

Drug Master File Procedures in the EU, the US and Japan

Taking into account the guidance on elemental (ICH Q3D) and genotoxic (ICH M7) impurities

Speakers



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22-23 November 2018, Berlin, Germany

HIGHLIGHTS:

- Requirements of the European ASMF procedure
- Different types of Drug Master Files in the US
- How to document drug substance stability
- Compiling data for residual solvents and impurities taking into account metal and genotoxic impurities
- Special aspects of Drug Master Files in Japan
- Handling changes in European, US and Japanese Drug Master Files
- Maintaining Drug Master Files
- Comparison of ASMF and CEP procedure



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Objectives

This education course is intended to provide guidance on the procedures for the European ASMF, the US-DMF and the Japanese DMF. You will get to know

- how to describe manufacturing processes
- how to compile data for drug substance stability, impurities and residual solvents
- which are the important points to consider for US-DMFs
- which are the requirements for Japanese DMFs
- how to handle changes in European, US and Japanese DMFs
- which are the major differences and advantages of the ASMF and CEP procedure

Participants will have the opportunity to take part in one of two parallel workshops about

- 1. Description of the manufacturing process or
- 2. Managing changes in Drug Master Files

Background

Documentation of the drug substance quality is an integral part of any marketing authorisation application. In Europe the most common document for this purpose is the Active Substance Master File (ASMF) as long as the applicant has no Certificate of Suitability of the pharmacopoeial monograph (CEP). The European ASMF procedure differs significantly from the US-DMF procedure and for strategic reasons it is very important to take these differences into account. Moreover, there are particular requirements for DMFs in Japan. For global acting companies it is a big challenge to handle the different procedures of compiling, submitting, changing and maintaining Drug Master Files in an efficient way.

Target Audience

The education course is designed for all persons involved in the compilation of pharmaceutical dossiers for marketing authorisations, especially for Drug Master Files, who want to become familiar with the different DMF procedures. Furthermore, the course will be of interest to personnel from Quality Units of the pharmaceutical and the API industry.

Programme

The European Active Substance Master File procedure - An Introduction

- Regulatory background and Scope
- The revised ASMF guideline
- Open and closed parts points to consider
- Comparison of ASMF and CEP procedure

Drug Master File Procedures in the US

- Types of Drug Master Files
- Drug Master Files under GDUFA
- Submissions of DMFs
- Holder obligations
- Maintenance of Drug Master Files
- US vs EU DMF differences in the procedure

How to document drug substance stability

- Stability Guidelines
- Stability Testing of new drug substances and drug products
- Storage Conditions
- Bracketing and Matrixing Designs
- Stability data from new drug dosage forms
- How to document evaluation of stability data
- Optimising the submission

Residual solvents and Impurities: synthesis derived Impurities, Metals and genotoxic Impurities

- Guidelines
- Impact of the new guidelines ICH Q3D and ICH M7
- Sources of Impurities
- Setting and justification of specifications
- Residual solvents, solvent classes
- Content and scope of data documentation requirements
- Frequent mistakes

Parallel Workshops

Please choose one out of two Parallel Workshops:

Description of the Active Substance manufacturing process Managing changes in Drug Master Files -Case Studies-

Post Approval Changes in the US

- Post approval activities
- Reporting requirements to the FDA (CBE 0, CBE 30, Annual Report)
- Post approval commitments and post approval reporting requirements
- Risk evaluation and mitigation strategies (REMS)

Handling Changes in European Drug Master Files

- Why is there a need for changes
- Types of changes
- How to communicate with the MA holders and how to get feed back
- Differences between ASMF and CEP
- When to implement a specific change
- Version management of the ASMF

Requirements of the Drug Master File Procedure in Japan

- Regulatory procedures in Japan:
 - Site accreditation | GMP paper-based inspection | Drug Master File
- Drug Master File format
- Specific points to consider for the J-DMF
- Communication with the Japanese authorities

Changes and Maintenance of Japanese Drug Master Files

- Change procedures and communication with the Japanese authority
- Types of changes
- Notification of changes

Comparison of the CEP and ASMF Procedure

- The certification scheme of the PhEur
- Advantages and disadvantages of the CEP procedure compared to the ASMF procedure
- Handling of variations in the CEP procedure
- Countries accepting CEPs

Speakers



Marieke van Dalen, Aspen Oss B.V., The Netherlands

Marieke van Dalen is the global regulatory specialist in the regulatory group dedicated to APIs, with almost 30 years of experience

in the regulatory field. She is a board member of APIC, the European API Industry organization, and she participates in the Japan task force, Emerging markets task force and the Quality metrics task force. She frequently represents APIC in meetings and conferences organized by EMA, EDQM, ICH etc.



