METHODS IN CLINICAL CANCER RESEARCH

The 14th intensive Workshop for junior clinical oncologists in any clinical research specialty area, to learn the essentials of clinical trial design

Waldhaus Flims, Switzerland

Applications Open: 14 December 2011
Applications Close: 13 February 2012

www.ecco-org.eu
WORKSHOP DIRECTORS

Representing ECCO
Patrick Schöffski   University Hospital Gasthuisberg, Leuven, Belgium

Representing AACR
Stephen M. Hahn   University Hospital of Pennsylvania, Philadelphia, USA

Representing EORTC
Jan Bogaerts   EORTC Headquarters, Brussels, Belgium

Representing ESMO
Johann de Bono   Royal Marsden Hospital, Sutton, United Kingdom

Workshop Faculty

Representing ECCO
Peter Naredi   Riccardo Riccardi

Representing AACR
Patricia M. LoRusso   Ignacio Wistuba

Representing EORTC
Sandrine Marreaud   Stefan Sleijfer

Representing ESMO
Christoph Dittrich   Chris H. Takimoto

Additional Faculty
Powel H. Brown   Susan Percy Ivy   Yolande Lievens   Laura Luisa   David MacPhin   Xavier Paletti   Lesley K. Seymour   Sally Steinig   Stefan Michals   Piotr Rutkowski

Additional Faculty will be announced. Faculty disclosures will be provided in the Workshop Syllabus

I have worked over the years with fellows and junior faculty members who have attended the Flims course and have found them to be exceptionally well-prepared to perform high quality clinical research.

I have attended Flims in 1999 and it changed my career and my understanding of cancer research.

The most stimulating, interactive and multidisciplinary educational activity on clinical trial methodology in oncology. Outstanding, top-ranking international faculty. Highly motivated, selected workshop participants with very heterogeneous clinical, scientific and cultural background. A unique, once-in-a-lifetime opportunity for young clinical cancer researchers.

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KEY BENEFITS OF ATTENDING THE WORKSHOP

- Exclusive access to and mentoring by up to 40 highly experienced clinical experts in the field of oncology from Europe and the U.S.A;
- Exceptional opportunity to meet and network with an elite group of 80 junior clinical oncologists from all over the world;
- Outstanding educational experience in a unique setting conducive to professional relationship building, learning and development;
- Access to a variety of educational tools designed to enable participants to develop their initial protocol proposal into a complete protocol;
- Active promotion of productive dialogues between young cancer specialists and the European and non-European Cancer Societies;
- Establishment of a network for educational exchanges between young cancer clinicians worldwide.

WHY DO WE NEED A WORKSHOP?

The presence of a strong research base is essential to the future of good quality cancer care. Clinical scientists who are able to set up and run high-quality clinical trials are vital to the advancement of new therapies. This Workshop was established to reverse the decline in numbers of clinical scientists. The ultimate goal: to develop a robust, expanding base of well-trained clinical researchers by providing them with the training they need to conduct better clinical/translational trial designs.

ワークショップの目的

良好な癌治療の将来に必要な強い研究の基盤は重要です。高品質の臨床試験を設立および実施する能力を持つ臨床科学者たちは新薬の進展に非常に重要です。このワークショップは、臨床科学者の数の減少を逆転することを目的に設けられました。最終的な目標：訓練を受けた臨床研究者を育てることで、より良い臨床/臨床トランスレーション試験を行う能力を培います。
The workshop starts on Saturday 23 June 2012 so participants are asked to arrive in Flims no later than 04.00pm as the workshop will start with a mandatory pre-test at 04.00 p.m.

Saturday 23 June 2012

12:00 – 16:00  Registration
16:00 – 17:00  Pre-test (mandatory for all fellows)
17:00 – 17:35  Welcome and Workshop overview
17:35 – 18:15  Introductory lecture: Questions to ask yourself in designing a clinical trial
18:30 – 20:30  Protocol development session 1

Protocol Development Sessions
These sessions form the core activity of this Workshop and allow students to complete the writing of their protocol by applying the knowledge acquired during the Workshop. Students will receive detailed critiques of their proposals from the experts in the field in small groups comprising a maximum of 10 people.

Meet Your Expert Sessions
One-to-one sessions where students will have access to experts providing individual counselling and advice on protocol and career development.

