Introduction...
by Bill Oetting

The PanAmerican Society for Pigment Cell Research has two major sources of information about our society, the PASPCR Newsletter, and the PASPCR Web Site. The internet presence of the PASPCR began as a Gopher site in February 28, 1994 (the PASPCR Gopher site still exists), and was administered by DeWayne Townsend, at the University of Minnesota. Vince Hearing created the first PASPCR Web Site in early 1995, and later that year the site was moved to the same server that contained the PASPCR Gopher site. Since then, the PASPCR Web Site has undergone numerous alterations. At present, the web site receives over 16 ‘hits’ per day, and has accumulated over 30,000 visits since 1996. This number does not include multiple hits from the same IP address, but represents different individuals who are visiting the PASPCR Web Site.

The PASPCR Web Site is the major, up-to-date source of current information for the PASPCR membership and for individuals who are interested in the PASPCR. The PASPCR Web site is not only includes new information, but it is also a repository on the history of our society, our goals and our by-laws. This site also includes a registration page for potential new members and I have received well over 30 requests over the last two years. This is a very cost effective means to disseminate information about the PASPCR. If there is additional information that you would like to see on the Web site, or you would like to include information of past PASPCR activities, please let me know and I will add them. A visit to our web site may be the first time that an individual has heard about the PASPCR, and we should present ourselves in the best possible light. I look forward to your comments and inclusions.

The PASPCR Web Site can be found at:
http://www.paspcr.org

The PASPCR Newsletter is published quarterly and is intended to serve as a means of communication for the members of our Society. You are invited to contribute articles, or other information you feel will be of interest to members of the PASPCR. If you attend a scientific meeting and have heard results which you think will be of interest to the membership of the PASPCR, please write a few paragraphs summarizing what was presented and share it with us. Any information on upcoming meetings of interest will be added to the “Calendar of Events”. We also want to note any change of affiliation or address that you may have had to help us keep our membership list up-to-date. This is your Newsletter, and we depend upon you to help us make sure it best serves the Society’s needs. Contributions and comments can be sent to me, preferably by E-mail, to bill@lenti.med.umn.edu.

Due to problems with the domain names obtained for the IFPCS, the addresses IFPCS.info and IPCC.info are currently not active. I am trying to get these addresses back, but there is a possibility that they will be lost. I will let you know the results when they become available.

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The PanAmerican Society for Pigment Cell Research

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Richard A. King

Calendar of Events:

July 14-19, 2002  XIXth IUPAC Conference on Photochemistry will be held in Budapest.
Contact: The Hungarian Chemical Society, (MKE), Fu u. 68. Hungary, H-1027 Budapest,
Phone: 36-1-201-6886
Fax: 36-1-201-8056
E-Mail: mail.mke@mtesz.hu.

Sept. 9 - 13, 2002  The XVIIIth International Pigment Cell Conference, to be held in The Hague, Holland.
Contact: Dr. Stan Pavel, President ESPCR,
University Hospital Leiden, Dept of Dermatology,
PO Box 9600, NL - 2300 RC LEIDEN
Phone: 31-(71) 526 1952
Fax: 31-(71) 524 8106;
E-mail: SPavel@algemeen.azl.nl
Website: http://www.ipcc.info

Dec. 14-18, 2002  The American Society for Cell Biology 42nd Annual Meeting is to be held in San Francisco, CA
For more information: www.ascb.org

Sept. 3-7, 2003  XIth Annual Meeting of the PanAmerican Society for Pigment Cell Research, to be held in Wood’s Hole, MA.
Contact: Dr. Jean Bologna.
E-mail: jean_bologna@qm.yale.edu.

Sept. 17-20, 2003  XIth Annual Meeting of the European Society for Pigment Cell Research, to be held in Gent.
Contact: Prof. JeanMarie Naeyaert.
E-mail: JeanMarie.Naeyaert@rug.ac.be.

The PASPCR Newsletter is published quarterly by the PanAmerican Society for Pigment Cell Research. All views are those of the authors. For further information or to submit articles, please contact members of the Publications Committee.

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Corporate Sponsors
by Raymond E. Boissy

The PASPCR would like to acknowledge and thank our Corporate Sponsors; the list below reflects contributions over the past 2 years. Financial gifts from these sponsors have allowed our Society to increase benefits to the membership far out of proportion to the actual dues collected from members. Monies contributed by these sponsors have been used over the years to support various PASPCR functions including our Young Investigator Award program, meeting travel stipends, annual meeting expenses and this Newsletter.

