



MEN 2010

12th International Workshop
on Multiple Endocrine
Neoplasia

Gubbio, Italy
September 16-18, 2010



CIC Edizioni Internazionali

MEN 2010



12th International Workshop
on Multiple Endocrine
Neoplasia

September 16 • 18 2010
Gubbio • Park Hotel ai Cappuccini

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Piazza Adua, 1/4 - 50123 Firenze
Tel. +39 055 2608941 - Fax +39 055 2608948 - e-mail: info@men2010.com



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Scientific program

September 16

- 15.00-15.30 **Opening Ceremony**
SYMPOSIUM I: Partnering for Rare Hereditary Cancers
Chairs: Jérôme Bertherat and Rajesh Thakker
- 15.30-16.00 Hereditary cancer: commonalities and differences with other rare diseases
Paolo Casali, Italy
- 16.00-16.30 Clinical research, therapy development and regulatory considerations in rare diseases - A View from the Patients
Alastair Kent, United Kingdom
- 16.30-17.00 **PLENARY LECTURE I**
Chair: Paolo Miccoli
"SPECT/CT and PET/CT in the diagnosis and follow-up of pancreatic neuroendocrine tumors"
Drazo Schillaci, Italy
- 17.00-17.30 **PLENARY LECTURE II**
Chair: Robert F. Gagel
Preimplantation genetic diagnosis in hereditary endocrine tumors
Steven A. Lietman, USA
- 17.30-18.00 **PLENARY LECTURE III**
Chair: Diego Ferone
Dopastatin: from basic to clinics
Annamaria Colao, Italy
Presentation of the "Parathyroid Medal" of the F.I.R.M.O. Foundation
- 18.00-18.30 Lecture: Hyperparathyroid genes: sequencing reveals answers and questions
- 18.30 **Welcome Reception**

September 17

- 8.00-8.30 **SYMPOSIUM II: Pituitary in Hereditary Tumor Syndromes**
Chairs: Gaetano Lombardi and Anna Spada
Familial pituitary tumor syndromes
Albert Beckers, Belgium
- 8.30-9.00 AIP-related molecules in pituitary adenomas
Auli Karhu, Finland
- 9.00-9.30 VEGF-A inhibition, a potential novel therapy for pituitary tumors
Napoleone Ferrara, USA
- 9.30-10.00 **Coffee Break and Poster Viewing**
SYMPOSIUM III: Parathyroids in Hereditary Tumor Syndromes
Chairs: Stephen Marx and Samuel A. Wells
- 10.00-10.30 Menin interactions: tumor-suppressive and pro-oncogenic effects
Sumita Agarwal, USA
- 10.30-11.00 The epidemiology and genetics of parathyroid carcinoma
Bin T. Teh, USA
- 11.00-11.30 Interactions of the MEN1 Gene Product with Nuclear Receptors and Histone Methylation Pathways
H. Th. Mark Timmers, The Netherlands
- 11.30-12.00 Regulation of Cell Proliferation by Menin
Xianxin Hua, Usa
- 12.00-12.30 Surgery of hereditary hyperparathyroidism
Bruno Niederle, Austria

