Updated Programme

DEVELOPING MEDICINES FOR COMPANION ANIMALS

Three Day Course with a Residential Option

Key topics to be covered on this three day course:

- EU Regulatory Framework
- Product Development
- Process Scale Up and Validation
- How to 'Block Out Studies'
- Part III: the Toxicological Package
- User Safety: Risk Assessment
- Environmental Risk Assessment

- 'Safety' DACS
- Pre-Clinical Development
- Clinical Trial Applications
- Clinical Development
- Target Species Tolerance
- 'Clinical' DACS
- Regulatory Strategies and Procedures
- Writing the Regulatory Submission

Chairman:

Ray Harding Consultant, Cyton Biosciences Ltd

Tutors:

Julian Braidwood Managing Director, Triveritas Ltd

Dr Raymond Munden Pharmaceutical Consultant



28 February, 1 & 2 March 2011 The Rembrandt Hotel, London

INTRODUCTION AND OBJECTIVES

The development of a veterinary medicine is complex, time consuming and expensive. It requires teamwork from individuals with different scientific training and a wide range of skills. Everyone involved must be aware of the main stages in the development programme and be able to relate his or her responsibilities to the expertise and needs of the scientists and commercial members of the team.

This course has been designed to demonstrate how pharmaceutical, pharmacological, toxicological and clinical investigations and regulatory management are brought together in the development programme. It will take the participants through all of the stages in the development of a veterinary medicinal product for companion animals, for which a marketing authorisation is sought in the European Union. Presentations will cover pharmaceutical and process development, toxicological, pharmacological, safety risk assessments, clinical development and regulatory submissions. An important part of the course will be devoted to working on case studies in the workshop sessions.

The course will provide a comprehensive introduction to the entire development programme for companion animal medicines.

WHO SHOULD ATTEND?

The course has been designed for anyone who has limited experience in only one of the disciplines in veterinary medicine development, such as pharmaceutical or analytical development, clinical trials, regulatory affairs or quality assurance. Anyone learning the role of project manager, as well as more experienced personnel seeking to review special problems encountered in product development, will benefit from the comprehensive programme delivered by experienced professionals. There will be ample opportunity for informal discussion during the proceedings.

If possible please bring a laptop with you to use during the workshop sessions.

CHAIRMAN

Ray Harding, Consultant, Cyton Biosciences Limited Ray established Cyton Biosciences Limited in 1997 to provide specialist services in new product development and registration for the bioscience industries in Europe. After managing the company since it was founded, he has now taken the role of consultant.

TUTORS

Julian Braidwood, Managing Director of Triveritas which is a contract product development company offering a comprehensive range of services to the Animal Health industry and related companies.

Dr Raymond Munden, has over 30 years experience in pharmaceutical research and development and was formerly Head, Analytical Services Europe for GlaxoSmithKline.

Day One

28 February 2011

Session One: Tutor - Ray Harding

09.30 Introduction and Objectives of the Course

09.45 ► EU Regulatory Framework

- · Understanding the regulatory objectives
- EU legal framework
- Legal base of regulatory procedures and dossier requirements
- 10.30 **▶ Coffee**
- 10.45 ► EU Regulatory Framework (cont.)

11.30 Assessing Development Candidates

· New chemical entities and older products

11.45 Project Team: R&D and Marketing

- · Defining marketing objectives
- · Costs and profitability
- · The Project Manager and Project Team
- 12.15 Discussion
- 12.30 **Lunch**

Session Two: Tutor - Dr Raymond Munden

13.30 Product Development

- Physical and chemical characteristics of the drug substance
- Process development
- Requisites of an ideal product
- Formulation development
- Optimising performance of problem drug substances
- Analytical development and setting specifications
- · Stability testing

15.15 **Tea**

15.30 Process Scale up and Validation

- Effect of product type
- Facilities/Personnel
- Regulatory issues

16.30 Workshop: Understanding How to "Block Out Studies"

- · Pharmaceutical studies
- Toxicological studies
- · Pharmacokinetic studies
- · Clinical studies

Led by Ray Harding

17.15 **End of Day One**

19.30 Conference Dinner at a Local Restaurant

Day Two	1 March 2011

Session Three: Tutor: - Ray Harding

09.00 Part III: The Toxicological Package

- · The objective of Part III
- The data required

09.45 User Safety: Risk Assessment

- · Reviewing toxicology studies
- · Setting the scenario
- Risk assessment
- Risk management

10.30 Pharmacokinetics and Bioequivalence

- Role of pharmacokinetic studies in product development
- · Bioequivalence studies
- Exemptions from bioequivalence studies
- 11.00 **Coffee**

11.15 Environmental Risk Assessment

• Phase I Assessment

11.30 • 'Safety' Detailed and Critical Summary

12.00 ► Introduction to Workshop

- Objectives of the workshop and review case studies
- Planning approach to workshop and workshop teams
- 12.30 **Lunch**

