

Modification Form for Permit BIO-UWO-0023

Permit Holder: Martin Sandig

Approved Personnel

(Please stroke out any personnel to be removed)

~~Ying Xia~~

Additional Personnel

(Please list additional personnel here)

Please stroke out any approved Biohazards to be removed below

Write additional Biohazards for approval below. Give the full name - do not abbreviate.

Approved Microorganisms

~~E. coli dHS-alpha~~

Approved Primary and Established Cells

~~Human (primary): coronary arteries, umbilical veins, Lung microvasculature. Human (established): THP-1, PG3, DU145, LnGAP, HEK-293~~

mouse or rodent tissues (human later)

Approved Use of Human Source Material

~~Human Blood (whole) or other body fluid: Healthy donors / LHSC-UH~~

Approved Genetic Modifications (Plasmids/Vectors)

~~Plasmid: pcDNA3.1, pIRES2-EGFP, Vector: pLNGX2~~

Approved Use of Animals

Approved Biological Toxin(s)

* PLEASE ATTACH A MATERIAL SAFETY DATA SHEET OR EQUIVALENT FOR NEW BIOHAZARDS.
** PLEASE ATTACH A BRIEF DESCRIPTION OF THE WORK THAT EXPLAINS THE BIOHAZARDS USED AND HOW THEY WILL BE STORED, USED AND DISPOSED OF..

As the principal investigator, I have ensured that all of the personnel named on the form have been trained. 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 of Permit Holder: 

Current Classification: 2 Containment Level for Added Biohazards: _____

Date of Last Biohazardous Agents Registry Form: Apr 27, 2009

Date of Last Modification (if applicable): _____

BioSafety Officer(s): _____

Chair, Biohazards Subcommittee: _____ Date: _____

Research do be performed in the labs MSB475 and MSB479

While there have been significant advances in developing 3D teaching tools for anatomy, the teaching of histology on the basis of 3D reconstructions has not received equal attention. The work to be done in the lab involves the processing of perfusion-fixed animal tissues for serial histological sectioning and staining. Although human tissues will be used in the future the initial project will use tissues obtained from small mammals. Mouse or rat tissues will be obtained from collaborators that use these animals for other research purposes. Tissues will be fixed by perfusion in a solution containing 2% formaldehyde and 2.5% glutaraldehyde and processed for embedding in Epon 812 (lab M475). This will involve the use of serial dilutions of ethanol and propylene oxide. Since tissue integrity is essential for 3D reconstruction from serial sections, complete semi-thin (1 μm thick) serial sections will be obtained with a Histo Jumbo diamond knife and a Reichert/Jung Ultracut E microtome (lab M479). Sections will be transferred to microscope slides and stained with Methylene blue and azure II, a polychromatic stain that has been shown to give outstanding differential staining of various cellular components in plastic embedded tissues (lab M475). Stained sections will be photographed using a 40X lens on a Light Microscope equipped with a digital color camera. Cellular structures of interest will be marked and outlined and 3D volume rendering will be performed using the digital animation software in Amira.

Alcohols and organic solvents, as well as fixatives such as glutaraldehyde and formaldehyde, will be stored and disposed of according to WHMIS and Biosafety Guidelines.

The lab facility in DSB00060 has been vacated by Drs. Belliveau and Sandig.

**THE UNIVERSITY OF WESTERN ONTARIO
 BIOHAZARDOUS AGENTS REGISTRY FORM
 Approved Biohazards Subcommittee: March 27, 2009
 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 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 also 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. 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>Martin Sandig</u>
SIGNATURE	
DEPARTMENT	<u>Anatomy and Cell Biology</u>
ADDRESS	<u>DSB 00075</u>
PHONE NUMBER	<u>ext. 86815</u>
EMERGENCY PHONE NUMBER(S)	<u>519-434-4617</u>
EMAIL	<u>martin.sandig@schulich.uwo.ca</u>

Location of experimental work to be carried out: Building(s) DSB Room(s) 00060

*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: HSFO: Integrin signaling in monocyte transendothelial migration
 GRANT TITLE(S): HSFO: Vascular smooth muscle cell phenotype switching and elastin
synthesis in 3D tissue engineered coronary artery substitutes
NSERC: Tumor cell transendothelial migration

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

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

<u>Ying Xia, Ph.D.</u>	_____
_____	_____
_____	_____
_____	_____

1.0 Microorganisms

1.1 Does your work involve the use of microorganisms or biological agents of plant or animal origin (including but not limited to viruses, prions, parasites, bacteria)? YES NO
 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)*	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
<i>E. coli</i> dHS alpha	<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	40L	lab cloning strain	<input checked="" type="radio"/> 1 <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"/> 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"/> 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"/> 3

98
pl-e-mail attached

*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 in the table below

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	Coronary Arteries Umbilical Veins Lung Microvasculature	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 the table below.

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	THP-1, PC3, DU145, LnCAP	ATCC
Rodent	<input type="radio"/> Yes <input type="radio"/> No	HEK-293	AT confirmed by email - see attached
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 types(s) indicate PHAC or CFIA containment level required 1 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 Known to Be 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	Healthy donors/ LHSC UH	<input type="radio"/> Yes <input checked="" type="radio"/> No		<input type="radio"/> 1 <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"/> 1 <input type="radio"/> 2 <input type="radio"/> 3
Human Organs or Tissues (unpreserved)		<input type="radio"/> Yes <input type="radio"/> No		<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 3
Human Organs or Tissues (preserved)		<input type="radio"/> Yes <input type="radio"/> No		<input type="radio"/> 1 <input type="radio"/> 2 <input type="radio"/> 3

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
E-coli	pcDNA3.1, pIRES2-EGFP	Invitrogen Clontech	RhoA, Rac1 cDNAs excised from pcDNA3.1 and cloned into pIRES2-EGFP	Construction of viral vectors

* 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 done? YES, complete table below NO

Virus Used for Transduction *	Vector(s) *	Source of Vector	Gene Transfected	Describe the change that results
Phoenix helper-free packaging cells	pLNCX2	Clontech	RhoAG14V, RhoAT19N, Rac1G12V, Rac1T17N, Cdc42G12V, Cdc42T17N	defects in microfilament assembly dynamics and cell motility

* Please attach a Material Safety Data Sheet or equivalent.

4.4 Will genetic sequences from the following be involved?

- ◆ 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 using the viral vector in 4.0? YES NO
 If no, please proceed to Section 6.0 If YES attach a full description of the make-up of the virus.