- 12.30-13.00 **PLENARY LECTURE IV**
Chair: Silvano Adami
Medical management of primary hyperparathyroidism
Aliya Khan, Canada
- 13.00-14.00 **Lunch Break and Poster Viewing**
PLENARY LECTURE V
Chair: Gennaro Favia
Cinacalcet in the treatment of primary hyperparathyroidism
Annamaria Colao, Italy
- 14.00-14.30 **PLENARY LECTURE VI**
Chair: Laura Masi
Parathyroid hormone substitution therapy in hyperparathyroidism
Lars Reynmark, Denmark
- 14.30-15.00 **ORAL PRESENTATIONS I**
Chairs: Diego Ferone and Barbara Pasini
- 15.00-16.30 **SYMPOSIUM IV: Neuroendocrine Tumors in Hereditary Tumor Syndromes**
Chairs: Annamaria Colao and Bertram Wiedenmann
Carcinoids: eticopathogenesis and reflections on therapy
Guido Rindi, Italy
- 16.30-17.00 Von Hippel-Lindau-associated pancreatic manifestations
Alberto Falchetti, Italy
- 17.00-17.30 Surgical approaches for duodenal-pancreatic tumors in MEN 1
Francesco Tonelli, Italy
- 17.30-18.00 **Coffee Break and Poster Viewing**
SYMPOSIUM V: Somatostatin Analogs
Chairs: Alberto Falchetti and Piero Ferolla
- 18.00-18.30 Somatostatin analogs in gastro-entero-pancreatic tumors
Kjell Öberg, Sweden
- 18.30-19.00 PRRT with Lu-177/IDOTA, Tyr310Octreotate
Dirk Jan Kwekkeboom, The Netherlands
- 19.00-19.30 **PLENARY LECTURE VII**
Chair: Bruno Niederle
- 19.30-20.00 Novel pharmacotherapies of neuroendocrine tumors
Bertram Wiedenmann, Germany
- 20.30-21.30 **THE MEN ADVOCACY GROUP**
Sponsored by AIMEN 1 & 2
Associazione Italiana Neoplasia Endocrina Multiple 1 & 2
Chairmen: Maria Luisa Brandi and C.J.M. Lips
 - Welcome of the President of AIMEN 1 & 2
 - The role of MEN Associations
 - In Italy - In Belgium - In Holland
 - In England - In USA
- 21.30 The International Federation of MEN Associations (FederMEN)
Questions and answers
Cocktail

September 18

- 8.00-8.30 **SYMPOSIUM VI: Thyroid in Hereditary Tumor Syndromes**
Chairs: Friedhelm Raue and Furio Pacini
Novel signaling mechanisms of oncogenic RET kinases
Massimo Santoro, Italy
- 8.30-9.00 Targeting Mutant RET
Clinical Advances in the Use

- 9.00-9.30 of Tyrosine Kinase Inhibitors in Multiple Endocrine Neoplasia Type II and Sporadic Medullary Thyroid Carcinoma
Robert Gagel, USA
- 9.30-10.00 Surgical approaches to hereditary medullary thyroid cancer
Henning Dralle, Germany
Coffee Break and Poster Viewing
- 10.00-10.30 **SYMPOSIUM VII: Adrenal Glands in Hereditary Tumor Syndromes**
Chairs: Hartmut P.H. Neumann and Bruce G. Robinson
Functional consequences of SDH mutations
Giuseppe Opocher, Italy
- 10.30-11.00 MENX - Associated rat pheochromocytomas
Natalia S. Pellegata, Germany
- 11.00-11.30 Mice models of adrenal PRAK1A inactivation
Antoine Martinez, France
- 11.30-12.00 The transcriptome that mediates the PRAK1A effects in Carney complex and other settings: the Wnt signaling pathway
Constantine Stratakis, USA
- 12.00-12.30 Strategy to identify the genetic basis of aldosterone producing adrenal tumors
Felix Beuschlein, Germany
- 12.30-13.00 **PLENARY LECTURE VIII**
Chair: Claudio Marcolucci
Role of MEN1 mutational analysis: resolving challenges due to phenocopies and identifying children with hereditary endocrine tumors
Rajesh Thakker, United Kingdom
- 13.00-14.30 **Lunch Break and Poster Viewing**
PLENARY LECTURE IX
Chair: Francesco Tonelli
Thoracic Neuroendocrine Tumours in MEN 1
Piero Ferolla, Italy
- 14.30-15.00 **ORAL PRESENTATIONS II**
Chairs: Rocco Bellantone and Rossella Elisei
- 15.00-16.30 **SYMPOSIUM VIII: Non Classical Targets of the Hereditary Endocrine Tumor Syndromes**
Chairs: Henning Dralle and Gabriella Nesi
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Andrea Saggini, Italy
- 16.30-17.00 Interactions of hereditary endocrine tumors' genes with bone growth and metabolism
Maria Luisa Brandi, Italy
- 17.00-17.30 Reproductive disorders in hereditary endocrine tumors
Felice Petraglia, Italy
- 17.30-18.00 **Coffee Break and Poster Viewing**
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Chair: Paolo Beck-Peccoz
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Matthias Schott, Germany
- 18.30-19.00 **Farewell Dinner**
- 21.00

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