M038136: <a href="#">view genomic region</a> (about GC identifiers)</p> <p>Start:    38,136,719 bp from pter          End:    38,172,981 bp from pter          Size:    36,263 bases          Orientation: minus strand</p>

ProtoNet, UniProtKB, and/or BLOCKS, Sets of similar genes according to GeneDecks)  
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IPR011489 EMI\_domain  
 IPR016666 TGFb-ind\_bIGH3/osteoblast\_fac2  
 IPR000782 FAS1\_domain

[Graphical View of Domain Structure for InterPro Entry Q15063](#)

ProtoNet protein and cluster: [Q15063](#)

2 Blocks protein families:  
 IPB000782 Beta-Ig-H3/Fasciclin domain  
 IPB011489 EMI

UniProtKB/Swiss-Prot: [POSTN\\_HUMAN\\_Q15063](#)  
 Similarity: Contains 1 EMI domain  
 Similarity: Contains 4 FAS1 domains

Gene Function for POSTN gene  
 (According to MGI May 08 2010, UniProtKB, IUBMB, and/or Genatlas, shRNA from OriGene, Sigma-Aldrich, RNAi from Millipore, Abnova, siRNAs from Applied Biosystems, Sigma-Aldrich, Clones from Millipore, Sigma-Aldrich, OriGene, Sino Biological, Ontologies according to Gene Ontology Consortium 01 May 2010 via Entrez Gene.)  
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Inhib. RNA:  [Browse for Gene Knock-down Tools from Millipore](#)

 [Browse Abnova for Chimera RNAi Products](#)

 [Origene 29mer shRNA kits in GFP-retroviral vector \(see all 4\): POSTN](#)  
[Origene shRNA RFP \(see all 4\): POSTN](#)  
[Origene basic RS shRNA \(see all 4\): POSTN](#)

 [Applied Biosystems Silencer® siRNAs for POSTN](#)  
[Sigma-Aldrich siRNA for POSTN](#)  
[Sigma-Aldrich shRNA Panels and shRNA for POSTN](#)  
[Explore Sigma-Aldrich super-pooled esiRNAs](#)

Clones:  [Browse Clones for the Expression of Recombinant Proteins Available from Millipore](#)

 [Browse iPSC Reprogramming Factors at Sigma-Aldrich](#)

 [Origene GFP tagged cDNA clones in CMV expression vector \(see all 4\): POSTN](#)  
[Origene Myc/DDK tagged cDNA clones in CMV expression vector \(see all 4\): POSTN](#)  
[Origene untagged cDNA clones in CMV expression vector \(see all 4\): POSTN](#)

 [Browse Sino Biological Human cDNA Clones](#) 

UniProtKB/Swiss-Prot: [POSTN\\_HUMAN\\_Q15063](#)

Function: Binds to heparin. Induces cell attachment and spreading and plays a role in cell adhesion. May play a role in extracellular matrix mineralization

2 Gene Ontology (GO) molecular function terms (GO ID links to tree view):

GO ID	Qualified GO term	Evidence	PubMed IDs
<a href="#">GO:0005515</a>	protein binding	IEA	--
<a href="#">GO:0008201</a>	heparin binding	ISS	--

[About this table](#)

 [POSTN for ontologies](#) [About GeneDecksing](#)

Animal Models: 14 MGI mutant phenotypes (inferred from 3 alleles ) ([MGI details for Postn](#)):

[cardiovascular system](#) [cellular](#) [craniofacial](#) [endocrine/exocrine gland](#) [growth/size](#)  
[homeostasis/metabolism](#) [immune system](#) [lethality-postnatal](#) [life span-post-weaning/aging](#) [limbs/digits/tail](#)  
[muscle](#) [reproductive system](#) [skeleton](#) [tumorigenesis](#)

 [POSTN for phenotypes](#) [About GeneDecksing](#)

Pathways & Interactions for POSTN gene  
 (Pathways according to Millipore, Cell Signaling Technology, Sigma-Aldrich, Applied Biosystems GeneAssist, KEGG and/or UniProtKB, (map by GeneGo), Sets of similar genes according to GeneDecks, Proteins Network according to SABiosciences, Interactions according to <sup>1</sup>UniProtKB, <sup>2</sup>MINT, and/or <sup>3</sup>STRING, with links to IntAct and Ensembl, Ontologies