**GOLD Corporate Patrons**
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**SILVER Corporate Patrons**
- Avon Products, Inc.
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- Stiefel Laboratories
- Combe, Inc.

Some Items of Interest:
The American Society for Cell Biology 42nd Annual Meeting is in San Francisco, December 14-18, 2002. Full details about the meeting are on the ASCB website www.ascb.org. If you would like to receive a copy of the Call for Abstracts send an e-mail to ascbinfo@ascb.org The Abstract deadline is August 1. The meeting opens on Saturday afternoon, December 14 with Special Interest Subgroups. The opening Reception is Saturday Evening at 6:00 PM and is titled, “Opportunities & Challenges in Cell Biology”. The Chairs are Steven M. Block, R. Alta Charo, Ron McKay, and Andrew W. Murray. Sunday, December 15-Wednesday December 18 there will be two morning Symposia, and each afternoon will be your choice of 8 Minisymposia. Over 3,000 Poster Sessions will be presented. The Exhibit Hall will feature products and services for use by Cell Biologists. Over 350 companies will be Exhibiting. If you are looking for a job, or have a job to fill, there will be a career center located next to the Exhibit Hall. Advanced Registration ends October 1. Minorities travel Award applications and Predoctoral travel Award Applications must be received by August 1.

From Miri Seiberg

Johnson & Johnson Skin Research Center training grant program

The Skin Research Center of the Johnson & Johnson Consumer Companies, Inc. investigates the molecular, cellular and physiological mechanisms involved in skin homeostasis and diseases. The Skin Research Center is committed to enhancing global scientific interactions and collaborations in the area of skin research and to enabling young scientists to become a part of the skin research community.

The Johnson & Johnson SRC training grant program supports postdoctoral training of basic research in skin biology or physiology in key universities, colleges and research institutions around the world. A key component of the Johnson & Johnson SRC training grant is the promotion of mutually beneficial relationships between Johnson & Johnson SRC scientists, grant recipients and their laboratories. The SRC training grant program seeks new scientific developments that could, over time, prove beneficial to Johnson & Johnson. More information on this program could be found at the PASPCR Web site: www.paspcr.org/P&Pskingrant.pdf.

From Lynn Lamoreux

We will discontinue the following stocks at the end of June. Anyone who wants these stocks should contact Lynn Lamoreux. All are congenic with C57BL/6J

- Belted
- Silver with black (this is not available elsewhere on B6)
- Silver with brown
- Slaty with chinchilla
- Slaty with brown
- Chinchilla with brown
Model Organisms

by Lynn Lamoreux

メダカ
(Medaka)

The Japanese medaka is a vertebrate research model that offers particular advantages to pigment cell research. This hardy little fish produces a dozen or more eggs at a predictable time each day, and it retains them attached to the female for several hours. Medaka is easy and inexpensive to raise and to transport; thus is suitable for research projects at all levels from the classroom to the sophisticated research laboratory. The domesticated fish models, of course, avoid problems faced in working with mammals or species that must be collected in the wild. At the same time, medaka uniquely satisfies some specific needs for biological research. For example, the See-Through Medaka is essentially transparent throughout its lifetime because it lacks most differentiated pigment cells. Thus, the See-Through Medaka is a unique and convenient organism for any research that will benefit from continual observation of a living vertebrate at all levels of organization from the whole organism to fluorescent-tagged cells/molecules.

The See-Through Medaka was created by Mendelian genetics, using mutations that prevent normal function of the four different types of medaka chromatophores. The mutations used to make the see-through medaka are available in all combinations. Thus the embryonic and adult development of the four types of melanophores can be observed in presence or absence of differentiated representatives of any or all of the other melanophore types. The rich resources of medaka mutants, combined with data obtainable from other species, especially zebra fish and mouse, offer information not otherwise readily available regarding the chromatophore lineage.

Mouse News

by Lynn Lamoreux

Congenic inbred mice are important to evaluation of gene functions and genic interactions because the functions and interactions can be evaluated in absence of confounding changes in the background genome. However, the exclusive use of C57BL/6J congenics would eventually lead us to an unnaturally narrow view of the range of variability that can be expressed as a result of change in a given gene. Furthermore, genes to not function in isolation, and the expression of a gene against one inbred background will not tell us the rest of the story regarding its relationship with the genome as a whole. One method of evaluating the influence of background genome upon a specific gene function or genic interaction is place the same specific gene(s) in a contrasting genetic background.

