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Q5C	Quality of Biotechnological Products: Stability Testing of Biotechnological/Biological Products	Nov. 1995			
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Q6A	Specifications: Test Procedures and Acceptance Criteria for New Drug Substances and New Drug Products: Chemical Substances	Oct. 1999			
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	BATCH S: Safety				
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S1A	Guideline on the Need for Carcinogenicity Studies of Pharmaceuticals	Nov. 1995			
S1B	Testing for Carcinogenicity of Pharmaceuticals	July 1997			
S1C(R2)	Dose Selection for Carcinogenicity Studies of Pharmaceuticals	March 2008			
S2(R1)	Guidance on Genotoxicity Testing and Data Interpretation for Pharmaceuticals Intended for Human Use	Nov 2011			
S3A	Note for Guidance on Toxicokinetics: The Assessment of Systemic Exposure in Toxicity Studies	Oct. 1994			
S3B	Pharmacokinetics: Guidance for Repeated Dose Tissue Distribution Studies	Oct. 1994			
S4	Duration of Chronic Toxicity Testing in Animals (Rodent and Non Rodent Toxicity Testing)	Sept. 1998			
S5(R2)	Detection of Toxicity to Reproduction for Medicinal Products and Toxicity to Male Fertility (the Addendum dated November 1995 has been incorporated into the core guideline in November 2005)	June 1993			
S6(R1)	Preclinical Safety Evaluation of Biotechnology-Derived Pharmaceuticals	June 2011			
S7A	Safety Pharmacology Studies for Human Pharmaceuticals	Nov 2000			
S7B	The Non-clinical Evaluation of the Potential for Delayed Ventricular Repolarization (QT Interval Prolongation) by Human Pharmaceuticals	May 2005			
S8	Immunotoxicity Studies for Human Pharmaceuticals	Sept. 2005			
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	BATCH E: Efficacy				
	Finalised Guidelines (Step 4)				
E1	The Extent of Population Exposure to Assess Clinical Safety for Drugs Intended for Long- Term Treatment of Non-Life-Threatening Conditions	Oct. 1994			
E2A	Clinical Safety Data Management: Definitions and Standards for Expedited Reporting	Oct. 1994			
E2B(R2)	Clinical Safety Data Management: Data Elements for Transmission of Individual Case Safety Reports (This guideline is re-opened for revision under Step 2. See E2B(R3)).	Feb. 2001			
E2C(R1)	Clinical Safety Data Management: Periodic Safety Update Reports for Marketed Drugs (<i>The Addendum dated February 2003 has been incorporated into the core guideline in November 2005</i>).	Nov. 1996			
E2D	Post-Approval Safety Data Management: Definitions and Standards for Expedited Reporting	Nov. 2003			
E2E	Pharmacovigilance Planning	Nov. 2004			
E2F	Development Safety Update Report	Aug. 2010			
E3	Structure and Content of Clinical Study Reports	Nov. 1995			
E4	Dose-Response Information to Support Drug Registration	March 1994			
E5(R1)	Ethnic Factors in the Acceptability of Foreign Clinical Data	March 1998			
E6(R1)	Good Clinical Practice: Consolidated Guideline	May 1996			
E7	Studies in Support of Special Populations: Geriatrics	June 1993			
E7 Q&A	Studies in Support of Special Populations: Geriatrics Questions & Answers	July 2010			
E8	General Considerations for Clinical Trials	July 1997			
E9	Statistical Principles for Clinical Trials	Feb. 1998			
E10	Choice of Control Group and Related Issues in Clinical Trials	July 2000			
E11	Clinical Investigation of Medicinal Products in the Pediatric Population	July 2000			
E14	The Clinical Evaluation of QT/QTc Interval Prolongation and Proarrhythmic Potential for Non-Antiarrhythmic Drugs	May 2005			
E15	Definitions for Genomic Biomarkers, Pharmacogenomics, Pharmacogenetics, Genomic Data and Sample Coding Categories	Nov. 2007			
E16	Biomarkers Related to Drug or Biotechnology Product Development: Context, Structure and Format of Qualification Submissions	Aug. 2010			
	Documents released for consultation (Step 2)				
E2B(R3) Implementation Guide	Implementation Guide for Electronic Transmission of Individual Case Safety Reports (ICSRs), Data Elements and Message Specification including key parts of the updated E2B(R3) Guideline	June 2011			
E2C(R2)	Periodic Benefit-Risk Evaluation Report (PBRER)	Feb. 2012			
Consensus Draft Principle					
E12	Principles for Clinical Evaluation of New Antihypertensive Drugs	March 2000			

BATCH M: Multidisciplinary			
Finalised Guidelines (Step 4)			
M2 ICSR (R2)	Electronic Transmission of Individual Case Safety Reports Message Specification (ICH ICSR DTD Version 2.1) companion document to E2B(R3)	Feb. 2001	
M3(R2)	Guidance on Nonclinical Safety Studies for the Conduct of Human Clinical Trials and Marketing Authorization for Pharmaceuticals	June 2009	
M3(R2) Q&As	Guidance on Nonclinical Safety Studies for the Conduct of Human Clinical Trials and Marketing Authorization for Pharmaceuticals Questions & Answers	June 2011	
M4(R3)*	Organisation of the Common Technical Document for the Registration of Pharmaceuticals for Human Use (<i>Edited with Numbering and Section Header Changes, September 2002</i>) Including the Annex: the Granularity Document (Revised November 2003).	Nov 2000	
M4Q(R1)*	The Common Technical Document for the Registration of Pharmaceuticals for Human Use: Quality (Edited with Numbering and Section Header Changes, September 2002)	Nov 2000	
M4S(R2)*	The Common Technical Document for the Registration of Pharmaceuticals for Human Use: Safety (Edited with Numbering and Section Header Changes, September 2002)	Nov 2000	
M4E(R1)*	The Common Technical Document for the Registration of Pharmaceuticals for Human Use: Efficacy (Edited with Numbering and Section Header Changes, September 2002)	Nov 2000	
Guidelines released for consultation (Step 2)			

* Notice for Clarification:

Within the ICH regions, local versions are published. The wording of the core CTD (Modules 2, 3, 4 and 5) in the local versions might be slightly different from one region to another due to specific editing that takes into consideration regional regulations. It does not affect the common understanding by the six ICH parties of the CTD published on the ICH website (http://www.ich.org).

Questions & Answers:

In order to facilitate the implementation of the CTD, the ICH Experts have developed a series of Q&As which are continuously updated and can be downloaded from the ICH website directly from the following url: http://www.ich.org/products/ctd.html