 SABiosciences Gene Network Central™ Interacting Genes and Proteins Network for [POSTN](#)

4 Gene Ontology (GO) biological process terms (GO ID links to tree view):

GO ID	Qualified GO term	Evidence	PubMed IDs
<a href="#">GO:0001501</a>	skeletal system development	TAS	<a href="#">8363580</a>
<a href="#">GO:0007155</a>	cell adhesion	IDA	<a href="#">12235007</a>
<a href="#">GO:0009888</a>	tissue development	IEA	--
<a href="#">GO:0030198</a>	extracellular matrix organization	IEA	--

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 [POSTN for ontologies](#) [About GeneDecksing](#)

Periostin, osteoblast specific factor  
 Hs.136348 [show with all ESTs], Hs.721018 [show with all ESTs]  
 Unigene Representative Sequences: [NM\\_006475](#), [AK021444](#)

[GeneLoc Exon Structure](#)

**5/9 Alternative Splicing Database (ASD) splice patterns (SP) for POSTN (see all 9)**

ExUns: 1 ^ 2 ^ 3 ^ 4 ^ 5 ^ 6 ^ 7 ^ 8 ^ 9 ^ 10 ^ 11 ^ 12 ^ 13 ^ 14 ^ 15 ^ 16a · 16b ^ 17 ^ 18a · 18b · 18c ^ 19a · 19b · 19c ^ 20a · 20b ^

SP1: - - - - -  
 SP2: - - - - -  
 SP3: - - - - -  
 SP4: - - - - -  
 SP5: - - - - -

ExUns: 21a · 21b ^ 22 ^ 23a · 23b · 23c · 23d

SP1: -  
 SP2: -  
 SP3: -  
 SP4: - -  
 SP5: - -

[About this scheme](#)

**ECgene alternative splicing isoforms for POSTN**

**8 Ensembl transcripts including schematic representations:**

[ENST00000474646](#) [ENST00000473823](#) [ENST00000478947](#) [ENST00000497145](#) [ENST00000379749](#) [ENST00000379743](#)  
[ENST00000379742](#) [ENST00000379747](#)

**Expression for POSTN gene**  
 (Experimental results according to <sup>1</sup>GeneNote and GNF BioGPS, probe sets-to-genes annotations according to <sup>2</sup>GeneAnnot, <sup>3</sup>GeneTide, Sets of similar genes according to GeneDecks, Electronic Northern calculations according to data from UniGene (Build 223 Homo sapiens), SAGE tags according to CGAP, plus additional links to SOURCE, and/or GNF BioGPS, and/or EXPOLDB, and/or UniProtKB, Expression Assays from Applied Biosystems, Primers from OriGene and/or SABiosciences )  
[About This Section](#)

**POSTN expression in normal and diseased human tissues**

**AB** Applied Biosystems TaqMan® Gene Expression Assays for POSTN

[GeneNote](#) / [GeneAnnot](#) / [GeneTide](#)

