Sex Hormones and Antihormones in Endocrine Dependent Pathology: Basic and Clinical Aspects

Proceedings of an International Symposium, Milano, 10–14 April 1994

Editors:

Marcella Motta

Department of Endocrinology University of Milan Milan, Italy

Mario Serio

Endocrinology Unit Department of Clinical Physiopathology University of Florence Florence, Italy



1994

Elsevier

 $Amsterdam-Lausanne-New\ York-Oxford-Shannon-Tokyo$



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International Congress Series No. 1064 ISBN 0-444-81879-0

This book is printed on acid-free paper.

Published by: Elsevier Science B.V. P.O. Box 211 1000 AE Amsterdam The Netherlands

Library of Congress Cataloging in Publication Data:

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Sex hormones and antihormones in endocine dependent pathology
 proceedings of an international symposium, Milano, 10-14 April 1994
 / editors, Marcella Motta, Mario Serio.
     p. cm. -- (International congress series; no. 1064)
   Includes bibliographical references and indexes.
   ISBN 0-444-81879-0 (alk. paper)
   1. Breast--Cancer--Endocrine aspects--Congresses. 2. Prostate-
 -Cancer--Endocrine aspects--Congresses. 3. Hormones, Sex-
 -Antagonists--Therapeutic use--Congresses. 4. Hormones, Sex-
 -Therapeutic use--Congresses. 5. Uterus--Diseases--Endocrine
 aspects--Congresses. I. Motta, Marcella. II. Serio, Mario.
 III. Series.
   [DNLM: 1. Neoplasms, Hormone-Dependent--congresses. 2. Sex
 Hormones--congresses. 3. Prostatic Neoplasms--congresses.
 4. Breast Neoplasms--congresses. 5. Genital Diseases, Female-
 -congresses.
              W3 EX89 no. 1064 1994 / QZ 200 S518 1994]
 RC280.B8S48 1994
 616.99'449--dc20
DNLM/DLC
 for Library of Congress
                                                            94-32151
                                                                CIP
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In order to ensure rapid publication this volume was prepared using a method of electronic text processing known as Optical Character Recognition (OCR). Scientific accuracy and consistency of style were handled by the author. Time did not allow for the usual extensive editing process of the Publisher.

Printed in the Netherlands



©1994 Elsevier Science B.V. All rights reserved. Sex hormones and antihormones in endocrine dependent pathology: basic and clinical aspects M. Motta and M. Serio, editors,

Pure antioestrogens in breast cancer: experimental and clinical observations

R.I. Nicholson¹, J.M.W. Gee¹, C.L. Eaton¹, D.L. Manning¹, R.E. Mansel², A.K. Sharma², A. Douglas-Jones², M. Price-Thomas³, A. Howell⁴, D.J. DeFriend⁴, N.J. Bundred⁴, E. Anderson⁴, J.F.R. Robertson⁵, R.W. Blamey⁵, M. Dowsett⁶, M. Baum⁶, P. Walton⁷ and A.E. Wakeling⁷

¹Tenovus Cancer Research Centre, Cardiff, UK; ²Departments of Surgery (REM, AS) and Pathology (AD-J), UWCM, Cardiff, UK; ³Royal Gwent Hospital, Newport, UK; ⁴Withington and Christie Hospitals, Manchester, UK; ⁵Department of Surgery, City Hospital, Nottingham, UK; ⁶Royal Marsden Hospital, London, UK; and ⁷Zeneca Pharmaceuticals, Macclesfield, UK

Key words: ICI 182,780, ICI 164,384, oestrogen receptor, oestrogens, tamoxifen

Introduction

The last 10 years have seen the emergence of a new class of pharmacological agents termed pure antioestrogens (reviewed in [1]). These compounds, which were originally developed by ICI Pharmaceuticals Division in the UK, have the unique property of binding to the oestrogen receptor [2] and producing a receptor complex which lacks oestrogenic activity [3,4]. If we assume that the action of oestrogens on sensitive breast cancers favours cell proliferation and survival, and that they thereby act as a driving force for the growth and development of the disease [5], pure antioestrogens have the potential to fully negate these activities by producing a state of complete oestrogen withdrawal [6].