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

5.3 How will the virus 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 be used in live animals YES, specify: _____ NO

10.0 Plants Requiring CFIA Permits

10.1 Do you use plants that require a permit from the CFIA? 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

10.9 Please 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 _____
If no, please proceed to Section 10.0 NO

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 Prabh Jand

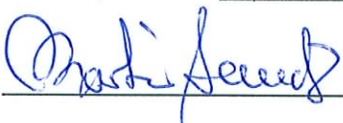
13.0 Containment Levels

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

13.2 Has the facility been certified by OHS for this level of containment?
 YES, permit # if on-campus BIO-UWO-0023
 NO, please certify
 NOT REQUIRED for Level 1 containment

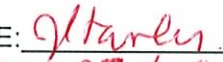
14.0 Procedures to be Followed

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  Date: April 3, 2009

15.0 Approvals

UWO Biohazard Subcommittee: SIGNATURE: 
Date: 27 April 2009

Safety Officer for Institution where experiments will take place: SIGNATURE: 
Date: April 29/09

Safety Officer for University of Western Ontario (if different from above): SIGNATURE: _____
Date: _____

Approval Number: Bio-UWO-0023 Expiry Date (3 years from Approval): Apr 26, 2012

Special Conditions of Approval:

Subject: Re: Biohazardous Agents Registry Form: Sandig
From: Martin Sandig <Martin.Sandig@schulich.uwo.ca>
Date: Mon, 27 Apr 2009 14:14:48 -0400
To: Jennifer Stanley <jstanle2@uwo.ca>

Hi Jennifer:

You are correct. We use the E-coli for small volume cloning and not for large scale protein expression.

Cheers,
Martin

Martin Sandig, PhD
Associate Professor
Associate Chair, Undergraduate Affairs
Department of Anatomy and Cell Biology
Schulich School of Medicine and Dentistry
University of Western Ontario
Dental Sciences Building, Room 00075
Phone: 519 661 2111 86815
Fax: 519 850 25662
<http://www.uwo.ca/anatomy/department/sandigm/msandig.html>

>>> Jennifer Stanley <jstanle2@uwo.ca> 4/27/2009 2:08 PM >>>

Thanks for the clarification, Dr. Sandig,
I will note this on your form - I assume that you culture less than 10 litres of the E. coli (if this is not the case, let me know).

Thanks again,
Jennifer

Martin Sandig wrote:

- > Hi Jennifer:
- > 1. The 293 cells are HEK293T cells being part of the commercially available Phoenix retroviral expression system.
- > 2. We use the dh5alpha E-coli strain.
- > Cheers,
- > Martin

5. A complete informative summary of the proposal on this page.
Présentez sur cette page un résumé complet du projet de recherche.

Integrin signaling in monocyte transendothelial migration

Atherosclerosis involves the extravasation of monocytes into the intima of arterial vessels, a process known as transendothelial migration or diapedesis. Cellular adhesion and motility during diapedesis is regulated largely by the monocyte integrins VLA-4 and LFA-1, binding to endothelial VCAM-1 and ICAM-1, respectively. Under the regulation of the chemokines MCP-1 and the intracellular signaling molecule PI3K these integrins activate signaling complexes that link integrin-mediated adhesion to the regulation of the actin cytoskeleton and cell motility. Specific actin assemblies are nucleated by the Arp2/3 protein complex. During leukocyte chemotaxis filopodia are formed through Cdc42/WASP activation of Arp2/3, while lamellipodia form through activation of Arp2/3 by Rac1/WAVE2. Our recent work demonstrated in monocytes that PI3K in the absence of MCP-1 induced the activation of VLA-4, enhanced adhesion and spreading on VCAM-1, and promoted diapedesis without affecting ICAM-1 dependent interactions. In the presence of MCP-1, in contrast, LFA-1/ICAM-1 interactions were stimulated in a PI3K-dependent manner, while VLA-4/VCAM-1 interactions were rendered PI3K independent. These experiments strongly suggested that activation of LFA-1 and VLA-4 is differentially regulated, involving both MCP-1 and PI3K. It is not known, however, how following ligand binding specific signaling pathways link LFA-1 and VLA-4 to specific changes in cytoskeletal assemblies resulting in coordinated motility during diapedesis. We hypothesize that during adhesion, motility and diapedesis VLA-4/VCAM-1 interactions largely regulate Arp2/3 activation through Cdc42/WASP initiating filopodia formation, while LFA-1/ICAM-1 interactions largely mediate lamellipodia formation through Rac1/WAVE2. To address this hypothesis we will use substrates coated with VCAM-1 and ICAM-1 protein, a well established culture model of diapedesis, and a hypercholesterolemic rat model of atherogenesis to dissect spatio-temporal changes in the association of monocyte integrins with key cytoskeletal regulators during monocyte adhesion, motility and diapedesis. We will focus on the following specific aims:

Aim 1: Characterize signaling complexes associated with VLA-4 and LFA-1 in response to MCP-1 in unbound monocytes and following binding to their ligands VCAM-1 and ICAM-1, respectively. We will expose untreated or MCP-1 treated monocytes to a) soluble VCAM-1 and ICAM-1 protein, b) VCAM-1 and ICAM-1 coated surfaces, and c) endothelial monolayers to examine at various times by immunoprecipitation and western blot analysis the composition and state of phosphorylation of signaling complexes associated with VLA-4 and LFA-1. We will focus on WASP, WAVE2, Arp3, RhoA, Rac1, Cdc42 and PI3K using commercially available antibodies. Experiments will be done in the absence and in the presence of chemical crosslinking to capture transient molecular complexes. The role of PI3K will be explored by pretreating monocytes with the PI3K inhibitor LY294002 or its inactive analogue LY303511. Epitope specific antibodies and flow cytometry will be employed to determine the relative activation state of VLA-4 and LFA-1

Aim 2: Determine whether VLA-4 and LFA-1 specifically colocalize with Cdc42/WASP or Rac1/WAVE2 in filopodia or lamellipodia, respectively, during adhesion to VCAM-1 or ICAM-1, and during diapedesis through endothelium in culture and in situ. Using multi-label immunocytochemistry and laser scanning confocal microscopy we will colocalize PI3K, Cdc42/WASP/Arp3 and Rac1/WAVE2/Arp3, as well as molecular components identified under Aim 1 with LFA-1 and VLA-4 in monocytes adhering to protein carpets, during diapedesis through endothelial monolayers in culture, and during atherogenesis in a hypercholesterolemic rat model in situ. Changes in adhesion, morphology, motility, and diapedesis will be quantified by imaging following F-actin staining and by time-lapse video microscopy.

Aim 3: Determine the effects of WASP and WAVE2 inhibition on monocyte interactions with VCAM-1, ICAM-1 and endothelial monolayers. Pharmacological inhibition of WASP using the WASP inhibitor Wiskostatin and siRNA inhibition of WASP and WAVE2 expression will be used to determine their respective roles during adhesion and spreading on VCAM-1 and ICAM-1 as well as during diapedesis through endothelial monolayers. Following F-actin labeling monocyte filopodia and lamellipodia formation will be scored as well as spatio-temporal changes in morphology during diapedesis. This aim will explore to what extent

The proposed experiments will allow us to gain a better understanding of the molecular mechanisms that regulate monocyte motility during extravasation and may help us to design novel therapeutic strategies to treat pathogenic inflammatory responses such as

5. A complete informative summary of the proposal on this page.
Présentez sur cette page un résumé complet du projet de recherche.

Vascular smooth muscle cell phenotype switching and elastin synthesis in 3D tissue engineered coronary artery substitutes.