**7 probe-sets matching POSTN gene**

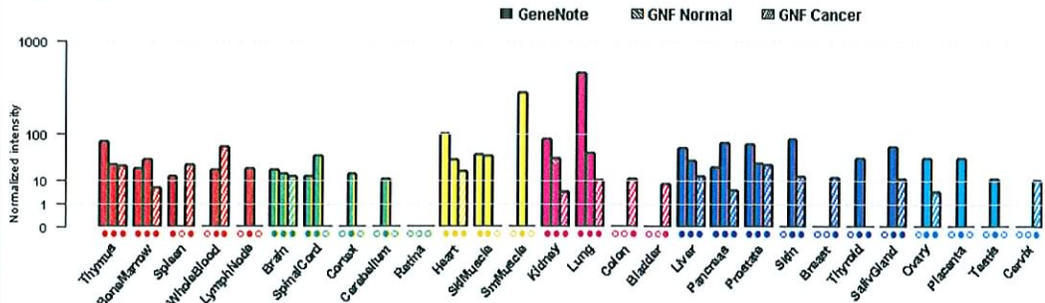
Affymetrix probe-set	Array	GeneAnnot data			GeneNote data		GeneTide data				
		# genes	Sensitivity	Specificity	Correlation	Length	Gb_Accession	Consensus	Uniqueness	Score	Rank
1451_s_at <sup>2,3</sup>	U95-A	1	1.00	1.00	1.00	1.00	D13666	1.00	1.00	1.00	1
210809_s_at <sup>2,3</sup>	U133-A	1	1.00	1.00	--	--	D13665	0.80	1.00	0.91	1
214981_at <sup>2,3</sup>	U133-A	1	0.55	1.00	--	--	AW137148	0.40	1.00	0.76	1
1555778_a_at <sup>2</sup>	U133Plus2	1	1.00	1.00	--	--	--	--	--	--	--
210809_s_at <sup>2</sup>	U133Plus2	1	1.00	1.00	--	--	--	--	--	--	--
214981_at <sup>2</sup>	U133Plus2	1	0.55	1.00	--	--	--	--	--	--	--
1555777_at <sup>2</sup>	U133Plus2	1	0.36	1.00	--	--	--	--	--	--	--

[About this table](#)

**GeneDecks** POSTN for expression [About GeneDecksing](#)

**Data from Genenote (Publications) and GNF BioGPS**

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**Genomic Variants for POSTN gene**

10/274 NCBI SNPs in POSTN are shown (see all 274 )  
(Click **AB** for Applied Biosystems TaqMan® Genotyping Assay) (see all 116)

(SNPs according to the <sup>1</sup>NCBI SNP Database, <sup>2</sup>Ensembl, <sup>3</sup>PupaSUITE, and UniProtKB, Linkage Disequilibrium by HapMap, Structural Variations (CNVs/InDels /Inversions) from the Database of Genomic Variants, Genotyping Reagents from Applied Biosystems)  
[About This Section](#)

AB	Genomic Data				Transcription Related Data				Allele Frequencies				
	SNP ID	Valid	Chr 13 pos	Sequence	Recs	AA Chg	Type	More	Recs	Allele freq	Pop	Total sample	More
Sort	-	1st	-	--	--	--	2nd	--	--	-	-	-	--
<b>AB</b>	rs9547951 <sup>1,2</sup>	C,F,A,H	18936145(+)	cctctgC/Aqctta	4	--	ng3 <sup>1</sup>		9		NS EA NA WA	532	
<b>AB</b>	rs3812842 <sup>1,2</sup>	C,F,A,H	18974259(-)	ATTACG/ACAAGT	4	--	ng5 <sup>1</sup>		13		NA NS EA WA	686	
<b>AB</b>	rs9532087 <sup>1,2</sup>	C,F,A,H	18974391(+)	CTTAAC/TATTGC	4	--	ng5 <sup>1</sup>		14		NA EA WA	718	
<b>AB</b>	rs9547952 <sup>1,2</sup>	C,F,A,H	38138689(+)	CCTCAC/TGGGTG	8	M V	mis <sup>1</sup> ref <sup>1</sup> ese <sup>3</sup>		10		NS EA NA	998	
<b>AB</b>	rs1028728 <sup>1,2</sup>	C,F,O	18973294(+)	TTTCAA/TTGTTA	4	--	ng5 <sup>1</sup>		91		NS EA PA EU CA WA NA MN	5136	
<b>AB</b>	rs3829365 <sup>1,2</sup>	C,F,A	38172896(-)	GTTCTC/GTTCGG	4	--	ut5 <sup>1</sup>		15		MN EA NS NA WA	1410	
<b>AB</b>	rs9576302 <sup>1,2</sup>	C,F,A	18936042(+)	cggccG/ATCTAA	4	--	ng3 <sup>1</sup>		11		NS EA NA	848	
<b>AB</b>	rs6750 <sup>1,2</sup>	C,F,A	38136832(-)	AAATTG/CAGTAA	4	--	ut3 <sup>1</sup> ese <sup>3</sup>		10		MN NA NS EA	848	
<b>AB</b>	rs9594223 <sup>1,2</sup>	C,F	38158945(+)	TTACTA/C/G/T TTATA	16	K T R I	mis <sup>1</sup> ref <sup>1</sup>		9		NS EA NA	908	
<b>AB</b>	rs769131 <sup>1,2</sup>	F,H	18973113(+)	CACCTT/AATGIG	4	--	ng5 <sup>1</sup>		4		NS EA	420	