One such opportunity exists when white-spotting genotypes are compared on JUCtLm and C57BL/6J. Differences become apparent. Dr. Tomohisa Hirobe has reported differences in skin pigmentation when standard genes that affect melanogenesis are placed on a C57BL/10 background. For readers who are interested in using JU, please contact Lynn Lamoreux or check out the stocks maintained at the Wellcome Trust Functional Genomics Mouse Pigmentary Mutants Repository. http://www.sghms.ac.uk/depts/anatomy/pages/WTFGMPMR.htm

Dr. Hirobe’s stocks are listed below.

- C57BL/10JHir (black)
- C57BL/10JHir-A/A (agouti)
- C57BL/10JHir-b/b (brown) Tyrp1 locus
- C57BL/10JHir-c/c (albino) Tyr locus
- C57BL/10JHir-d/d (dilute)
- C57BL/10JHir-p/p (pink-eyed dilution)
- C57BL/10JHir-Wsh/Wsh (sash) Kit locus
- C57BL/10JHir-e/e (recessive yellow) Mc1r locus
- C57BL/10JHir-slt/slt (slaty)
- C3H/HeJ/msHir

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For more information on the Medaka fish and different mutants that are available, visit The Medaka Homepage at http://bioll.bio.nagoya-u.ac.jp/
Keep the membership informed.

If you have news about a member of the PASPCR, please let us know. Contact a member of the publications committee and we will make sure that it is in the next issue.

Positions - Wanted and Available

Postings for Positions Available will be open to all individuals and institutions so long as the position is related to pigment cell research. Postings for Positions Wanted will be open only to members of the PanAmerican Society for Pigment Cell Research or its sister societies (JSPCR and ESPCR). Send postings to Bill Oetting at bill@lenti.med.umn.edu. Please provide an expiration date for any submitted postings. Final decisions will be made by the Publications Committee of the PASPCR.

Postdoctoral Position
Polarized Kit-ligand expression in the epidermis: Its role in human melanocyte homeostasis

A postdoctoral position (fully funded for the first year with the possibility of a 2 year extension) is immediately available in the Department of Pathology, Centre Medical Universitaire at the University of Geneva, Switzerland. The project is supervised by Dr. Bernhard Wehrle-Haller and Prof. Beat Imhof and is within the frame of a collaboration between the University of Geneva and Industry.

The aim of this project is to understand the role of kit-ligand in melanocyte homeostasis in the adult epidermis and how manipulation of kit-ligand expression or localization in keratinocytes affect melanocyte behavior. The project will employ cell-biological, pharmaceutical, biochemical as well transgenic approaches (mouse) to develop methods to modify Kit-ligand localization (polarity and cell surface expression) in vivo and to study melanocyte behavior in response to such altered Kit-ligand presentation. For references and rational see Wehrle-Haller and Imhof (2001, J. Biol. Chem. 276, 12667-74) and Grichnik et al., (1998, J. Invest. Dermatol. 111, 233-38).

The Centre Medical Universitaire provides a stimulatory research environment located within the City of Geneva. Research in the department is centered around problems of autoimmunity, wound healing, inflammation, cell-cell junctions and cell migration. Geneva, located at the lake of Geneva in close proximity to the French Alps, provides a rich multicultural environment facilitating social integration.

Interested candidates preferably having experience in one or more of the aforementioned domains should send their CV (e.g. e-mail) including names and contacting information of two references to:

Bernhard Wehrle-Haller PhD
Department of Pathology
Centre Medical Universitaire
1. Rue Michel-Servet
1211 Geneva 4
Switzerland
Tel/Fax: 0041 22 702 5735 / 5746
Bernhard.Wehrle
Haller@medecine.unige.ch

Postdoctoral Research Position

A postdoctoral position is available immediately to study the transcriptional co-repressor and co-activator activities of the oncogenic protein Ski in human melanomas (PNAS (USA) 97:5924-5929, 2000). Seeking individuals with experience in EMSA, in vitro transcription-translation, site-directed mutagenesis and yeast two-hybrid screening. Interested individuals should send inquiries and applications (including CV, a brief description of past experience and future research interests, and the name of three references) to:

Estela E. Medrano, Ph.D.
Huffington Center on Aging
Baylor College of Medicine
One Baylor Plaza N-803.01
Houston, TX 77030