Background: Diseases of the cardiovascular system are the major causes of mortality in Canada. Current surgical interventions to replace diseased blood vessels using prosthetic materials often fail due to host immune rejection. Other alternatives such as xenografts face host rejection and inter-species disease transmission. Tissue engineering has emerged as a promising technology in the design of responsive living blood vessels with properties similar to that of the native tissue. In tissue engineering of blood vessels, biodegradable porous 3D scaffolds are seeded with vascular cells and cultured in a bioreactor to remodel with extracellular matrix proteins. Vascular smooth muscle cells (VSMCs) and elastin constitute the main components of the tunica media of arterial vessels. In arteries, elastin confers elasticity preventing dynamic tissue creep by stretching under load and recoiling to their original configurations after the load is released. In addition to the mechanical responsiveness, elastin is a potent autocrine regulator of VSMC activity for preventing fibrocellular pathology. Elastin induces VSMC actin stress fiber organization, inhibits proliferation and regulates migration. In vivo elastin knockout studies and clinical observations have revealed an essential regulatory function since, in the absence of extracellular elastin, VSMC proliferation stenoses arteries. Thus to ensure appropriate mechanical function of the vessel and to prevent vessel stenosis, successful engineered vascular tissues must incorporate an elastic component and as such, represents a critical design goal. In spite of its critical role in vessel integrity, elastin is conspicuously absent from previous tissue-engineered vascular substitutes. Although progress has been made towards understanding the underlying principles of in vivo elastin biosynthesis and incorporation into fibers, many of the conditions and mechanisms required to form viable, elastin-containing engineered vessels are still elusive. Our main goal is to induce VSMC phenotype shifts to engineer coronary artery substitutes. We hypothesize that under specific biochemical and biomechanical signals, VSMCs seeded into porous 3D biodegradable polyurethane scaffolds will modulate their phenotype. In a synthetic phenotype, they will proliferate and produce elastin while in a subsequent contractile phenotype, they will express contractile markers similar to that observed in vivo. In order to test the above hypothesis we propose to focus on the following specific objectives: (i) Establish the role of fibronectin immobilization on 3D porous biodegradable polyurethane scaffolds on attachment, proliferation and synthetic phenotypic expression of human coronary artery smooth muscle cells (HCASMCs).

(ii) Elucidate the effects of TGF-beta and retinoic acid on elastin biosynthesis by HCASMCs grown on 3D biodegradable polyurethane scaffolds under static and dynamic (pulsatile distension) culture conditions. (iii) Determine the effect of serum withdrawal on the induction of the contractile phenotype of HCASMCs grown on 3D biodegradable polyurethane scaffolds. Research plan: Porous biodegradable polyurethane scaffolds will be fabricated and fibronectin immobilization will be carried out by a known ethylcarbodiimide (EDC)/N-hydroxy succinimide (NHS) activation method. ELISA methods will be used to ascertain fibronectin immobilization. Following HCASMC seeding, cell attachment and proliferation will be studied by fluorescence microscopy. Cellular differentiation will be evaluated by specific VSMC differentiation markers (h-caldesmon and smoothelin). We anticipate to promote long-term attachment of HCASMCs and modulate the synthetic phenotype. The effect of exogenous biomechanical signals (TGF-beta and retinoic acid) on elastin synthesis will be assessed by Laser Scanning Confocal Microscopy (LSCM) and Western blot analyses. Further quantification of elastin, following alkali extraction, will be determined by colorimetric absorbance. We anticipate HCASMCs to retain their synthetic phenotype and enhance elastin synthesis in response to these signals. We will also investigate the effect of biomechanical signals on cell phenotype and elastin synthesis by subjecting the scaffold to pulsatile distention. After a period of elastin production, we will induce the contractile phenotype of HCASMCs by serum withdrawal. The switching of media conditions in the bioreactor can easily be done since our bioreactor is capable of exchanging media under flow conditions. Following serum starvation, we will analyze expression of contractile proteins using Western blots. We anticipate upregulation of contractile proteins in response to serum withdrawal which is vital for engineered vessels to function properly.

Significance: Key information regarding vascular smooth muscle cell phenotype and elastin synthesis in 3D culture is important for understanding the factors that regulate tissue-engineered vessel maturation. The proposed study may lead to the fabrication of elastin containing vascular tissues.

SUMMARY OF PROPOSAL / RÉSUMÉ DE LA PROPOSITION

In the space provided below, state the objectives of the proposed research program and summarize the scientific approach, highlighting the novelty and expected significance of the work.

Dans l'espace prévu ci-dessous, énoncez les objectifs du programme de recherche proposé et résumez la démarche scientifique, en soulignant l'originalité et l'importance prévue des travaux.

Tumor metastasis follows a cascade of events that involves the departure of individual malignant cells from the primary tumor, their dissemination through the vasculature and their extravasation at preferred secondary sites in distant organs, where subsequently metastases form and impede normal organ function. This complex process is largely regulated by adhesive interactions of tumor cells between themselves, with the extracellular matrix and with vascular endothelial cells during transendothelial migration (diapedesis). These interactions are functionally linked to the ability of tumor cells to change shape, to migrate through extracellular matrix and to extend motile processes that breach the endothelial barrier during diapedesis. This research proposal focuses on understanding the molecular mechanisms that regulate interactions between breast tumour cells and endothelial cell, and specifically addresses the question whether and how communication via gap junctions between tumor cells and endothelial cells regulates cellular motility during diapedesis. Using a well characterized in vitro assay, that combines high-resolution laser scanning confocal microscopy (LSCM), and time-lapse video microscopy, with quantitative analysis of diapedesis our previous work revealed that introducing connexin 43 (Cx43) into communication-deficient HBL100 breast tumor cells increased diapedesis. This increase was dependent on functional Cx43 and gap junction intercellular communication between tumor cells and endothelial cells. The main goal of this proposal is to explore the molecular mechanisms controlling gap junction dependent regulation of diapedesis and to explore the role of RhoGTPases in tumor cell diapedesis. We hypothesize that connexin expression and/or heterocellular gap junction dependent signalling modulates the expression and activation of specific RhoGTPases that influence tumor cell motility and interactions with the endothelium required for diapedesis, and that different connexins may play differential roles in this process. In Aim 1 we will determine if ectopic expression of the endothelial connexins Cx43, Cx37, and Cx40 in HBL100 tumor cells changes the expression levels and state of activation of RhoA, Rac-1 and CDC42. We will use western blot analysis and pull-down assays to quantify levels of total and active forms of these RhoGTPases. Our in vitro diapedesis assay will be used to correlate connexin expression, RhoGTPase levels and activity with changes in cellular behaviour at different stages during diapedesis. To confirm that observed changes in cellular behaviour is due to changes in RhoGTPase activity we will in Aim 2 differentially activate or inhibit RhoA, Rac-1 and CDC42 by expressing constitutively active and dominant negative forms of the GTPases in HBL-100 cells and by using pharmacological inhibitors. The effects on cell motility, cell shape and diapedesis will be examined by time-lapse microscopy and LSCM following labelling for F-actin and VE-cadherin. Cell adhesion to endothelium and extracellular matrix components will be quantified using adhesion assays and by quantifying focal contacts and cellular shape changes. To examine if tumour cell diapedesis is mediated by contact dependent modulation of interendothelial cell adhesion and motility through endothelial Rho GTPases we will in Aim 3 explore if tumor cells secrete factors that modulate RhoGTPase activity in endothelial cells and lead to modulation of endothelial cell-cell junctions. We will treat endothelial monolayers with tumour cell conditioned media and examine, using biochemical approaches, the composition and state of activation of adherens junctions and complexes known to affect endothelial contractility such as MLCK and RhoGTPases. In Aim 4 we will examine if Rho GTPases participate in endothelial responses required for tumour cell diapedesis. We will express constitutively active and dominant negative forms of the small GTPases RhoA, Rac-1 and Cdc42 in endothelial monolayers and examine the effect on connexin expression, gap junctional communication and tumor cell diapedesis. Using inhibitors and activators of MLCK, PI3-kinase and src kinases under these conditions will dissect signalling pathways that play a role in the endothelial response during diapedesis. Combining state of the art imaging techniques with molecular biology, biochemical and functional single cell analysis the experiments proposed focus on molecular events that regulate adhesion, motility, and diapedesis of breast tumor cells. We believe that these studies will therefore further our understanding of the molecular mechanisms that regulate tumour cell transendothelial migration and may point to novel avenues to design treatment regimen during early stages of cancer metastasis.