Small Group Discussion Sessions
These sessions focus on topics that are essential to the success of clinical trials as well facilitating discussion on and around the difficulties and challenges of a particular type of trial. They are limited in size to maximise exchange of information.

Lectures and Panel Discussions
Presentations by experts in the field on specific topics will provide participants with an overview of the design and conduct of high-quality clinical trials. This will be followed by a panel discussion or round table session during which Faculty and students can explore issues raised during the talks in greater depth.

The Scientific Sessions have been specially formulated to cater for all learning needs and will use one of the following four formats:

Sunday 24 June 2012

08:30 – 10:00  Lecture session 1
- Basic biostatistics for the clinical trialist I
- Basic biostatistics for the clinical trialist II
- Common errors in statistics
10:15 – 11:30  Small Discussion Groups - session 1
11:30 – 12:30  Small Discussion Groups - session 2
13:30 – 15:30  Protocol development session 2
16:00 – 17:00  Small Discussion Groups - session 3
17:15 – 18:15  Small Discussion Groups - session 4
18:15 – 19:30  Individual work on protocols
20:45  Individual work on protocols
20:45 – 22:45  Meet-your-expert session (office hours)
Tuesday 26 June 2012

08:00 – 09:30 Individual work on protocols
09:30 – 11:00 Protocol development session 4
11:00 – 12:30 Lecture session 5
   • Role of pharmacokinetics in clinical trials
   • What can imaging contribute to your trial?
   • CRM & Bayesian designs – practical aspects of implementation

13:30 – 14:30 Lecture session 6
   • Ethical principles in the conduct of clinical trials
   • Patient-oriented endpoints / Quality of Life
   • Round table – Ethical issues and informed consent - a case-based discussion

14:45 – 16:45 Meet-your-expert session in parallel with Small Discussion Group sessions and individual work on protocols

14:45 – 15:45 Small Discussion Groups - session 5
15:45 – 16:45 Small Discussion Groups - session 6
17:00 – 19:00 Individual work on protocols
20:30 – 21:30 Meet-your-expert session (continued)

Wednesday 27 June 2012

08:00 – 09:00 Individual work on protocols
09:00 – 10:00 Small Discussion Groups - session 7
10:00 – 11:00 Lecture session 7
   • Reading the literature with a critical eye
   • Data and safety monitoring and independent study review

11:15 – 12:30 Lecture session 8
   • Barriers to successful implementation
   • Regulatory and other practical issues
   • Improving patient participation in cancer clinical trials
   • Panel discussion

13:30 – 15:30 Protocol development session 5
15:45 – 19:00 Individual work on protocols
20:30 – 21:30 Individual work on protocols

Thursday 28 June 2012

08:30 – 10:00 Lecture session 9
   • Molecular targeting and the prevention-therapy convergence
   • Translating cancer research into targeted therapeutics
   • Haematology as a role model for drug development

10:15 – 11:00 Small Discussion Groups - session 8
11:15 – 13:15 Protocol development session 6
13:30 – 17:00 Individual work on protocols
14:15 – 16:00 Break (mandatory for all fellows)
16:00 – 17:00 Panel discussion
17:00 – 19:00 Group activity
19:00 Final protocol due
20:00 Reception and dinner

Friday 29 June 2012

Departure

Although I’m not a radiation oncologist, I enjoyed this lecture. I learnt about the difference between radiation oncology study and medical oncology study.

Ogita Shin

Monday 25 June 2012

08:15 – 09:15 Lecture session 2
   • Special considerations in trials of radiation therapy – implications for design, endpoints and quality control
   • Special considerations in combined treatment trials (Chemo-radiation) – implications for design, endpoints and quality control

09:15 – 10:15 Lecture session 3
   • Integrating surgery in multi-modality trials – implications for design, endpoints and quality control
   • Design of studies with immunological agents

10:30 – 11:30 Lecture session 4
   • Prognostic and predictive markers for patient selection
   • Individual work on protocols

14:45 – 16:45 Meet-your-expert session in parallel with Small Discussion Group sessions and individual work on protocols

14:45 – 15:45 Small Discussion Groups - session 5
15:45 – 16:45 Small Discussion Groups - session 6
17:00 – 19:00 Individual work on protocols
20:30 – 21:30 Meet-your-expert session (continued)
ONLINE APPLICATION PROCEDURE

Applications to participate in the ECCO-AACR-EORTC-ESMO Workshop on Methods in Clinical Cancer Research can only be submitted electronically. Paper submissions will NOT be accepted.