The perceived importance of pure antioestrogens, therefore, is as alternatives to antihormonal treatments which are designed to reduce the synthesis of oestrogens, but which currently fail to nullify oestrogenic signals arising from other sources [7,8], and as potential successors to "tamoxifen-like" antioestrogens, which although widely and successfully used in the therapy of primary and advanced disease [6], possess partial oestrogenic activity [9] which may negate aspects of their effectiveness as antitumour agents.

Since pure antioestrogens are now entering clinical development, the current paper seeks to outline some of their basic cellular and antitumour properties on human

Address for correspondence: R.I. Nicholson, Tenovus Cancer Research Centre, University of Wales, College of Medicine, Heath Park, Cardiff CF4 AXX, UK.



breast cancer cells in vitro [1,10] primarily using the lead compound ICI 164,384, and to compare this information with data derived from a phase I study of ICI 182,780 in primary breast cancer patients [11]. In each instance, emphasis will be placed on immunohistochemical data as it was our original hope that such an approach would facilitate an assessment of the degree to which pure antioestrogens were fulfilling their potential as complete antagonists of oestrogen action in clinical breast cancer and thereby aid in defining the importance of oestrogens in the regulation of breast cancer growth.

Figure 1 shows the structure of ICI 164,384 and ICI 182,780 which are 7α long-chain analogues of oestradiol. The ER binding affinity and potency of ICI 182,780 are greater than that observed for ICI 164,384 due to the substitution of the amide function by a sulphoxide group and the fluorination of the terminal chain [12]. Such differences, however, do not alter the intrinsic biological behaviour of the drugs which are identical to other pure antioestrogens, based on substitutions in the oestradiol nucleus [13,14] or nonsteroidal forms [15].

Properties of pure antioestrogens in vitro

One of the most important early observations arising from the functional disablement of ER signalling by pure antioestrogens in oestrogen-sensitive human breast cancer cell lines was that treated cells frequently became very efficiently growth-arrested [10,16,17]. This property is illustrated in Fig. 2a and shows the growth of MCF-7 cells in 10⁻⁹M oestradiol in the presence or absence of a 100-fold excess of ICI

Fig. 1. Structures of ICI 164,384 and ICI 182,780.



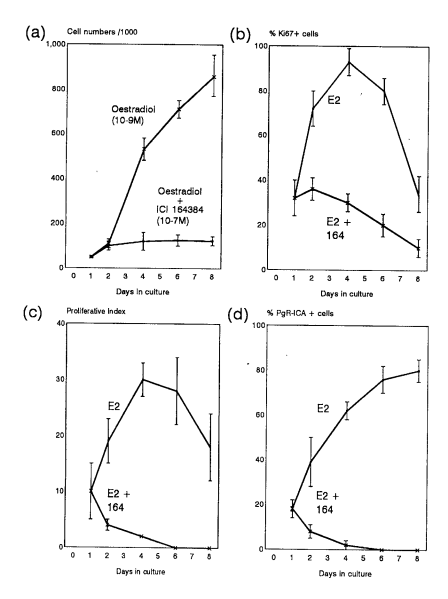


Fig. 2. Growth and immunohistochemical characterisation of MCF-7 cells. The cells were grown in white RPMI tissure culture medium with 5% DCC-stripped FCS (medium A) containing oestradiol \pm ICI 164,384. (a) Cell numbers were assessed using a Coulter Counter; (b,c) Ki67; and (d) PR assays were performed according to the methods of Bouzubar et al. [19] and Walker et al. [20], respectively. The Ki67 proliferative index was calculated as the proportion of cells showing intense nucleoplasmic and nucleolar staining patterns [21]. Results are shown as the mean \pm SD of six replicates.

164,384. In contrast to the expansion of the cell population that occurs in the presence of the steroid, the pure antioestrogen virtually abolishes the growth of the MCF-7 cells, allowing at best one doubling of the initial cell number with the cells



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