Cell Biology

ATCC® Number: **CRL-1740™** Price: **\$256.00**

Designations: LNCaP clone FGC Depositors: JS Horoszewicz

Biosafety Level: 1 Shipped: frozen

Medium & Serum: See Propagation Growth Properties: adherent, single cells and loosely attached clusters epithelial

Organism: *Homo sapiens* (human) Morphology: 

Source: **Organ:** prostate
Disease: carcinoma
Derived from metastatic site: left supraclavicular lymph node

Cellular Products: human prostatic acid phosphatase; prostate specific antigen [21889]

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.

[Related Cell Culture Products](#)

Restrictions: Distribution of this material for commercial purposes will require execution of a Non-exclusive License Agreement. At the time of placing an order, customers must send a request to licensing@ATCC.org. Orders will be shipped when Customer Service receives confirmation from our Licensing officer.

Isolation: **Isolation date:** 1977

Applications: transfection host ([technology from amaxa Roche FuGENE® Transfection Reagents](#))

Receptors: androgen receptor, positive; estrogen receptor, positive [23045]

Tumorigenic: Yes

Cytogenetic Analysis: This is a hypotetraploid human cell line. The modal chromosome number was 84, occurring in 22% of cells. However, cells with chromosome counts of 86 (20%) and 87 (18%) also occurred at high frequencies. The rate of cells with higher ploidies was 6.0%.

Age: 50 years adult

Gender: male

Ethnicity: Caucasian

LNCaP clone FGC was isolated in 1977 by J.S. Horoszewicz, et al., from a needle aspiration biopsy of the left supraclavicular lymph node of a 50-year-old Caucasian male (blood type B+) with confirmed diagnosis of metastatic prostate carcinoma. [21889]

These cells are responsive to 5-alpha-dihydrotestosterone (growth modulation and acid phosphatase production). [23045]

The cells do not produce a uniform monolayer, but grow in clusters which should

be broken apart by repeated pipetting when subcultures are prepared. They attach only lightly to the substrate, do not become confluent and rapidly acidify the medium.

Growth is very slow.

The cells should be allowed to incubate undisturbed for the first 48 hours after subculture.

Comments:

When flask cultures are shipped, the majority of the cells become detached from the flask and float in the medium.

Upon receipt, incubate the flask (in the usual position for monolayer cultures) for 24 to 48 hours to allow the cells to re-attach.

The medium can then be removed and replaced with fresh medium.

If desired, the contents of the flask can be collected, centrifuged at 300 X g for 15 minutes, resuspended in 10 ml of medium and dispensed into a single flask.

Propagation:

ATCC complete growth medium: The base medium for this cell line is ATCC-formulated RPMI-1640 Medium, Catalog No. 30-2001. To make the complete growth medium, add the following components to the base medium: fetal bovine serum to a final concentration of 10%.

Atmosphere: air, 95%; carbon dioxide (CO₂), 5%

Temperature: 37.0°C

Protocol:

1. Remove and discard culture medium.
2. Briefly rinse the cell layer with 0.25% (w/v) Trypsin- 0.53 mM EDTA solution to remove all traces of serum that contains trypsin inhibitor.
3. 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 15 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.

Subculturing:

4. Add 6.0 to 8.0 ml of complete growth medium and aspirate cells by gently pipetting.
5. Add appropriate aliquots of the cell suspension to new culture vessels. Maintain cultures at a cell concentration between 1 X 10⁴ and 2 X 10⁵ cells/cm².
6. Incubate cultures at 37°C.

Subcultivation Ratio: A subcultivation ratio of 1:3 to 1:6 is recommended

Medium Renewal: Twice per week

Preservation:

Freeze medium: Complete growth medium supplemented with 5% (v/v) DMSO

Storage temperature: liquid nitrogen vapor phase

Doubling Time:

about 34 hours

Recommended medium (without the additional supplements or serum described under ATCC Medium): ATCC 30-2001

Related Products:

recommended serum: ATCC 30-2020

derivative: ATCC CRL-10995

purified DNA: ATCC CRL-1740D

21889; . Models for prostate cancer. 37New York: Liss; 1980.

References:

- 22410: Gibas Z, et al. A high-resolution study of chromosome changes in a human prostatic carcinoma cell line (LNCaP). *Cancer Genet. Cytogenet.* 11: 399-404, 1984. PubMed: [6584201](#)
- 23045: Horoszewicz JS, et al. LNCaP model of human prostatic carcinoma. *Cancer Res.* 43: 1809-1818, 1983. PubMed: [6831420](#)
- 32283: Hu SX, et al. Development of an adenovirus vector with tetracycline-regulatable human tumor necrosis factor alpha gene expression. *Cancer Res.* 57: 3339-3343, 1997. PubMed: [9269991](#)
- 33090: Boffa LC, et al. Invasion of the CAG triplet repeats by a complementary peptide nucleic acid inhibits transcription of the androgen receptor and TATA-binding protein genes and correlates with refolding of an active nucleosome containing a unique AR gene sequence. *J. Biol. Chem.* 271: 13228-13233, 1996. PubMed: [8662737](#)

[Return to Top](#)

Cell Biology

ATCC® Number:	HTB-81™	Order this Item	Price:	\$264.00
Designations:	DU 145		Depositors:	KR Stone
Biosafety Level:	1		Shipped:	frozen
Medium & Serum:	See Propagation		Growth Properties:	adherent
Organism:	<i>Homo sapiens</i> (human)		Morphology:	epithelial

Source: **Organ:** prostate
Disease: carcinoma
Derived from metastatic site: brain

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.