Baylor College of Medicine is an Equal Opportunity Employer

Research Associate/Post Doctoral Fellow Position Available

Position available for either an entry level postdoctoral fellow or a more senior research associate to study the molecular and cellular biology
of the melanocyte in general and the pathophysiology of vitiligo in specific. The research project will focus globally on the role of survival factors and apoptotic regulators on the viability of melanocytes in the skin and in culture. In addition, the project will focus on the genetic and cellular susceptibility of melanocytes from patients with vitiligo to under apoptosis in response to various stimuli. Postdoctoral fellow candidate should have experience with routine molecular and cellular techniques including cell culturing, site directed mutagenesis, and protein biochemistry. Research Associate candidate should have similar experiences utilizing the melanocyte system. Candidate will become part of an interactive research group focusing of various aspects of pigmentation in the Department of Dermatology and on skin physiology in the Skin Sciences Institute within the University of Cincinnati College of Medicine. Send curriculum vitae and list of three references to:

Raymond E. Boissy, Ph.D.
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Principal Scientist- Clinical Research - Skin Science Research

Unilever employs over 200 scientists at our New Jersey Laboratory who are dedicated to innovative and scientifically rigorous skin research programs. Our world sales exceed $40 billion so our programs have solid financial funding allowing for an innovative and challenging research culture. We currently have a full time opening that provides a unique opportunity to apply your basic science skills to human studies that impact the condition of skin for hundreds of millions people worldwide. We are seeking an expert in pigment biology or photobiology who can advance our knowledge and link laboratory research to clinically defined improvements of consumer skin problems. As a member of our skin research team, you will have an opportunity to work with other scientific experts in many fields including cell biology, biochemistry, measurement science and physical chemistry. You will also be encouraged to establish and maintain close ties to research in academic and government research communities.

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Postdoctoral Fellows - Cancer and Developmental Biology - Two NIH-funded positions are available for fellows interested in studying the Hedgehog signaling pathway in development and disease using skin as a model system. One project centers on defining the function of the Hedgehog pathway during skin appendage morphogenesis (Dev. Biol. 205: 1-9, 1999); a second project focuses on understanding how deregulated activation of this pathway gives rise to basal cell carcinomas (Nature Genet. 24: 216-7, 2000). Applicants should have a solid background in molecular and cell biology, with experience in transgenic animal models desirable but not required. Interested individuals should send a CV, letter of interest, and names of three references to: Dr. Andrzej Dlugosz, University of Michigan, Department of Dermatology and Comprehensive Cancer Center, 3310 CC03, Box 0932, 1500 East Medical Center Drive, Ann Arbor, MI 48109-0932 Email: dlugosza@umich.edu. The University of Michigan is an Equal Opportunity Employer.

And now for the rest of the story.

Therer is no story for this issue. Please look in the September issue for the next installment.

If you wish to know how a particular line of investigation got started, or know of a story that would be interesting to readers of the PASPCR Newsletter, please email me at bill@lenti.med.umn.edu, and I will try to get the rest of the story.
2002 IFPCS Conference

The XVIIIth IPCC will take place at the Hotel Zuiderduin in Egmond aan Zee. Hotel Zuiderduin is situated in the dunes of Egmond aan Zee, one hundred meters from the North Sea. This beautiful, modern hotel is provided with a pub, swimming pool, sauna, beauty centre and fitness centre. Egmond aan Zee is a small fishermen town, with a lot of tourist attractions. Most prominent is the Jan van Speyck lighthouse.

Near Egmond aan Zee, the larger city of Alkmaar is found. Alkmaar is a medieval town which is still visible via the old town centre with a labyrinth of small streets. The most prominent buildings in town are The Great St. Laurens Church and the Weigh house. On Friday morning till half September there is a cheese market (for almost 400 years now) around the Weigh house.

SOCIAL PROGRAM

Welcome Reception

Sunday 8th September 2002, 17:00 - 19:00
Delegates and their accompanying persons are invited to attend this reception, which will take place in Hotel Zuiderduin

Boat Trip and Banquet in Grand Hotel Krasnapolsky, Amsterdam

Wednesday 11th September 2002, 16:30 - 23:30
The participants will have the opportunity to take part in an excursion to Amsterdam, where they will be taken on a boat tour through the canals. The Conference Banquet will take place at the Golden Tulip Grand Hotel Krasnapolsky.

The Golden Tulip Grand Hotel Krasnapolsky is situated directly opposite the Royal Palace on the Dam right in the heart of Amsterdam. The breathtaking Winter Garden, the glass palace dating from 1879, where the dinner will take place, is listed as National Monument.

Please note that the capacity in the Winter Garden is limited; tickets will be available on a first come, first served basis. (Dress code: informal)

Farewell Party

Thursday 12th September 2002, 19:00 - 23:00
Delegates and their accompanying persons are invited to attend the farewell party, which will take place in Hotel Zuiderduin.