Session Four: Tutor - Ray Harding

- 13.30 Workshop Session
- 15.15 **Tea**

15.30 Planning Pre-Clinical and Clinical Development

- · Preparing pilot batches
- · PKs and target animal tolerance
- · Approaches to dose selection
- · Clinical trial applications

16.00 EU Regulatory Strategies and Procedures

- Full and abbreviated applications
- · Generic applications
- · Centralised Procedure
- Decentralised, MRP and National Procedures

17.00 **▶ End of Day Two**

Day Three 2 March 2011

Session Five: Tutor: - Julian Braidwood

09.00 Clinical Development

- Selecting a dose
- · Field studies
 - Multi-centred
 - Controlled
 - Randomised
 - Blinded
- · Efficacy guidelines
- 10.30 **▶ Coffee**

PROGRAMM

10.45 ► **Target Species Tolerance**

11.15 • 'Clinical' Detailed and Critical Summary

- Pre-clinical, clinical, PK and clinical data
- · Target species tolerance
- · Risk/benefit ratio

12.00 **Lunch**

Session Six: Tutor - Ray Harding

13.00 Final Workshop Session

14.30 Workshop Presentations

- · Presentation by each team
- · Review and discussion
- 15.30 **► Tea**

15.45 Writing the Regulatory Submission

- Writing the dossier
- Summary of Product Characteristics and labelling
- Working with Writers of Detailed and Critical Summaries
- 16.15 **▶ Discussion**
- 16.30 Concluding Remarks
- 16.45 **End of Course**

Register on-line at www.management-forum.co.uk or telephone +44 (0) 1483 730071

A Certificate of Attendance for Professional Development will be given to each participant who completes the course.



DEVELOPING MEDICINES FOR **COMPANION ANIMALS**



Application

28 February, 1-2 March 2011, Conf. No. V2-8011

Please PRINT your details:	
Title	
Position	
Department	
Company	
Company VAT No.	
Address	
CityPost Code	
Country	
Tel No Fax No	
E-mail	
Secretary's name	
Signature	
Substitutions may be made at any time at no extra charge Payment by either: VISA MASTERCARD AMEX	
Card No.	
Card Security No. AMEX	
Expiry date/	
☐ Cheque enclosed payable to Management Forum Limited ☐ Bank transfer on receipt of invoice ₩	
+44 (0) 1483 730008 Register	
Maybury Road, Woking, Surrey GU21 5JL, UK	
www.management-forum.co.uk	
E-mail: registrations@management-forum.co.uk If you have NOT received confirmation seven days after	
registering, please contact Registration Department.	

REGISTRATION INFORMATION

Registration Information

Dates 28 February 1- 2 March 2011

Times 28 February 2011 Start 09.30 - Finish 17.15 28 February 2011 **Conference Dinner Start 19.30** 1 March 2011 Start 09.00 - Finish 17.00

2 March 2011 Start 09.00 - Finish 16.45

Registration & Coffee

28 February 2011 09.00

Venue

The Rembrandt Hotel, 11 Thurloe Place, London SW7.

Opposite V&A Museum.

Nearest Underground station: South Kensington. Map available on Website under Hotels and Venues.

Accommodation

A limited number of bedrooms have been reserved at The Rembrandt Hotel, 11 Thurloe Place, London SW7, at a special rate of £133.92 (Superior) £152.18 (Executive) both including English breakfast, excluding VAT – subject to availability.

Hotel Tel: +44(0)20 7589 8100. Hotel Fax: +44(0)20 7225 3476.

Email: reservations_rembrandt@sarova.co.uk

All bookings should be made directly with the hotel or online at www.sarova.com/rembrandt, quoting promo code 'manforum'.

Conference Fee: (please tick)

Non Residential Fee (excluding conference dinner):

£1.650 + VAT

Residential Fee (including two nights accommodation and all meals including conference dinner) £2,100 + VAT

Conference No. V2-8011

Discounted Rates

Available on application for personnel from non-profit making organisations and registered charities. Group discount available on request.

Cancellation Policy:

Over 14 days prior to the Seminar: Cancellation fee of £75. 7/14 days prior to the Seminar: 50% of the fee. Fewer than 7 days or if no notification received: Registrant liable to pay FULL seminar fee.

NB: Cancellations must be received in writing by registrations@management-forum.co.uk.

In the event of circumstances beyond its control, Management Forum reserves the right to alter the programme, the speakers, the date or the venue.

If you do not want to receive future mailings from Management Forum please contact nick@management-forum.co.uk

If you do not wish to receive selected third party mailings please contact nick@management-forum.co.uk

Exhibition spaces and promotional opportunities will be available at this meeting. For further information please contact Judith Black (email: judith.black@management-forum.co.uk)