[Related Cell Culture Products](#)

Applications: transfection host (technology from amaxa Roche FuGENE® Transfection Reagents)

Tumorigenic: YES

Antigen Expression: Blood Type O; Rh+

Cytogenetic Analysis: This is a hypotriploid human cell line. Both 61 and 62 chromosome numbers had the highest rate of occurrence in 30 metaphase counts. The rate of higher ploidies was 3%. The t(11q12q), del(11)(q23), 16q+, del(9)(p11), del(1)(p32) and 6 other marker chromosomes were found in most cells. The N13 was usually absent. The Y chromosome is abnormal through translocation to an unidentified chromosomal segment. The X chromosome was present in single copy.

Isoenzymes: AK-1, 1
ES-D, 1
G6PD, B
GLO-I, 2
Me-2, 1-2
PGM1, 1
PGM3, 2

Age: 69 years

Gender: male

Ethnicity: Caucasian

Comments: The line is not detectably hormone sensitive, is only weakly positive for acid phosphatase and isolated cells form colonies in soft agar. The cells do not express prostate antigen. Ultrastructural analyses of both the cell line and original tumor revealed microvilli, tonofilaments, desmosomes, any mitochondria, well developed Golgi and heterogenous lysosomes.

Propagation: **ATCC complete growth medium:** The base medium for this cell line is ATCC-formulated Eagle's Minimum Essential Medium, Catalog No. 30-2003. To make the complete growth medium, add the following components to the base medium: fetal bovine serum to a final concentration of 10%.

Atmosphere: air, 95%; carbon dioxide (CO₂), 5%

Temperature: 37.0°C

Protocol:

1. Remove and discard culture medium.
2. Briefly rinse the cell layer with 0.25% (w/v) Trypsin - 0.53 mM EDTA solution to remove all traces of serum which contains trypsin inhibitor.
3. 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 15 minutes).

Subculturing:

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 37C to facilitate dispersal.

4. Add 6.0 to 8.0 ml of complete growth medium and aspirate cells by gently pipetting.
5. Add appropriate aliquots of the cell suspension to new culture vessels.
6. Incubate cultures at 37C.

Subcultivation Ratio: A subcultivation ratio of 1:4 to 1:6 is recommended

Medium Renewal: 2 to 3 times per week

Preservation:

Freeze medium: Complete growth medium, 95%; DMSO, 5%

Storage temperature: liquid nitrogen vapor temperature

recommended serum: ATCC 30-2020

Recommended medium (without the additional supplements or serum described under ATCC Medium): ATCC 30-2003

Related Products:

purified DNA: ATCC HTB-81D

0.25% (w/v) Trypsin - 0.53 mM EDTA in Hank' BSS (w/o Ca⁺⁺, Mg⁺⁺): ATCC 30-2101

Cell culture tested DMSO: ATCC 4-X

22289: Papsidero LD, et al. Prostate antigen: a marker for human prostate epithelial cells. *J. Natl. Cancer Inst.* 66: 37-42, 1981. PubMed: 6935463

22858: Stone KR, et al. Isolation of a human prostate carcinoma cell line (DU 145). *Int. J. Cancer* 21: 274-281, 1978. PubMed: 631930

23028: Mickey DD, et al. Heterotransplantation of a human prostatic adenocarcinoma cell line in nude mice. *Cancer Res.* 37: 4049-4058, 1977.

PubMed: 908039

23226: Pollack MS, et al. HLA-A, B, C and DR alloantigen expression on forty-six cultured human tumor cell lines. *J. Natl. Cancer Inst.* 66: 1003-1012, 1981.

PubMed: 7017212

References:

32283: Hu SX, et al. Development of an adenovirus vector with tetracycline-regulatable human tumor necrosis factor alpha gene expression. *Cancer Res.* 57: 3339-3343, 1997. PubMed: 9269991

32341: Sheng S, et al. Maspin acts at the cell membrane to inhibit invasion and motility of mammary and prostatic cancer cells. *Proc. Natl. Acad. Sci. USA* 93: 11669-11674, 1996. PubMed: 8876194

32460: Carter RE, et al. Prostate-specific membrane antigen is a hydrolase with substrate and pharmacologic characteristics of a neuropeptidase. *Proc. Natl. Acad. Sci. USA* 93: 749-753, 1996. PubMed: 8570628

32486: Nupponen NN, et al. Genetic alterations in prostate cancer cell lines

detected by comparative genomic hybridization. *Cancer Genet. Cytogenet.* 101: 53-57, 1998. PubMed: 9460501

32768: Robinson D, et al. A tyrosine kinase profile of prostate carcinoma. *Proc. Natl. Acad. Sci. USA* 93: 5958-5962, 1996. PubMed: 8650201

32916: Su ZZ, et al. Surface-epitope masking and expression cloning identifies the human prostate carcinoma tumor antigen gene PCTA-1 a member of the galectin gene family. *Proc. Natl. Acad. Sci. USA* 93: 7252-7257, 1996. PubMed: 8692978

32925: Zhu X, et al. Cell cycle-dependent modulation of telomerase activity in tumor cells. *Proc. Natl. Acad. Sci. USA* 93: 6091-6095, 1996. PubMed: 8650224

33090: Boffa LC, et al. Invasion of the CAG triplet repeats by a complementary peptide nucleic acid inhibits transcription of the androgen receptor and TATA-binding protein genes and correlates with refolding of an active nucleosome containing a unique AR gene sequence. *J. Biol. Chem.* 271: 13228-13233, 1996. PubMed: 8662737

[Return to Top](#)

Cell Biology

ATCC® Number:	CRL-1435™ <input type="button" value="Order this Item"/>	Price:	\$256.00
Designations:	PC-3	Depositors:	ME Kaighn
Biosafety Level:	1	Shipped:	frozen
Medium & Serum:	See Propagation	Growth Properties:	adherent (The cells form clusters in soft agar and can be adapted to suspension growth) epithelial
Organism:	<i>Homo sapiens</i> (human)	Morphology:	
Source:	Organ: prostate Tumor Stage: grade IV Disease: adenocarcinoma Derived from metastatic site: bone		
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.		

Related Cell Culture Products

Applications:	transfection host (technology from amaxa Roche FuGENE® Transfection Reagents)
Tumorigenic:	YES
Antigen Expression:	HLA A1, A9 Amelogenin: X CSF1PO: 11 D13S317: 11 D16S539: 11
DNA Profile (STR):	D5S818: 13 D7S820: 8,11 THO1: 6,7 TPOX: 8,9 vWA: 17
Cytogenetic Analysis:	The line is near-triploid with a modal number of 62 chromosomes. There are nearly 20 marker chromosomes commonly found in each cell; and normal N2, N3, N4, N5, N12, and N15 are not found. No normal Y chromosomes could be detected by Q-band analysis.
Age:	62 years adult
Gender:	male
Ethnicity:	Caucasian
Comments:	The PC-3 was initiated from a bone metastasis of a grade IV prostatic adenocarcinoma from a 62-year-old male Caucasian. [22363] The cells exhibit low acid phosphatase and testosterone-5-alpha reductase activities.