Active Jazz players: a dixieland band from the medical faculty from Praque will entertain us during the farewell party and a jam-session will be organized. Take your instrument with you !!!

The following is the scientific program for The XVIIIth International Pigment Cell Conference, to be held in The Hague, Holland. For the most current information, go to the IPCC web site at http://users.raketnet.nl/ipcc/

SCIENTIFIC PROGRAM

Sunday 8th September 2002

Council Meetings
Building up exhibitions
Poster mounting
14:30 - 17:00 Registration
17:00 - 19:00 Welcome Reception

Monday 9th September 2002

Morning:
Welcome and Opening Remarks
Special Lecture: Videomicroscopy in Pigment Cell Research (P. Friedl, Germany)
Plenary Symposium: Biochemistry and Molecular Biology of Melanogenesis
Special Lecture: What is wrong in xeroderma pigmentosum (J.H.J. Hoeijmakers, The Netherlands)
Plenary Symposium: Photochemistry and Skin Pigmentation
Lunch and poster viewing

Afternoon:
Plenary Symposium: Epidemiology of Melanoma and Precursor Lesions
Press Conference: Nevi and UV radiation as risk factors for melanoma

Evening:
Concurrent workshops
Workshop I: Immunobiology of Melanocytes and Melanoma.
Workshop II: Melanocyte Microscopy; Imaging and Ultrastructural Studies
Dinner at Hotel Zuiderduin
Tuesday 10th September 2002

Morning:
Plenary Symposium IV: Genetics of Pigmentation
Special Lecture: Molecular genetics of pigmented disorders (R.A. Spritz, U.S.A.)
Seiji Memorial Lecture
Lunch and Poster viewing / Women forum

Afternoon:
Plenary Symposium: Developmental Biology of Pigmentary System
Concurrent Workshops
Workshop I: Chemistry of Eumelanin and Pheomelanin
Workshop II: Comparative Biology; Other Pigmentary Systems
Dinner at Hotel Zuiderduin

Wednesday 11th September 2002

Morning:
Presidential Address: Melanin and melanogenesis: a fascinating target for chemists (S. Ito, Japan)
Plenary Symposium: Cutaneous and Ocular Melanoma: Genetics and Basic Biology
Concurrent workshops
Workshop I: Extracutaneous Pigmentation
Workshop II: (Patho)physiological Aspects of Melanosomes
Lunch and poster viewing

Afternoon:
Special lecture: The role of melanocortin-1-receptor polymorphism in skin cancer risk (R.A. Sturm, Australia).
Plenary Symposium: Cell Biology and Cell Signalling in Melanocytes
Excursion to Amsterdam, Boat Trip and Banquet, 16:30 - 23:30
Banquet at Hotel Krasnapolsky, Amsterdam

Thursday 12th September 2002

Morning:
Plenary Symposium: Cutaneous and Ocular Melanoma: Diagnosis and Treatments
Plenary Symposium: Vitiligo and Other Pigmentary Disturbances
Lunch and poster viewing

Afternoon:
Plenary Symposium: Hot Topics
Special lecture: Identification of Melanocyte and Melanoma Biomarkers using DermArray DNA Microarrays (T.P. Dooley, U.S.A.)
Plenary Symposium: Genomics and Proteomics in Pigment Cell Research
Closing remarks
19:00-23:00 Farewell Party (Hotel Zuiderduin)

Note:
The following persons have been invited to present their recent work in the above mentioned plenary symposia:
Z. Abdel-Malek (U.S.A.); R. Balloti (France); J. Bolognia (U.S.A.); L. Brochez (Belgium); A.M. Eggermont (The Netherlands); Garcia-Borron (Spain); V.J. Hearing (U.S.A.); M. Herlyn (U.S.A.); Imokawa, G. (Japan), R. Mackie (U.K.); S. Shibahara (Japan); K. Toyofuku (Japan).

Friday 13th September 2002

SATellite Symposia

I. The Congenital Giant Naevus: Research and Treatment
Organiser: H. Etchevers (France).
The symposium may be of particular interest for dermatologists, plastic and reconstructive surgeons and scientists involved in the molecular and cell biology of congenital (giant) naevi. Sessions of grouped short presentations will be followed by round table exchanges with the audience. Presentation abstracts and lunch will be provided. The following topics are anticipated:
dermabrasion, dermal and epidermal substitutes, epidemiology, fetal naevogenesis, malignant transformation, neurocutaneous melanosis, phototherapy, scarring in and near naevi, satellite naevi, skin mobilisation, targeted immunotherapy, the naevocyte in vitro, tissue expanders. The first Annual Meeting of Naevus 2000 France-Europe’s Committee on Medicine and Science will follow the symposium.