For the online application please go to Workshop website at: www.ecco-org.eu (select ‘Education>Flims>Flims 14’) and follow the instructions on the screen.

Deadline for receipt of applications: Monday 13 February 2012.

MINIMUM SELECTION CRITERIA

1. Candidates must be in at least the 2nd year of training and within 5 years of completion of Residency/Fellowship training in one of the following disciplines:
   - Junior physician specialising in oncology;
   - Junior clinical professional managing cancer patients (i.e. urologist, gynaecologist, neuro-oncologist, haematologist);
   - Junior radiologist or pathologist with a strong involvement in cancer care.

2. Have a major interest in clinical research and intend to develop a career in that field.

3. Aim to write and conduct a clinical protocol for a study not previously performed, nor written, which is also considered feasible within the institutional setting and the time of completion of the candidate’s clinical training.

4. Be fluent in written and spoken English and have good computer skills.

5. Have support from the Supervisor/Head of Department and sustained commitment in the years following the Workshop.

GENERAL INFORMATION & CONDITIONS OF PARTICIPATION

Selection of Participants

Participation to the Workshop is limited to 80 participants. The Workshop Review Committee will evaluate the applications and base its decision on a number of factors including:

- Quality and feasibility of the proposed protocol concept and the letters of commitment submitted;
- Individual career path in medical training and competence in clinical cancer research;
- Support of relevant departments and/or institutions to help conduct the clinical trial.

The Workshop Review Committee’s decision is final and whilst we welcome your feedback about the application process, the Workshop Review Committee will not enter into any discussions regarding the final decision.

For further details on application requirements, the selection criteria and process, please visit www.ecco-org.eu

Workshop Materials

As of May 2012, selected participants will have access to the Flims Intranet, an online resource platform for all educational Workshop material. The Flims Intranet will also be used as a message centre and as a platform for all organisational aspects of the Workshop.

Participation Fee

In order to attend the Workshop, all selected participants will be required to pay the Workshop Participation Fee of 2,000 EUR (local VAT incl.).

Applicants from countries with limited resources may apply for an exemption of the Workshop Participation Fee. Each application will be assessed on a case-by-case basis in accordance with the evaluation criteria.

The Workshop Participation Fee offsets only part of the actual Workshop costs per student, which includes the following:

- Round-trip travel arrangements from closest home airport to Zurich or travel reimbursement as specified in the Workshop Reimbursement Policy for trips arranged by the participant;
- Shuttle bus service from Zurich airport to the Workshop Venue on Saturday 23 June 2012;
- Shuttle bus service from the Workshop Venue to Zurich airport on Friday 29 June 2012;
- Accommodation in the Workshop Venue from 23-29 June 2012 (for single room accommodation a supplement applies)
- Food and beverages (5 meals/day) throughout the duration
- Access to Flims Intranet, the online resource platform for all Workshop material.

Please note: This Workshop is supported by generous grants from national, European and international cancer organisations and educational grants from corporate sponsors.

The Flims Alumni Club

http://www.ecco-org.eu/Flims

The Flims Alumni Club (FAC) is a non-profit organisation and an Advisory Member of ECCO, the European CanCer Organisation. It was established in 2001 and is open solely to young professionals and Faculty who have participated in the ECCO-AACR-EORTC-ESMO Workshops on ‘Methods in Clinical Cancer Research’ in Flims, Switzerland.

The FAC responds to the interests and needs of highly driven junior clinical oncologists by offering an expanding range of benefits exclusive to its Members. These include access to the FAC Members Directory and a direct networking opportunity via the Flims Alumni Club LinkedIn Group. Through these channels the FAC aims to develop a sense of community by fostering interactions amongst its members and also promote a productive dialogue between young cancer specialists and the European and non-European Cancer Societies.
Workshop Venue

Waldhaus Flims Hotel
Via dil Parc
7018 Flims
Switzerland
www.waldhaus-flims.ch

Workshop Secretariat

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