Propagation:	<p>ATCC complete growth medium: The base medium for this cell line is ATCC-formulated F-12K Medium, Catalog No. 30-2004. To make the complete growth medium, add the following components to the base medium: fetal bovine serum to a final concentration of 10%.</p> <p>Atmosphere: air, 95%; carbon dioxide (CO₂), 5%</p> <p>Temperature: 37.0°C</p> <p>Protocol:</p> <ol style="list-style-type: none"> 1. Remove and discard culture medium. 2. Briefly rinse the cell layer with 0.25% (w/v) Trypsin- 0.53 mM EDTA solution to remove all traces of serum that contains trypsin inhibitor. 3. 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 15 minutes).
Subculturing:	<p>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.</p> <ol style="list-style-type: none"> 4. Add 6.0 to 8.0 ml of complete growth medium and aspirate cells by gently pipetting. 5. Add appropriate aliquots of the cell suspension to new culture vessels. 6. Incubate cultures at 37°C.
Preservation:	<p>Subcultivation Ratio: A subcultivation ratio of 1:3 to 1:6 is recommended</p> <p>Medium Renewal: 2 to 3 times per week</p> <p>Freeze medium: Complete growth medium supplemented with 5% (v/v) DMSO</p> <p>Storage temperature: liquid nitrogen vapor phase</p>
Related Products:	<p>Recommended medium (without the additional supplements or serum described under ATCC Medium): ATCC 30-2004</p> <p>recommended serum: ATCC 30-2020</p>
References:	<p>22363: Kaighn ME, et al. Establishment and characterization of a human prostatic carcinoma cell line (PC-3). <i>Invest. Urol.</i> 17: 16-23, 1979. PubMed: 447482</p> <p>22470: Chen TR. Chromosome identity of human prostate cancer cell lines, PC-3 and PPC-1. <i>Cytogenet. Cell Genet.</i> 62: 183-184, 1993. PubMed: 8428522</p> <p>26302: Ohnuki Y, et al. Chromosomal analysis of human prostatic adenocarcinoma cell lines. <i>Cancer Res.</i> 40: 524-534, 1980. PubMed: 7471073</p> <p>32341: Sheng S, et al. Maspin acts at the cell membrane to inhibit invasion and motility of mammary and prostatic cancer cells. <i>Proc. Natl. Acad. Sci. USA</i> 93: 11669-11674, 1996. PubMed: 8876194</p> <p>32344: Umekita Y, et al. Human prostate tumor growth in athymic mice: inhibition by androgens and stimulation by finasteride. <i>Proc. Natl. Acad. Sci. USA</i> 93: 11802-11807, 1996. PubMed: 8876218</p> <p>32460: Carter RE, et al. Prostate-specific membrane antigen is a hydrolase with substrate and pharmacologic characteristics of a neuropeptidase. <i>Proc. Natl. Acad. Sci. USA</i> 93: 749-753, 1996. PubMed: 8570628</p> <p>32486: Nupponen NN, et al. Genetic alterations in prostate cancer cell lines detected by comparative genomic hybridization. <i>Cancer Genet. Cytogenet.</i> 101: 53-57, 1998. PubMed: 9460501</p> <p>32488: Geiger T, et al. Antitumor activity of a PKC-alpha antisense oligonucleotide in combination with standard chemotherapeutic agents against</p>

various human tumors transplanted into nude mice. *Anticancer Drug Des.* 13: 35-45, 1998. PubMed: [9474241](#)
32916: Su ZZ, et al. Surface-epitope masking and expression cloning identifies the human prostate carcinoma tumor antigen gene PCTA-1 a member of the galectin gene family. *Proc. Natl. Acad. Sci. USA* 93: 7252-7257, 1996. PubMed: [8692978](#)

[Return to Top](#)

Cell Biology

ATCC® Number: **TIB-202™** Order this Item Price: **\$264.00**
 Designations: THP-1 Depositors: S Tsuchiya
 Biosafety Level: 1 Shipped: frozen
 Medium & Serum: See Propagation Growth Properties: suspension
 Organism: *Homo sapiens* (human) Morphology: 

Source: **Organ:** peripheral blood
Disease: acute monocytic leukemia
Cell Type: monocyte;

Cellular Products: lysozyme [58053]

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.

Related Cell Culture Products

Applications: transfection host (technology from amaxa Roche FuGENE® Transfection Reagents)

Receptors: complement (C3), expressed [58053]
 Fc, expressed

Antigen Expression: HLA A2, A9, B5, DRw1, DRw2 [58053]

Amelogenin: X,Y

CSF1PO: 11,13

D13S317: 13

D16S539: 11,12

DNA Profile (STR): D5S818: 11,12

D7S820: 10

THO1: 8,9,3

TPOX: 8,11

vWA: 16

Age: 1 year infant

Gender: male

Comments: The cells are phagocytic (for both latex beads and sensitized erythrocytes) and lack surface and cytoplasmic immunoglobulin. [58053]

Monocytic differentiation can be induced with the phorbol ester 12-O-tetradecanoylphorbol-13-acetate (TPA). [22193]

Propagation: **ATCC complete growth medium:** The base medium for this cell line is ATCC-formulated RPMI-1640 Medium, Catalog No. 30-2001. To make the complete growth medium, add the following components to the base medium: 2-mercaptoethanol to a final concentration of 0.05 mM; fetal bovine serum to a final concentration of 10%.

Atmosphere: air, 95%; carbon dioxide (CO₂), 5%

Temperature: 37.0°C

- Protocol:** Cultures can be maintained by the addition of fresh medium or replacement of medium. Alternatively, cultures can be established by centrifugation with subsequent resuspension at 2-4 X 10⁴ viable cells/ml. Subcultre when cell concentration reaches 8X10⁵ cells/ml. Do not allow the cell concentration to exceed 1 X 10⁶ cells/ml.
- Medium Renewal:** Every 2 to 3 days
- Freeze medium:** Complete growth medium supplemented with 5% (v/v) DMSO
- Storage temperature:** liquid nitrogen vapor phase
- Subculturing:** approximately 26 hrs
- Preservation:** purified DNA:ATCC TIB-202D
purified RNA:ATCC TIB-202R
- Doubling Time:** approximately 26 hrs
- Related Products:** 22193: Tsuchiya S, et al. Induction of maturation in cultured human monocytic leukemia cells by a phorbol diester. *Cancer Res.* 42: 1530-1536, 1982. PubMed: [6949641](#)
22285: Skubitz KM, et al. Human granulocyte surface molecules identified by murine monoclonal antibodies. *J. Immunol.* 131: 1882-1888, 1983. PubMed: [6619543](#)
32286: Cuthbert JA, Lipsky PE. Regulation of proliferation and Ras localization in transformed cells by products of mevalonate metabolism. *Cancer Res.* 57: 3498-3504, 1997. PubMed: [9270019](#)
32351: Huang S, et al. Adenovirus interaction with distinct integrins mediates separate events in cell entry and gene delivery to hematopoietic cells. *J. Virol.* 70: 4502-4508, 1996. PubMed: [8676475](#)
32395: Clark RA, et al. Tenascin supports lymphocyte rolling. *J. Cell Biol.* 137: 755-765, 1997. PubMed: [9151679](#)
32466: Hambleton J, et al. Activation of c-Jun N-terminal kinase in bacterial lipopolysaccharide-stimulated macrophages. *Proc. Natl. Acad. Sci. USA* 93: 2774-2778, 1996. PubMed: [8610116](#)
33031: Hsu HY, et al. Inhibition of macrophage scavenger receptor activity by tumor necrosis factor-alpha is transcriptionally and post-transcriptionally regulated. *J. Biol. Chem.* 271: 7767-7773, 1996. PubMed: [8631819](#)
33088: Lucas M, Mazzone T. Cell surface proteoglycans modulate net synthesis and secretion of macrophage apolipoprotein E. *J. Biol. Chem.* 271: 13454-13460, 1996. PubMed: [8662812](#)
33134: Sando GN, et al. Induction of ceramide glucosyltransferase activity in cultured human keratinocytes. *J. Biol. Chem.* 271: 22044-22051, 1996. PubMed: [8703011](#)
33141: Ollivier V, et al. Elevated cyclic AMP inhibits NF-kappaB-mediated transcription in human monocytic cells and endothelial cells. *J. Biol. Chem.* 271: 20828-20835, 1996. PubMed: [8702838](#)
58053: Tsuchiya S, et al. Establishment and characterization of a human acute monocytic leukemia cell line (THP-1). *Int. J. Cancer* 26: 171-176, 1980. PubMed: [6970727](#)
- References:**