For more information regarding the scientific program, please contact:
IPCC Naevus Symposium
c/o Dr. Heather Etchevers
Institut d’Embryologie Cellulaire et Moléculaire
49bis avenue de la Belle Gabrielle
F-94736 Nogent-sur-Marne Cedex, France
Phone: +33-1 45 14 15 11, Fax: +33-1 48 73 43 77
E-mail: etchever@infobiogen.fr

II. Melanoma Biology

Organisers: D. J. Ruiter (The Netherlands), G.N.P. van Muijen (The Netherlands), M. Herlyn (U.S.A.)

The aim of the satellite symposium is to discuss new concepts in melanoma biology, new techniques that enable further observation in animal and human tumors, new approaches for diagnostic and prognostic purposes, and new strategies for immunologic therapy. The symposium will consist of invited lectures that will be divided in four successive sessions. Presentations will be given by leading experts in the field. The meeting will take place from 9:00 until 17:30 and will be followed by cocktails and dinner. Coffee / tea and lunch will also be provided.

A tentative program with the speakers who have been invited is given below:

New Concepts:

Stem cells and stromagenesis; (M. Herlyn; Philadelphia)

Lymphangiogenesis (R. de Waal; Nijmegen)

Dynamics of tumor invasion (P. Friedl, Wuerzburg)

New Technologies:

Transgenic mouse model (P. Krimpenfort; Amsterdam)

Spectral imaging of transcription factor expression (D. Becker; Pittsburg)

Tetramerestaining of cytotoxic lymphocytes (M. Bernsen; Nijmegen)

New Approaches in Diagnosis and Prognosis:

Comparative genomic hybridization-based microarrays (B. Bastian; San Francisco)

Cytokine profiles (D. Schadendorf; Mannheim)

Uveal melanoma as a paradigm (M. Jager; Leiden)

New Strategies for Immunologic Therapy:

Melanosomal proteins (V. Hearing; Bethesda)

Dendritic cells (G. Adema; Nijmegen)

New targets (M. Bar-Eli; Houston)

For more information regarding the scientific program, please contact

Prof. Dr. D. J. Ruiter
Department of Pathology
University Medical Center St Radboud
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NL-6500 HB Nijmegen
The Netherlands
Phone: +31-24-3614825; Fax:+31-24-3540520
E-Mail: d.ruiter@pathol.azn.nl

III. Pigment Cell Societies Development Group

Organiser: Dr. W. Pavan

The International Federation of Pigment Cell Societies Development Group is hosting a one-day satellite meeting designed to coalesce the remarkable recent progress in the field of pigment cell development with research aimed at gaining new insights into differentiation, gene expression and embryology. This meeting will explore these themes by gathering experts in a broad range of fields spanning basic transcriptional mechanisms through the clinical impact of pigment cell research. The program will include talks from invited group members and talks selected from abstract submissions. Our goal is to bring together a diverse group of dynamic speakers and participants who can together explore the current findings in pigment cell development.
IV. Hyperpigmentation and hypopigmenting agents

Co-chair James J. Nordlund (PASPCR), Jean-Paul Ortonne (ESPCR) and Masako Mizoguchi (JSPCR)

1. Introduction: The problem of hyperpigmentation - J Nordlund (USA)

2. Melasma and hyperpigmentation: histological studies - Won Hyoung Kang, Ajou (Korea)

3. New studies on the causes of melasma; role of contact and photodermatitis in melasma - Verena Verallo-Rowell Manila, Philippines

4. Dermal melanosis and melanocytosis: current therapies in the Far East - Masako Mizoguchi, Japan

5. Review of therapies in Europe and results with lightening agents for treatment of pigmentary problems - Wiete Westerhof (The Netherlands)

6. Review of new therapies in US and work on anisole and combination therapies - Jean Bolognia (USA)

Coffee break

7. Review of treatment of hyperpigmentation in South America - Doris Hexsel

8. Review of arbutin and derivatives - Ray Boissy (USA)

9. Theoretical Approaches to hypopigmenting skin - a basis in cell biology - Matsumitsu Ichihashi (Japan)

10. Theoretical Approaches to hyperpigmenting skin - a basis in cell biology - Jean Paul Ortonne (France)

11. Metabolism of melanosomes: Is melanin biodegradable? - Jan Borovansky (Czech Republic)

12. Open Discussion
This is a tentative agenda for the first Annual Melanoma Research Congress. It does not contain names but the workshops etc. Please contact Meenhard or DuPont Guerry for more information.