Dr Hiltrud Horn, Managing Director of Horn Pharmaceutical Consulting, Germany

Dr Hiltrud Horn is managing director of HORN PHARMACEUTICAL CONSULTING

with focus on CMC, GMP and Regulatory Affairs (EU and US). She started in pharma industry in 1990 and held several managerial positions within Hoffmann-La Roche in Basel and Knoll (now Abbott) in Ludwigshafen with global responsibility within QC/QA/Regulatory Affairs / Project Management / Medical Writing. She was consultant for the biotech and life science industry at Cap Gemini Ernst & Young prior to starting her own business more than 13 years ago. She is pharmacist with a Ph.D. and holds a Diploma in Pharmaceutical Medicine (Basel). Furthermore she is specialised pharmacist for pharmaceutical analytics and for drug information.



Dr Usfeya A. Muazzam, Bonn, Germany

Dr. Usfeya A. Muazzam worked as Senior Assessor for Quality, Division: Quality, Department: Scientific Quality Assurance, Staff Unit:

Strategy and Planning of BfArM. He left the agency in 2012. He is co-author of "Gute Regulatorische Praxis, Arzneimittelzulassung - Pharmazeutische Qualität", Wissenschaftliche Verlagsgesellschaft, Stuttgart, Germany and "Guide to Drug Regulatory Affairs", Editio Cantor Verlag, Aulendorf, Germany.



Dr Boris Pimentel, Antibioticos de León,

Dr Pimentel is Quality Director of Antibioticos de León in Spain. Before he joined ADL he was manager of the consulting company

for global regulatory services –Pi-Consulting in Switzerland. Before that he worked on the Dutch company DSM Nutritional Products in Switzerland, focusing in Pharma and Food regulations. Since 2010 as a member of the APIC board he chaired the task force for Japan regulations, and Emerging Market Regulations.



Dr Wilhelm Schlumbohm, Berlin, Germany Dr Schlumbohm worked more than 25 years with German drug licensing authorities in the field of assessment of the CMC parts of new drug applications. He is a member of the

Working Group on Active Substance Master File procedures.

Easy Registration









Date

Thursday, 22 November 2018, 9.00 h-17.45 h (Registration and coffee 8.30 h - 9.00) Friday, 23 November 2018, 8.30 h-15.15 h

Venue

InterCityHotel Berlin Hauptbahnhof Katharina-Paulus-Straße 5 10557 Berlin, Germany Phone +49 (0)30 288 755 0

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berlin-hauptbahnhof@intercityhotel.de

Fees (per delegate plus VAT)

ECA Members: € 1,590 APIC Members: € 1,690 EU GMP Inspectorates: € 895 Non-ECA Members: € 1,790

The conference fee is payable in advance after receipt of invoice and includes conference documentation, dinner on the first day, lunch on both days and all refreshments.

VAT is reclaimable.

Accommodation

CONCEPT HEIDELBERG CONCEPT has reserved a limited number of rooms in the conference hotel. You will receive a room reservation form/POG when you have registered for the conference. Please use this form for your room reservation to receive the specially negotiated rate for the duration of your stay. Reservation should be made directly with the hotel. Early reservation is recommended.

Conference language

The official conference language will be English.

Organisation and Contact

ECA has entrusted Concept Heidelberg with the organisation of this event.

CONCEPT HEIDELBERG

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For questions regarding content:

Dr Gerhard Becker (Operations Director) at +49-62 21/84 44 65 or per e-mail at becker@concept-heidelberg.de.

For questions regarding reservation, hotel, organisation etc.: Ms Susanne Ludwig (Organisation Manager) at +49-62 21/84 44 44,

or per e-mail at ludwig@concept-heidelberg.de.



Social Event

In the evening of the first conference day, you are cordially invited to a social event. This is an excellent opportunity to share your experiences with colleagues from other companies in a relaxed atmosphere.

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