[Return to Top](#)

Cell Biology

ATCC® Number: **CRL-1573™** Price: **\$256.00**
 Designations: 293 [HEK-293] Depositors: FL Graham
 Biosafety Level: 2 [CELLS CONTAIN ADENOVIRUS] Shipped: frozen
 Medium & Serum: See Propagation Growth Properties: adherent
 epithelial
 Organism: *Homo sapiens* (human) Morphology: 

Source: **Organ:** kidney
Cell Type: transformed with adenovirus 5 DNA

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.

[Related Cell Culture Products](#)

Restrictions: These cells are distributed for research purposes only. 293 cells, their products, or their derivatives may not be distributed to third parties.

Applications: efficacy testing [92587]
 transfection host (technology from amaxa Roche FuGENE® Transfection Reagents)
 viruscide testing [92579]

Receptors: vitronectin, expressed

Tumorigenic: YES

Amelogenin: X
 CSF1PO: 11,12
 D13S317: 12,14
 D16S539: 9,13

DNA Profile (STR): D5S818: 8,9
 D7S820: 11,12
 TH01: 7,9.3
 TPOX: 11
 vWA: 16,19

Cytogenetic Analysis: This is a hypotriploid human cell line. The modal chromosome number was 64, occurring in 30% of cells. The rate of cells with higher ploidies was 4.2 %. The der(1)t(1;15) (q42;q13), der(19)t(3;19) (q12;q13), der(12)t(8;12) (q22;p13), and four other marker chromosomes were common to most cells. Five other markers occurred in some cells only. The marker der(1) and M8 (or Xq+) were often paired. There were four copies of N17 and N22. Noticeably in addition to three copies of X chromosomes, there were paired Xq+, and a single Xp+ in most cells.

Age: fetus

Although an earlier report suggested that the cells contained Adenovirus 5 DNA from both the right and left ends of the viral genome [RF32764], it is now clear that only left end sequences are present. [39768]
 The line is excellent for titrating human adenoviruses.

Comments:	<p>The cells express an unusual cell surface receptor for vitronectin composed of the integrin beta-1 subunit and the vitronectin receptor alpha-v subunit. [23406] The Ad5 insert was cloned and sequenced, and it was determined that a colinear segment from nts 1 to 4344 is integrated into chromosome 19 (19q13.2). [39768] ATCC complete growth medium: The base medium for this cell line is ATCC-formulated Eagle's Minimum Essential Medium, Catalog No. 30-2003. To make the complete growth medium, add the following components to the base medium: fetal bovine serum to a final concentration of 10%.</p>
Propagation:	<p>Atmosphere: air, 95%; carbon dioxide (CO₂), 5% Temperature: 37.0°C The cell line does not adhere to the substrate when left at room temperature for any length of time, therefore, live cultures may be received with the cells detached. The cells will re-attach to the flask over a period of several days in culture at 37C.</p>
Subculturing:	<p>Protocol:</p> <ol style="list-style-type: none"> 1. Remove and discard culture medium. 2. Briefly rinse the cell layer with 0.25% (w/v) Trypsin- 0.53 mM EDTA solution to remove all traces of serum that contains trypsin inhibitor. 3. 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 15 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. 4. Add 6.0 to 8.0 ml of complete growth medium and aspirate cells by gently pipetting. 5. Add appropriate aliquots of the cell suspension to new culture vessels. An inoculum of 2 X 10³ to 6 X 10³ viable cells/cm² is recommended. 6. Incubate cultures at 37°C. 6. Subculture when cell concentration is between 6 and 7 X 10⁴ cells/cm².
Preservation:	<p>Subcultivation Ratio: 1:10 to 1:20 weekly. Medium Renewal: Every 2 to 3 days Freeze medium: Complete growth medium supplemented with 5% (v/v) DMSO Storage temperature: liquid nitrogen vapor phase Recommended medium (without the additional supplements or serum described under ATCC Medium): ATCC 30-2003 derivative: ATCC CRL-10852 derivative: ATCC CRL-12006 derivative: ATCC CRL-12007 derivative: ATCC CRL-12013 derivative: ATCC CRL-12479 derivative: ATCC CRL-2029 derivative: ATCC CRL-2368 purified DNA: ATCC CRL-1573D</p>
Related Products:	<p>21624: Xie QW, et al. Complementation analysis of mutants of nitric oxide synthase reveals that the active site requires two hemes. Proc. Natl. Acad. Sci.</p>