First Annual Melanoma Research Congress

June 21-24, 2003

Wyndham Franklin Plaza Hotel
17th & Race Streets
Philadelphia, PA 19103
(Phone: 215-448-2852; Fax: 215-448-2853)

Sponsored by the Foundation for Melanoma Research and Other Foundations

Conference Co-Chairs
Meenhard Herlyn DuPont Guerry
The Wistar Institute University of Pennsylvania
Philadelphia, PA Philadelphia, PA

Saturday, June 21

3:00 pm to 10:00 pm REGISTRATION

7:00 pm to 9:00 pm Keynote Addresses

Steven Rosenberg
Rational Immunotherapy of Melanoma

Jeff Trent
Genomics approaches to Melanoma Biology

9:00 pm to 10:00 pm OPENING RECEPTION

Sunday, June 22

8:00 am to 10:30 am Plenary Session 1

Immunobiology and Immunotherapy of Melanoma
Giorgio Parmiani, Chair

Drew Pardoll
Thierry Boon
Soldano Ferrone
Ralph Reisfeld

11:00 am to 12:30 pm Workshops 1-3

Workshop 1
Experimental Melanocyte Transformation
Scott Lowe
Carola Berking
Marisol Soengas
Glenn Merlino

Workshop 2
Genetics of Familial Melanoma
Alisa Goldstein
David Hoog
Nick Hayward
Julia Newton Bishop

Workshop 3
Melanocyte Development and Melanoma
David Fisher
Heinz Arneheiter
Bill Pavan
Nishimura?

Saturday, June 21

3:00 pm to 10:00 pm REGISTRATION

7:00 pm to 9:00 pm Keynote Addresses

Steven Rosenberg
Rational Immunotherapy of Melanoma

Jeff Trent
Genomics approaches to Melanoma Biology

9:00 pm to 10:00 pm OPENING RECEPTION

Sunday, June 22

8:00 am to 10:30 am Plenary Session 1

Immunobiology and Immunotherapy of Melanoma
Giorgio Parmiani, Chair

Drew Pardoll
Thierry Boon
Soldano Ferrone
Ralph Reisfeld

11:00 am to 12:30 pm Workshops 1-3

Workshop 1
Experimental Melanocyte Transformation
Scott Lowe
Carola Berking
Marisol Soengas
Glenn Merlino

Workshop 2
Genetics of Familial Melanoma
Alisa Goldstein
David Hoog
Nick Hayward
Julia Newton Bishop

Workshop 3
Melanocyte Development and Melanoma
David Fisher
Heinz Arneheiter
Bill Pavan
Nishimura?

12:30 pm to 2:00 pm Break

2:00 pm to 4:00 pm Plenary Session 2

Molecular Epidemiology of Melanoma
Marianne Berwick, Chair
Margarete Tucker
Tim Rebbeck
Mann/Kefferd/
Landis/Bergman

4:00 pm to 4:30 pm Break

4:30 pm to 6:00 pm Workshops 4-6

Workshop 4
Immunobiology II
Susan Topalian
Franco Marincola
Ingegerd Hellstrom
Walter Storkus

Workshop 5
UV and Melanoma
Glenn Merlino
Vince Hearing
Rick Sturm
Frances Noonan
Ed DeFabo
Workshop 6  
Molecular and Functional Genomics of Melanoma  
David Speicher  
Katheryn Resing  
Boris Bastian  
8:00 pm to 10:00 pm  
Poster Session 1  
Immunobiology and Immunotherapy of Melanoma

Monday, June 23  
8:00 am to 10:00 am  
Plenary Session 3  
Biology of Melanoma  
Dorothea Becker, Chair  
Meenhard Herlyn  
Ze’ev Ronai  
Ruth Halaban  
Lynda Chin  
10:00 am to 10:30 am  
Break

11:00 am to 12:30 pm  
Workshops 7-9  
Biomarkers  
David Hoon  
Anja Bosserhoff  
Dirk Ruiter  
DuPont Guerry  
Workshop 8  
Tumor Progression  
David Elder  
Martin Mym  
Alistar Cochran  
Workshop 9  
Bioinformatics for Genomics and Proteonomics  
Todd Golub  
Towia Liebermann  
Phyllis Gimotty  
Louise Showe  
12:30 pm to 2:00 pm  
Break