[About this table](#)

HapMap Linkage Disequilibrium images for POSTN (up to first 250kb)

Structural Variations (Copy Number Variations, Insertions/Deletions, Inversions)   
Database of Genomic Variants (DGV): 2 variations for POSTN  
2 CNVs: 9709 47992

**Disorders & Mutations for POSTN gene**



POSTN for disorders

[About GeneDecksing](#)

(in which this Gene is Involved, According to OMIM, UniProtKB, Novoseek, PharmGKB, Genatlas, GeneTests, Blood group antigen gene mutations by BGMUT, LSDB, HGMD, GAD, HuGE Navigator, BCGD, and/or TGDB.)  
[About This Section](#)

OMIM: 608777

10/127 Novoseek disease relationships for POSTN gene (see all 127 )

Disease	Score	Articles	PubMed IDs for Articles with Shared Sentences (# sentences)
pterygia	10.10	6	19661231 (5), 17652725 (1)
carcinoma papillary thyroid	9.34	10	18434370 (8), 19321256 (2)
dupuytren's disease	7.57	5	19619531 (4), 19121738 (1)
bladder cancer	7.44	15	15880581 (7), 18097555 (6), 19578758 (2)
cholangiocarcinoma	6.10	7	20096135 (7)
nsclc	5.98	13	11550156 (6), 18949745 (4), 11509119 (3)
fibrous dysplasia	5.68	5	18799196 (5)
pancreatic cancer	5.22	12	18381746 (6), 17043657 (6)
keloids	4.64	5	17649947 (3), 19799038 (1), 18560459 (1)
adpkd	4.29	5	18753297 (5)

[About this table](#)

Human Genome Epidemiology Navigator: POSTN (4 documents)

**Medical News for POSTN gene**

(Possibly Related Articles in Doctor's Guide)  
[About This Section](#)

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**Publications for POSTN gene**

(in PubMed. Associations of this gene to articles via <sup>1</sup>Novoseek, <sup>2</sup>HGNC, <sup>3</sup>Entrez Gene, <sup>4</sup>UniProtKB/Swiss-Prot,

10/132 PubMed articles for POSTN gene (see all 132 ):

- Periostin secreted by epithelial ovarian carcinoma is a ligand for alpha(V)beta(3) and alpha(V)beta(5) integrins and promotes cell motility. (PubMed id 12235007)<sup>1, 2, 3, 4</sup> Gillan L....Chang D.D. (2002)
- Osteoblast-specific factor 2: cloning of a putative bone adhesion protein with homology with the insect protein fasciclin I. (PubMed id 8363580)<sup>2, 3, 4</sup> Takeshita S.... Amann E. (1993)
- Serum level of the periostin, a homologue of an insect cell adhesion molecule, as a prognostic marker in non-small cell lung carcinomas. (PubMed id 11550156)<sup>1, 3, 4</sup> Sasaki H.... Chen L.B. (2001)
- Periostin, a member of a novel family of vitamin K-dependent proteins, is expressed by mesenchymal stromal cells. (PubMed id

**Services for POSTN gene**

(Reagents available from Applied Biosystems, Antibodies and assays by Cell Signaling Technology, Novus Biologicals, Epitomics, Sigma-Aldrich, R&D Systems, SABiosciences, Millipore, Abnova, Clones available from OriGene, Sigma-Aldrich, Sino Biological, Drugs and/or compounds by Sigma-Aldrich, Enzo Life Sciences, and/or Tocris Bioscience) [About This Section](#)

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**Products for POSTN:**

- › [TaqMan® Gene Expression Assays](#)
- › [TaqMan® Genotyping Assays](#)
- › [Free SNP selection tool](#)



- › [Millipore Custom Antibody & Bulk Services](#)
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