- USA 93: 4891-4896, 1996. PubMed: 8643499
- 21631: Da Costa LT, et al. Converting cancer genes into killer genes. Proc. Natl. Acad. Sci. USA 93: 4192-4196, 1996. PubMed: 8633039
- 22282: Graham FL, et al. Characteristics of a human cell line transformed by DNA from human adenovirus type 5. J. Gen. Virol. 36: 59-72, 1977. PubMed: 886304
- 22319: Graham FL, et al. Defective transforming capacity of adenovirus type 5 host-range mutants. Virology 86: 10-21, 1978. PubMed: 664220
- 22699: Harrison T, et al. Host-range mutants of adenovirus type 5 defective for growth in HeLa cells. Virology 77: 319-329, 1977. PubMed: 841862
- 23406: Bodary SC, McLean JW. The integrin beta 1 subunit associates with the vitronectin receptor alpha v subunit to form a novel vitronectin receptor in a human embryonic kidney cell line. J. Biol. Chem. 265: 5938-5941, 1990. PubMed: 1690718
- 27819: Goodrum FD, Ornelles DA. The early region 1B 55-kilodalton oncoprotein of adenovirus relieves growth restrictions imposed on viral replication by the cell cycle. J. Virol. 71: 548-561, 1997. PubMed: 8985383
- 28301: Loffler S, et al. CD9, a tetraspan transmembrane protein, renders cells susceptible to canine distemper virus. J. Virol. 71: 42-49, 1997. PubMed: 8985321
- 32283: Hu SX, et al. Development of an adenovirus vector with tetracycline-regulatable human tumor necrosis factor alpha gene expression. Cancer Res. 57: 3339-3343, 1997. PubMed: 9269991
- 32396: Kolanus W, et al. alphaLbeta2 integrin/LFA-1 binding to ICAM-1 induced by cytohesin-1 a cytoplasmic regulatory molecule. Cell 86: 233-242, 1996. PubMed: 8706128
- 32490: Stauderman KA, et al. Characterization of human recombinant neuronal nicotinic acetylcholine receptor subunit combinations alpha 2 beta 4, alpha 3 beta 4 and alpha 4 beta 4 stably expressed in HEK293 cells. J. Pharmacol. Exp. Ther. 284: 777-789, 1998. PubMed: 9454827
- 32514: Bartz SR, et al. Human immunodeficiency virus type 1 cell cycle control: Vpr is cytostatic and mediates G2 accumulation by a mechanism which differs from DNA damage checkpoint control. J. Virol. 70: 2324-2331, 1996. PubMed: 8642659
- 32726: Sandri-Goldin RM, Hibbard MK. The herpes simplex virus type 1 regulatory protein ICP27 coimmunoprecipitates with anti-sm antiserum, and the C terminus appears to be required for this interaction. J. Virol. 70: 108-118, 1996. PubMed: 8523514
- 32829: Ansieau S, et al. Tumor necrosis factor receptor-associated factor (TRAF)-1, TRAF-2, and TRAF-3 interact in vivo with the CD30 cytoplasmic domain; TRAF-2 mediates CD30-induced nuclear factor kappa B activation. Proc. Natl. Acad. Sci. USA 93: 14053-14058, 1996. PubMed: 8943059
- 32893: Zhang J, et al. Dynamin and beta-arrestin reveal distinct mechanisms for G protein-coupled receptor internalization. J. Biol. Chem. 271: 18302-18305, 1996. PubMed: 8702465
- 32914: Oppermann M, et al. Monoclonal antibodies reveal receptor specificity among G-protein-coupled receptor kinases. Proc. Natl. Acad. Sci. USA 93: 7649-7654, 1996. PubMed: 8755530
- 32921: Xia Y, et al. Nitric oxide synthase generates superoxide and nitric oxide in arginine-depleted cells leading to peroxynitrite-mediated cellular injury. Proc. Natl. Acad. Sci. USA 93: 6770-6774, 1996. PubMed: 8692893

- 32925: Zhu X, et al. Cell cycle-dependent modulation of telomerase activity in tumor cells. *Proc. Natl. Acad. Sci. USA* 93: 6091-6095, 1996. PubMed: 8650224
- 32971: Uebele VN, et al. Functional differences in Kv1.5 currents expressed in mammalian cell lines are due to the presence of endogenous Kvbeta2.1 subunits. *J. Biol. Chem.* 271: 2406-2412, 1996. PubMed: 8576199
- 33003: Abell A, et al. Deletions of portions of the extracellular loops of the lutropin/choriogonadotropin receptor decrease the binding affinity for ovine luteinizing hormone, but not human choriogonadotropin, by preventing the formation of mature cell surface receptor. *J. Biol. Chem.* 271: 4518-4527, 1996. PubMed: 8626807
- 33010: Tiberi M, et al. Differential regulation of dopamine D1A receptor responsiveness by various G protein-coupled receptor kinases. *J. Biol. Chem.* 271: 3771-3778, 1996. PubMed: 8631993
- 33022: Shahrestanifar M, et al. Studies on inhibition of mu and delta opioid receptor binding by dithiothreitol and N-ethylmaleimide. His223 is critical for mu opioid receptor binding and inactivation by N-ethylmaleimide. *J. Biol. Chem.* 271: 5505-5512, 1996. PubMed: 8621408
- 33035: Boring L, et al. Molecular cloning and functional expression of murine JE (monocyte chemoattractant protein 1) and murine macrophage inflammatory protein 1alpha receptors. *J. Biol. Chem.* 271: 7551-7558, 1996. PubMed: 8631787
- 33036: Noonberg SB, et al. Evidence of post-transcriptional regulation of U6 small nuclear RNA. *J. Biol. Chem.* 271: 10477-10481, 1996. PubMed: 8631843
- 33050: Fox JC, Shanley JR. Antisense inhibition of basic fibroblast growth factor induces apoptosis in vascular smooth muscle cells. *J. Biol. Chem.* 271: 12578-12584, 1996. PubMed: 8647868
- 33056: Lee MJ, et al. The inducible G protein-coupled receptor edg-1 signals via the Gi/mitogen-activated protein kinase pathway. *J. Biol. Chem.* 271: 11272-11279, 1996. PubMed: 8626678
- 33123: Marchand P, et al. Cysteine mutations in the MAM domain result in monomeric meprin and alter stability and activity of the proteinase. *J. Biol. Chem.* 271: 24236-24241, 1996. PubMed: 8798668
- 33137: Arai H, Charo IF. Differential regulation of G-protein-mediated signaling by chemokine receptors. *J. Biol. Chem.* 271: 21814-21819, 1996. PubMed: 8702980
- 33138: Huang Q, et al. Substrate recognition by tissue factor-factor VIIa. *J. Biol. Chem.* 271: 21752-21757, 1996. PubMed: 8702971
- 33157: Monteclaro FS, Charo IF. The amino-terminal extracellular domain of the MCP-1 receptor, but not the RANTES/MIP-1alpha receptor, confers chemokine selectivity. *J. Biol. Chem.* 271: 19084-19092, 1996. PubMed: 8702581
- 33158: Keith DE, et al. Morphine activates opioid receptors without causing their rapid internalization. *J. Biol. Chem.* 271: 19021-19024, 1996. PubMed: 8702570
- 39768: Louis N, et al. Cloning and sequencing of the cellular-viral junctions from the human adenovirus type 5 transformed 293 cell line. *Virology* 233: 423-429, 1997. PubMed: 9217065
- 61259: Shaw G, et al. Preferential transformation of human neuronal cells by human adenoviruses and the origin of HEK 293 cells. *FASEB J.* 16: 869-871, 2002. PubMed: 11967234
- 92579: Standard Test Method for Determining the Virus-Eliminating Effectiveness of Liquid Hygienic Handwash and Handrub Agents Using the Fingerpads of Adult Volunteers. West Conshohocken, PA:ASTM International;ASTM Standard Test Method E 1838-02.

References:

92587: Standard Quantitative Disk Carrier Test Method for Determining the Bactericidal, Virucidal, Fungicidal, Mycobactericidal and Sporocidal Activities of Liquid Chemical Germicides. West Conshohocken, PA:ASTM International;ASTM Standard Test Method E 2197-02.

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