2:00 pm to 4:00 pm  
Plenary Session 5  
Invasion and Metastasis of Melanoma  
I. Fidler  
D. Ruiter  
R. Muschel  
D. Welsh  
4:00 pm to 4:30 pm  
Break

4:30 pm to 6:30 pm  
Poster Session B  
Etiology, Biology, Prevention, and Novel Therapy of Melanoma  
Advocates and Melanoma Research  
DuPont Guerry  
8:00 pm to 10:00 pm  
Reception/Dinner  
Inauguration of the Society for Melanoma Research  
Advocate Speaker

Tuesday, June 24  
8:00 am to 10:00 am  
Plenary Session 6  
Novel Concepts for Melanoma Etiology, Prevention, Diagnosis, and Prognosis  
Meenhard Herlyn  
Robert Benezra/Raffi  
Mary Hendrix  
Catherine Verfaille  
10:00 am to 10:30 am  
Break

10:30 am to 12:30 am  
Plenary Session 7  
Hot Topics in Non-immunological Therapy of Melanoma  
Bob Wittes  
Disease Gene Identification  
Genomics of Drug Discovery
Bibliography:

The Bibliography published in this issue covers the period March, 2002 through May, 2002. If you notice a paper that was not detected by this search that should be included, please send it to us and we will include it in the next issue. By its very nature, assignment of a reference to a particular category is arbitrary and we urge you to read through all categories to make sure you don’t miss any pertinent to your field.

MELANINS, MELANOGENS & MELANOGENESIS


MELANOSOMES, MELANOCYTES & KERATINOCYTES


Borodoker N, Cunningham ET, Yannuzzi LA, Nicoletti R: Peripheral curvilinear pigmentary streak in multifocal choroiditis. ARCH OPHTHALMOL 120:520-521 (2002).


Cleaver JE, Crowley E. UV damage, DNA repair and skin carcinogenesis. FRONT BIOSCI 7, D1024-D1043. 2002.


Donatien P, Jeffery G: Correlation between rod photoreceptor numbers and levels of ocular pigmentation. INVEST OPHTHALMOL VISUAL SCI 43:1198-1203 (2002).


Goud B. How Rab proteins link motors to membranes. NAT CELL BIOL 4[4], E77-E78. 2002.


Nagashima K, Torii S, Yi ZH, Igarashi M, Okamoto K, Takeuchi T et al. Melanophilin directly links Rab27a and myosin Va through its distinct coiled-coil regions. FEBS LETT 517[1-3], 233-238. 2002.


MELANOMA & METASTASIS

Time for a campaign on melanoma. EUR J CANCER 38:740 (2002).


Ge XK, Fu YM, Meadows GG: U0126, a mitogen-activated protein kinase kinase inhibitor, inhibits the invasion of human A375 melanoma cells. CANCER LETT 179:133-140 (2002).


Wilson D, Klein ML: Cryotherapy as a primary treatment for choroidal melanoma. ARCH OPHTHALMOL 120:400-403 (2002).


**MSH, POMC, GROWTH FACTORS & RECEPTORS**


Quto SS, Ng CE: Comparison of apoptotic, necrotic and clonogenic cell death and inhibition of cell growth following camptothecin and X-radiation treatment in a human melanoma and a human fibroblast cell line. CANCER CHEMOTHER PHARMACOL 49:167-175 (2002).

Retsas S, Henry K, Mohammed MQ, MacRae K: Prognostic factors of cutaneous melanoma and a new staging system proposed by the American Joint Committee on Cancer (AJCC): validation in a cohort of 1284 patients. EUR J CANCER 38:511-516 (2002).


Smalley KSM, Eisen TG: Farnesyl thiosalicylic acid inhibits the growth of melanoma cells through a combination of cytostatic and pro-apoptotic effects. INT J CANCER 98:514-522 (2002).


DEVELOPMENTAL BIOLOGY


Terry RB, Bailey E, Lear T, Cothran EG: Rejection of MITF and MGF as the genes responsible for appaloosa coat colour patterns in horses. ANIM GENET 33:82-84 (2002).


DIFFERENTIATION


Steingrimsson E, Tessarollo L, Pathak B, Hou L, Arnheiter H, Copeland NG, Jenkins NA: Mitf and Tfe3, two members of the Mitf-Tfe family of bHLH-Zip transcription factors, have important but functionally redundant roles in osteoclast development. PROC NAT ACAD SCI USA 99:4477-4482 (2002).


MISCELLANEOUS


